PV-WAVE 7.5

New Features Guide

Helping Customers SOLVE Complex Problems
The Visualization Toolkit

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# Table of Contents

**Preface**  
ix  
What’s in this Manual  ix  
Conventions Used in this Manual  x  
Technical Support  xi  

**Chapter 1: New Features Introduction**  
PV-WAVE 7.5 Major Enhancements  1  

**Chapter 2: New Commands**  
New PV-WAVE Commands  5  
AFFINE Function  5  
BLOB Function  6  
BLOBCOUNT Function  7  
BOUNDARY Function  8  
CPROD Function  9  
DERIVN Function  11  
DICM_TAG_INFO Function  12  
EUCLIDEAN Function  13  
EXPAND Function  14  
EXPON Function  16  
EXTREMA Function  17  
FACTOR Function  18  
GCD Function  19  
GREAT_INT Function  20
GRIDN Function 21
HISTN Function 22
INDEX_AND Function 24
INDEX_OR Function 25
INTERPOLATE Function 26
INTRP Function 27
JACOBIAN Function 28
LCM Function 29
LISTARR Function 30
MINIMIZE Function 31
MOLEC Function 32
MOMENT Function 33
NEIGHBORS Function 34
NORMALS Function 35
PRIME Function 36
PRODUCT Function 37
RENDER24 Function 38
REPLV Function 39
RESAMP Function 40
SAME Function 41
SGN Function 42
SHIF Function 43
SLICE Function 44
SMALL_INT Function 45
SORTN Function 46
vtkADDATTRIBUTE Procedure 47
vtkAXES Procedure 48
<table>
<thead>
<tr>
<th>Procedure</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>vtkCAMERA Procedure</td>
<td>50</td>
</tr>
<tr>
<td>vtkCLOSE Procedure</td>
<td>51</td>
</tr>
<tr>
<td>vtkCOLORBAR Procedure</td>
<td>52</td>
</tr>
<tr>
<td>vtkCOMMAND Procedure</td>
<td>53</td>
</tr>
<tr>
<td>vtkERASE Procedure</td>
<td>54</td>
</tr>
<tr>
<td>vtkGRID Procedure</td>
<td>55</td>
</tr>
<tr>
<td>vtkHEDGEHOG Procedure</td>
<td>56</td>
</tr>
<tr>
<td>vtkINIT Procedure</td>
<td>58</td>
</tr>
<tr>
<td>vtkLIGHT Procedure</td>
<td>59</td>
</tr>
<tr>
<td>vtkPLOTS Procedure</td>
<td>60</td>
</tr>
<tr>
<td>vtkPOLYDATA Procedure</td>
<td>62</td>
</tr>
<tr>
<td>vtkPOLYSHADE Procedure</td>
<td>63</td>
</tr>
<tr>
<td>vtkPPMREAD Function</td>
<td>65</td>
</tr>
<tr>
<td>vtkPPMWRITE Procedure</td>
<td>66</td>
</tr>
<tr>
<td>vtkRECTILINEARGRID Procedure</td>
<td>67</td>
</tr>
<tr>
<td>vtkRENDERWINDOW Procedure</td>
<td>68</td>
</tr>
<tr>
<td>vtkSCATTER Procedure</td>
<td>70</td>
</tr>
<tr>
<td>vtkSLICEVOL Procedure</td>
<td>72</td>
</tr>
<tr>
<td>vtkSTRUCTUREDGRID Procedure</td>
<td>74</td>
</tr>
<tr>
<td>vtkSTRUCTUREDPOINTS Procedure</td>
<td>75</td>
</tr>
<tr>
<td>vtkSURFACE Procedure</td>
<td>76</td>
</tr>
<tr>
<td>vtkSURFGEN Procedure</td>
<td>79</td>
</tr>
<tr>
<td>vtkTEXT Procedure</td>
<td>81</td>
</tr>
<tr>
<td>vtkTVRD Function</td>
<td>82</td>
</tr>
<tr>
<td>vtkUNSTRUCTUREDGRID Procedure</td>
<td>83</td>
</tr>
<tr>
<td>vtkWDELETE Procedure</td>
<td>84</td>
</tr>
<tr>
<td>vtkWINDOW Procedure</td>
<td>85</td>
</tr>
</tbody>
</table>
vtkWRITEVRML Procedure 87
vtkWSET Procedure 88
WgOrbit Procedure 89
WIN32_PICK_PRINTER Function 90

**New PV-WAVE:Database Connection Functions  91**
DB_GET_BINARY Function  91
NULL_PROCESSOR Function  92

**New PV-WAVE:IMSL Mathematics Commands  95**

Chapter 4: Quadrature  95
INTFCN_QMC Function 95

Chapter 9: Special Functions  97
CUM_INTR Function 97
CUM_PRINC Function 99
DEPRECIATION_DB Function 100
DEPRECIATION_DDB Function 102
DEPRECIATION_SLN Function 104
DEPRECIATION_SYD Function 105
DEPRECIATION_VDB Function 106
DOLLAR_DECIMAL Function 108
DOLLAR_FRACTION Function 109
EFFECTIVE_RATE Function 110
FUTURE_VALUE Function 111
FUTURE_VAL_SCHD Function 113
INT_PAYMENT Function 114
INT_RATE_ANNUITY Function 115
INT_RATE_RETURN Function 117
INT_RATE_SCHD Function 118
MOD_INTERN_RATE Function 120
NET_PRES_VALUE Function 121
NOMINAL_RATE Function 123
NUM_PERIODS Function 124
PAYMENT Function 125
PRESENT_VALUE Function 127
PRES_VAL_SCHD Function 128
PRINC_PAYMENT Function 130
ACCR_INT_MAT Function 131
ACCR_INT_PER Function 133
BOND_EQV_YIELD Function 135
CONVEXITY Function 137
COUPON_DAYS Function 139
COUPON_NUM Function 141
SETTLEMENT_DB Function 143
COUPON_DNC Function 145
DEPREC_AMORDEGRC Function 147
DEPREC_AMORLINC Function 148
DISCOUNT_PR Function 150
DISCOUNT_RT Function 152
DISCOUNT_YLD Function 154
DURATION Function 157
INT_RATE_SEC Function 159
DURATION_MAC Function 161
COUPON_NCD Function 163
COUPON_PCD Function 165
PRICE_PERIODIC Function 167
PRICE_MATURITY Function 170
MATURITY_REC Function 172
TBILL_PRICE Function 174
TBILL_YIELD Function 175
YEAR_FRACTION Function 176
YIELD_MATURITY Function 178
YIELD_PERIODIC Function 180

Chapter 10: Basic Statistics and Random Number Generation 183
FAURE_INIT Function 184
FAURE_NEXT_PT Function 187

New PV-WAVE:IMSL Statistics Commands 190

Chapter 8: Time Series and Forecasting 190
KALMAN Procedure 190

Chapter 11: Probability Distribution Functions and Inverses 200
BINOMIALPDF Function 200

Chapter 12: Random Number Generation 201
RANDOM_TABLE Procedure 201
RANDOM_NPP Function 205
RANDOM_ORDER Function 209
RAND_TABLE_2WAY Function 210
RAND_ORTH_MAT Function 212
RANDOM_SAMPLE Function 213
RAND_FROM_DATA Function 216
CONT_TABLE Procedure 218
RAND_GEN_CONT Function 219
DISCR_TABLE Function 221
RAND_GEN_DISCR Function 225
Chapter 3: Updates to Existing Functionality  239

Updated PV-WAVE Functions and Procedures  239
  JOURNAL Procedure  239
  NoBlock Keyword for PV-WAVE VDA Tool Procedures  240
  YLabelCenter Keyword  240
  !Version System Variable  240

Updated PV-WAVE:IMSL Statistics Functions  241
  RANDOM Function  241
  RANDOMOPT Function  241

Updated PV-WAVE:IMSL Mathematics Functions  242
  RANDOM Function  242
  RANDOMOPT Function  242

Index
Preface

This *New Features Guide* provides detailed information on major features added to PV-WAVE, PV-WAVE:IMSL Mathematics, and PV-WAVE:IMSL Statistics since version 7.0.

For additional information on this release, see the Release Notes, available in the PV-WAVE installation directory or on our Web site: www.vni.com.

**What’s in this Manual**


**Chapter 3: Updates to Existing Functionality** — Describes additions to previously existing functionality. This chapter already is integrated into the PV-WAVE Reference, PV-WAVE:IMSL Statistics Reference, and PV-WAVE:IMSL Mathematics Reference.

**Index** — Contains page references for an alphabetical list of subjects described in this guide.
You will find the following conventions used throughout this manual:

- Code examples appear in this typeface. For example:

  \begin{verbatim}
  PLOT, temp, s02, Title = 'Air Quality'
  \end{verbatim}

- Code comments are shown in this typeface, immediately below the commands they describe. For example:

  \begin{verbatim}
  PLOT, temp, s02, Title = 'Air Quality'
  ; This command plots air temperature data vs. sulphur
dioxide concentration.
  \end{verbatim}

- Variables are shown in lowercase italics ($myvar$), function and procedure names are shown in uppercase (XYOUTS), keywords are shown in mixed case italic ($XTitle$), and system variables are shown in regular mixed case type ($!Version$). For better readability, all GUI development routines are shown in mixed case (WwMainMenu).

- A $ at the end of a line of PV-WAVE code indicates that the current statement is continued on the following line. By convention, use of the continuation character ($) in this document reflects its syntactically correct use in PV-WAVE. This means, for instance, that strings are never split onto two lines without the addition of the string concatenation operator (+). For example, the following lines would produce an error if entered literally in PV-WAVE.

  \begin{verbatim}
  WAVE> PLOT, x, y, Title = 'Average $Air Temperatures by Two-Hour Periods'
  ; Note that the string is split onto two lines; an error
  ; message is displayed if you enter a string this way.
  \end{verbatim}

  The correct way to enter these lines is:

  \begin{verbatim}
  WAVE> PLOT, x, y, Title = 'Average ' + $'Air Temperatures by Two-Hour Periods'
  ; This is the correct way to split a string onto two
  ; command lines.
  \end{verbatim}

- Reserved words, such as FOR, IF, CASE, are always shown in uppercase.
Technical Support

If you have problems installing, unlocking, or running your software, contact Visual Numerics Technical Support by calling:

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Users outside the U.S., France, Germany, Japan, Korea, Mexico, Taiwan, and the U.K. can contact their local agents.

Please be prepared to provide the following information when you call for consultation during Visual Numerics business hours:

- Your license number, a six-digit number that can be found on the packing slip accompanying this order. (If you are evaluating the software, just mention that you are from an evaluation site.)
- The name and version number of the product. For example, PV•WAVE 7.5.
- The type of system on which the software is being run. For example, SPARC-station, IBM RS/6000, HP 9000 Series 700.
- The operating system and version number. For example, HP-UX 10.2 or IRIX 6.5.
- A detailed description of the problem.
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New Features Introduction

This chapter provides an introduction to the scope of the PV-WAVE 7.5 release, along with user information for major enhancements added to PV-WAVE since version 7.0.

For information on specific new PV-WAVE functions and procedures, see New PV-WAVE Commands in Chapter 2, New Commands.

For information on updates to existing PV-WAVE functionality, see Chapter 3, Updates to Existing Functionality.

PV-WAVE 7.5 Major Enhancements

PV-WAVE VTK Integration

PV-WAVE has been updated to include the integration of the Visualization Toolkit (VTK) to make all of its functionality available to PV-WAVE users. PV-WAVE users can now create high quality, interactive graphics through the use of the PV-WAVE link to the Visualization Toolkit (VTK). The Visualization Toolkit is an Open Source toolkit for creating both simple and complex visualizations in 3D using OpenGL for high-performance, accelerated graphics. The Visualization Toolkit and PV-WAVE complement each other well. PV-WAVE excels at data access, data manipulation, numerical algorithms, data filtering, user-interface development, and many interactive 2D graphical tasks. The Visualization Toolkit is a best-of-breed tool for creating complex 3D visualizations. Together they provide a simple and quick way to build tools for Visual Data Analysis. For detailed
information on the new PV-WAVE-VTK functions, see New PV-WAVE Commands in Chapter 2, New Commands.

**PV-WAVE: IMSL Mathematics and Statistics**

PV-WAVE: IMSL Mathematics and PV-WAVE: IMSL Statistics have been updated to include a complete implementation of Visual Numeric’s IMSL C Numerical Library 5.0.

*New Functionality includes:*

- A new time series routine useful in navigation, surveying, vehicle tracking (aircraft, spacecraft, missiles), geology, oceanography, fluid dynamics, steel/paper/power industries, and demographic estimation.
- 50 new functions in the area of finance and bonds.
- Routines to compute low discrepancy series of points using a generalized Faure sequence.
- A new algorithm for efficient multi-dimensional quadrature.
- More than twenty new random number routines, including Generalized Feedback Shift Register (GFSR) generator support.

This integration represents a major enhancement to the WAVE Family. For detailed descriptions of the new mathematics and statistics routines, see New PV-WAVE:IMSL Mathematics Commands and New PV-WAVE:IMSL Statistics Commands in Chapter 2, New Commands. For of the new mathematics and statistics functionality, see the PV-WAVE:IMSL Statistics Reference and the PV-WAVE:IMSL Mathematics Reference.

**PV-WAVE: Database Connection**

*New functionality includes:*

- **DB_GET_BINARY Function** - Returns binary large objects (BLOBS) from a DBMS (database management system) server.
- **NULL_PROCESSOR Function** - Facilitates the use of the resulting output from the Null_Info keyword for the DB_SQL function by extracting locations of NULL values in the database query.
- **Keyword - null_info_object** to DB_SQL function, which provides information on the location of NULL data in the result from a database query. For more information on keyword null_info_object, see NULL_PROCESSOR Function on page 92.
For more detailed information on the new database connection functions, see *New PV-WAVE: Database Connection Functions* on page 91.

**PV-WAVE: DICOM Reader**

PV-WAVE users can now read and write DICOM images with the implementation of IMAGE_READ and IMAGE_WRITE functions. For more details on IMAGE_READ and IMAGE_WRITE, see the *PV-WAVE Reference*.

*New Functionality includes:*

DICM_TAG_INFO Function — extracts the tag information from an image associative array that contains a DICOM image. For more details on DICM_TAG_INFO, see *New PV-WAVE Commands* on page 5 in Chapter 2, *New Commands*. 
New Commands

This chapter lists and describes new functions and procedures added to:

- PV-WAVE (page 5)
- PV-WAVE: Database Connection (page 91)
- PV-WAVE: IMSL Mathematics (see page 95)
- PV-WAVE: IMSL Statistics (page 190)

New PV-WAVE Commands

This section describes the new functions and procedures that have been added to PV-WAVE 7.5.

AFFINE Function

Standard Library function that applies an affine transformation to an array.

Usage

\[
\text{result} = \text{AFFINE}(a, b, [c])
\]

Input Parameters

- \(a\) — An n-dimensional array.
**b** — An invertible (n, n) array.

**c** — An n-element vector (optional).

**Returned Value**

**result** — An array representing \( a \) (and of the same dimensions as \( a \)) under the coordinate transformation \( y = b#x + c \), where \( y \) and \( x \) are coordinates for the \( result \) and for \( a \), respectively, which differ from array index coordinates by a simple translation to the array centroid.

**Keywords**

None.

**Example**

See `wave/lib/user/examples/affine_ex`.

**See Also**

ROT, ROTATE, ROT_INT

---

**BLOB Function**

Standard Library function that isolates a homogeneous region in an array.

**Usage**

\[
result = BLOB(a, i, b)
\]

**Input Parameters**

- **a** — An array of n dimensions.
- **i** — A vector of n integers giving a seed element for the region.
- **b** — A two-element vector giving bounds for values in the region.
**BLOBCOUNT Function**

Standard Library function that counts homogeneous regions in an array.

**Usage**

\[ \text{result} = \text{BLOBCOUNT}(a, b) \]

**Input Parameters**

- **a** — An array of \( n \) dimensions.
- **b** — A two-element vector of bounds for values in a region.

**Returned Value**

\( \text{result} \) — A list in which each element defines a distinct region whose values lie in the range \([b(0), b(1))\). \( \text{result}(j) \) is a \((m(j), n)\) array of \( m(j) \) \( n \)-dimensional indices into \( a \). If no such regions exist, then \( \text{result} \) is returned as \(-1\).
Keywords

$k$ — A positive integer (less than or equal to $n$) controlling connectivity: two cells are connected if they share a common boundary point, and if their centroids are within the square root of $k$ of each other. $k = 1$ by default, which implies that connected cells share a common face.

Example 1

```plaintext
a = ( image_read(!data_dir+'vni_small.tif') )('pixels')
a = bytscl( resamp(a,500,500) ) & tv, a
r = blobcount( a, [255,255] )
for i = 0, n_elements(r)-1 do a(index_conv(a,r(i))) = 50 & tv, a
```

Example 2

See `wave/lib/user/examples/blobcount_ex.pro`

See Also

BLOB, BOUNDARY, NEIGHBORS

---

**BOUNDARY Function**

Standard Library function that computes the boundary of a region in an array.

**Usage**

```plaintext
result = BOUNDARY(a,r)
```

**Input Parameters**

- $a$ — An array of $n$ dimensions.
- $r$ — A vector of indices defining the region of $a$.

**Returned Value**

- $result$ — A vector of indices defining the boundary of $r$. 
Keywords

\( k \) — A positive integer (less than or equal to \( n \)) defining connectivity. A boundary element of \( r \) is an element of \( r \) with neighbors not in \( r \); two array cells are neighbors if they share a common boundary point and their centroids are within the square root of \( k \) of each other. \( k = 1 \) by default, which implies neighbors share a common face.

Examples

```fortran
a = indgen( 5, 4 ) & pm, a
print, fix(boundary(a,[1,2,3,6,7,8,12,13]))
print, fix(boundary(a,[1,2,3,6,7,8,12,13],k=2))
a = bytscl( dist(500) ) & r = where( 150 le a and a le 200 )
a( boundary(a,r) ) = 0 & tv, a
```

See Also

BLOB, BLOBCOUNT, NEIGHBORS

---

**CPROD Function**

Standard Library function that returns the Cartesian product of some arrays.

**Usage**

\[ \text{result} = \text{CPROD}(a) \]

**Input Parameters**

\( a \) — A list of \( n \) arrays.

**Returned Value**

\( \text{result} \) — An \((m,n)\) array where \( \text{result}(i,*) \) is an element of the Cartesian product of the \( n \) arrays in \( a \), and where \( \text{result}(*,j) \) contains only elements from \( a(j) \); \( \text{result} \) is ordered so that \( \text{result}(*,j) \) cycles through the elements of \( a(j) \) in order, and does so faster than \( \text{result}(*,j+1) \) cycles through the elements of \( a(j+1) \).
Keywords
None.

Example
\texttt{pm, cprod( list([0,1], [0,1,2], [0,1,2,3]) )}

\textbf{CURVATURES Function}

Standard Library function that computes curvatures on a parametrically defined surface.

Usage
\[ c = \text{curvatures}(s) \]

Input Parameters
\( s \) — A 3-element list of 2-dimensional arrays of dimensions \( d \).

Returned Value
\( c \) — A 2-element list of 2-dimensional arrays of dimensions \( d \), where \( c(0) \) defines the distribution of minimum curvature and \( c(1) \) defines the distribution of maximum curvature.

Keyword
\( x \) - A 2-element list of vectors defining the independent variables. By default, \( x(i) = \text{findgen}(d(i)) \)

Example
See \texttt{wave/lib/user/examples/curvatures_ex.pro}.

See Also
EUCLIDEAN, JACOBIAN, NORMALS
**DERIVN Function**

Standard Library function that differentiates a function represented by an array.

**Usage**

\[ result = \text{DERIVN}(a, n) \]

**Input Parameters**

- \(a\) — An array of values of the dependent variable.
- \(n\) — An integer \((\geq 0)\) designating which dimension to differentiate.

**Returned Value**

\(result\) — An array of the same dimensions as \(a\), representing the derivative with respect to the \(n\)th independent variable.

**Keywords**

- \(x\) — A vector defining the independent variable of differentiation. \(x\) defaults to the indices into dimension \(n\) of \(a\).

**Examples**

- `pm, derivn( [0,2,1,0,1], 0 )`
- `pm, derivn( [[0,2,1,0,1],[0,1,0,0,2],[1,0,2,1,2]], 0 )`
- `pm, derivn( [[0,2,1,0,1],[2,1,0,2,0],[1,0,2,1,2]], 1 )`

**See Also**

DERIV
**DICM_TAG_INFO Function**

Extracts Digital Imaging and Communications in Medicine (DICOM) tags information from an image associative array.

**Usage**

\[ result = DICM\_TAG\_INFO (filename, image) \]

**Input Parameters**

- **filename** — On input, a string containing the name of the file which contains the descriptions for the DICOM tags.
- **image** — An associative array in image format.

**Returned Value**

- **result** — An associative array containing DICOM tags information.

**Discussion**

The DICM_TAG_INFO function extracts the tag information from an image associative array that contains a DICOM image. The tag information is returned as an associative array. The following table describes each key of the associative array:

<table>
<thead>
<tr>
<th>Array Key Name</th>
<th>Variable Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>tag</em></td>
<td>STRING</td>
<td>A 1-dimensional array containing the DICOM tags</td>
</tr>
<tr>
<td><em>description</em></td>
<td>STRING</td>
<td>A 1-dimensional array containing the DICOM tag descriptions</td>
</tr>
<tr>
<td><em>value</em></td>
<td>STRING</td>
<td>A 1-dimensional array containing the DICOM tag values</td>
</tr>
</tbody>
</table>

The DICM_TAG_INFO function needs a file containing the tag description as input. This file contains a tag followed by a description for this tag. The tags in this file must be in ascending order. For example:
Example

This example uses IMAGE_READ to read a DICOM image file. Then it extracts the DICOM tags and displays the information of the result variable.

```plaintext
e = euclidean (j)
```

**Input Parameters**

- `j` — A Jacobian defined by a `n`-element list of `m`-element lists of `m`-dimensional arrays of dimensions `d`.

**Returned Value**

- `e` — The Euclidean metric under a transformation with Jacobian `j`: an `m`-element list of `m`-element lists of `m`-dimensional arrays. `(e(p))(q)` is the `m`-dimensional array (of dimensions `d`) that represents the `(p, q)` component of the metric.
Keywords
None.

Example
See wave/lib/user/examples/euclidean_ex.pro.

See Also
CURVATURES, JACOBIAN, NORMALS

---

**EXPAND Function**

Standard Library function that expands an array into higher dimensions.

**Usage**

\[ result = \text{EXPAND}(a, d, i) \]

**Input Parameters**

- \( a \) — An array of \( n \) dimensions.
- \( d \) — A vector specifying the dimensions for the new array.
- \( i \) — A monotonically increasing vector of \( n \) indices into \( d \) specifying which of the new dimensions correspond to old dimensions:
  \( d(i) \) must equal \( \text{SIZE}(a, /\text{Dimensions}) \).

**Returned Value**

\( result \) — An array of dimensions \( d \), the expansion of the input array.

**Keywords**
None.

**Examples**

\[ pm, \text{EXPAND( [0,1], [2,3], [0] )} \]
pm, \( \text{EXPAND}( [0,1], [3,2], [1] ) \)

pm, \( \text{EXPAND}( [[0,1,2],[3,4,5]], [5,3,2], [1,2] ) \)

pm, \( \text{EXPAND}( [[0,1,2],[3,4,5]], [3,5,2], [0,2] ) \)

pm, \( \text{EXPAND}( [[0,1,2],[3,4,5]], [3,2,5], [0,1] ) \)

**See Also**

REBIN, REPLY
**EXPON Function**

Standard Library function that performs general exponentiation.

**Usage**

\[ \text{result} = \text{EXPON}(a, b) \]

**Input Parameters**

- **a** — An array (scalar) of any numerical data type.
- **b** — An array (scalar) of any numerical data type.

**Returned Value**

\[ \text{result} \] — A double complex array (scalar) containing the values \( a^b \).

**Keywords**

None.

**Examples**

\[ \text{pm, EXPON( [complex(0,1),-1], [complex(2,3),0.5] )} \]
**EXTREMA Function**

Standard Library function that finds the local extrema in an array.

**Usage**

\[ result = \text{EXTREMA}(\text{array}) \]

**Input Parameters**

*array* — The array for which the local extrema will be found.

**Returned Value**

*result* — A list containing two vectors of indices into *array*. *result*(0) contains the local minima and *result*(1) contains the local maxima.

**Keywords**

None.

**Examples**

\[ e = \text{EXTREMA}( [0,1,2,2,2,3,2,1,3] ) \text{ pm, e(0), ' ' & pm, e(1)} \]
\[ a = \text{bytscl( randomu(s,5,5), top=9 ) pm, a} \]
\[ e = \text{EXTREMA}( a ) \text{ & pm, e(0), ' ' & pm, e(1)} \]

**See Also**

MAX, MIN
**FACTOR Function**

Standard Library function that returns the prime factorization of an integer greater than 1.

**Usage**

\[ result = \text{FACTOR}(i) \]

**Input Parameters**

\( i \) — An integer greater than 1.

**Returned Value**

\( result \) — Sorted vector of longs containing the prime factorization of \( i \).

**Keywords**

\( a \) — If set, \( result \) contains all factors instead of just prime factors.

**Examples**

\[ \text{pm, FACTOR(12,}/a] \]
\[ \text{pm, FACTOR(12)} \]

**See Also**

GCD, LCM, PRIME


**GCD Function**

Standard Library function that returns the greatest common divisor of some integers greater than 0.

**Usage**

\[ \text{result} = \text{GCD}(i) \]

**Input Parameters**

\( i \) — An array of integers greater than 0.

**Returned Value**

\( \text{result} \) — An integer: the greatest common divisor of the integers \( i \).

**Keywords**

None.

**Examples**

\( \text{pm, GCD( [12,20,32] )} \)

**See Also**

FACTOR, LCM, PRIME
GREAT_INT Function

Greatest Integer Function. Standard Library function that returns the greatest integer less than or equal to the passed value. Also known as the Floor Function.

Usage

\[ \text{result} = \text{GREAT\_INT} (\text{values}) \]

Input Parameters

values — An array (scalar).

Returned Value

result — A long array (scalar) of the same dimensions as values: result(i) is the greatest integer less than or equal to values(i).

Keywords

None.

Examples

PM, GREAT\_INT ( [-0.5, 0, 0.5] )

See Also

SMALL\_INT
**GRIDN Function**

Standard Library function that grids n dimensional data.

**Usage**

\[ \text{result} = \text{GRIDN}(d, i) \]

**Input Parameters**

- \( d \) — An \((m,n+1)\) array of \(m\) datapoints in \(n\) independent variables and one dependent variable; \(d(*,n)\) is the dependent variable.
- \( i \) — A vector of \(n\) integers specifying the dimensions of the grid.

**Returned Value**

- \( \text{result} \) — An \(n\) dimensional array of values of the dependent variable on a regular grid over the independent variables.

**Keywords**

- \( r \) — A scalar specifying the order of the weighting function. The dependent variable at a grid point is computed as a weighted average of the variable over all neighborhood datapoints. The weighting function is \(1/e^r\) where \(e\) is the Euclidean distance between the grid point and the datapoint. \(r\) defaults to 2
- \( t \) — A scalar between 0 and 1 specifying neighborhood size. \(t=1\) gives a maximal neighborhood which includes all datapoints, while lower \(t\) values yield smaller neighborhoods. \(t\) defaults to 1
- \( b \) — A \(2 \times n\) array fixing the boundary of the grid. \(b(0,*)\) is the minimum corner and \(b(1,*)\) is the maximum corner. The default extent of the grid is the same as that of the data.
- \( f \) — The name of a user-supplied procedure describing voids in the independent variable space (datapoints and gridpoints within these regions are ignored in computation). Input to \(f\) is a \((p,n)\) array of \(p\) points in the independent variable space. \(f\) outputs two items where the first item is a vector of indices indicating which of the \(p\) input points are within bounds, and where the second item is a scalar that will appear as a place holder for the dependent variable at out-of-bounds gridpoints.
- \( c \) — (output) A list of \(n\) vectors defining the grid coordinates.
Examples

See wave/lib/user/examples/gridnex1.pro

wave/lib/user/examples/gridnex2.pro

See Also

FAST_GRID2, FAST_GRID3, FAST_GRID4, INTERPOLATE

---

**HISTN Function**

Standard Library function that computes an n dimensional histogram.

**Usage**

\[
\text{result} = \text{HISTN}(d[, \text{axes}])
\]

**Input Parameters**

\(d\) — An (m,n) array of m data points in n-space.

**Returned Value**

\(\text{result}\) — An n dimensional array of size \(\text{binnum}\).

\(\text{axes} = (\text{optional - use with }/\text{scale})\) an (n, \(\text{binnum}\)) array containing properly scaled axes with which to plot the results. For example:

CONTOUR, result, transpose(axes(0,*)), transpose(axes(1,*))

**Keywords**

**Binnum** — The number of bins for the histogram.

**Binsize** — The size of bins for the histogram. The default = 1.

---

**NOTE** Only 1 of \(\text{binnum}\) or \(\text{binsize}\) can be set.

**/Scale** — If set, the result is scaled so to have unit volume under the curve/surface when plotted against \(x.i/\text{stdev}(x.i)\)

**/Compatible** — If set, the result will align with HISTOGRAM. The default behavior of HISTN is “binnum-central” logic, while the default behavior of
HISTOGRAM is “binsize-central” logic. Setting /compatible will force HISTN to be “binsize-central.”

**NOTE** When using /compatible with 2D arrays and setting binsize manually, you may see poor results if the binsize is not appropriate for all variables. In this case, you should either set binnum or not use /compatible.

### Examples

Interpreting an n-dimensional histogram, a 2D example.

Consider two sets of 10 random numbers. If one computes 1D histograms with 3 bins, you may find these results (number of items in each bin):

```plaintext
x = RANDOMN(s,10) & y = RANDOMN(s,10)
xy = FLTARR(10,2) & xy(*,0) = x & xy(*,1) = y
PRINT, HISTN(x, binnum=3)
  4           5           1
PRINT, HISTN(y, binnum=3)
  3           4           4
```

The result for the 2D histogram may give:

```plaintext
PRINT, HISTN(xy, binnum=3)
  1           1           1
  2           2           0
  1           2           0
```

Summing this result vertically yields the 1D result for x. Summing this result horizontally yields the 1D result for y. To plot a probability distribution function with unit volume under the surface, one would call HISTN with the /Scaled keyword and plot the results against x/std(x) and y/STD(y). Using the axes output parameter with HISTN returns these vectors in a 2D array.

```plaintext
h2 = HISTN(xy, axes, binnum=3, /scaled)
shade_surf, h2, axes(0,*), axes(1,*)
```

### See Also

HISTOGRAM
INDEX_AND Function

Standard Library function that computes the logical AND for two vectors of positive integers.

Usage

\[ result = \text{INDEX}_\text{AND}(\text{array}_1, \text{array}_2) \]

Input Parameters

\( \text{array}_1 \) — A vector of positive integers.
\( \text{array}_2 \) — A vector of positive integers.

Returned Value

\( \text{result} \) — A vector containing all elements common to both input arrays. (\( \text{result} \) is not unique'd.)

Keywords

None.

Examples

\[ \text{PM INDEX_AND(} \{2, 0, 3\}, \{1, 2, 0, 2\} \text{) \]

See Also

INDEX_OR, WHEREIN
INDEX_OR Function

Standard Library function that computes the logical OR for two vectors of positive integers.

Usage

\[ \text{result} = \text{INDEX}_{-}\text{OR}(array_1, array_2) \]

Input Parameters

\( array_1 \) — A vector of positive integers.
\( array_2 \) — A vector of positive integers.

Returned Value

\( result \) — A vector consisting of all elements contained in either of the input arrays (\( result \) is unique’d).

Keywords

None.

Examples

PM, INDEX_OR( [2, 0, 3], [1, 2, 0, 2] )

See Also

INDEX_AND, WHEREIN
**INTERPOLATE Function**

Standard Library function that interpolates scattered data at scattered locations.

**Usage**

\[ \text{result} = \text{INTERPOLATE}(d, x) \]

**Input Parameters**

- \( d \) — An \((m, n+1)\) array of \(m\) datapoints in \(n\) independent variables and one dependent variable; \(d(*,n)\) is the dependent variable.
- \( x \) — A \((p,n)\) array specifying \(p\) interpolation points.

**Returned Value**

\( \text{result} \) — A 1d array of values of the dependent variable at points \(x\).

**Keywords**

- \( r \) — A scalar specifying the order of the weighting function. The dependent variable at an interpolation point is computed as a weighted average of the variable over all datapoints. The weighting function is \(1/e^r\) where \(e\) is the Euclidean distance between the interpolation point and the datapoint. \(r\) defaults to 2

**Examples**

\[ x = \text{findgen}(51) / 50 \quad \& \quad \text{plot}, x, \text{INTERPOLATE}([[0,1],[2,3]],x) \]

**See Also**

GRIDN
**INTRP Function**

Standard Library function that interpolates an array along one of its dimensions.

**Usage**

\[ \text{result} = \text{INTRP}(a, n, x) \]

**Input Parameters**

- \( a \) — An array.
- \( n \) — An integer (\( \geq 0 \)) designating the dimension to interpolate.
- \( x \) — A one-dimensional array giving the coordinates at which to interpolate.

**Returned Value**

\( \text{result} \) — The array of interpolated slices perpendicular to dimension \( n \).

**Keywords**

- \( z \) — A strictly increasing 1d array of coordinates for dimension \( n \) (defaults to the indices into this dimension).

**Examples**

See `wave/lib/user/examples/intrp_ex`.

**See Also**

REBIN, RESAMP
**JACOBIAN Function**

Standard Library function that computes the Jacobian of a function represented by \( n \) \( m \)-dimensional arrays

**Usage**

\[ j = \text{jacobian}(f) \]

**Input Parameters**

\( f \) — an \( n \)-element list of \( m \)-dimensional arrays all of the same dimensions \( d \); \( f \) represents a \( n \)-valued function of \( m \) variables.

**Returned Value**

\( j \) — an \( n \)-element list of \( m \)-element lists of \( m \)-dimensional arrays: \((j(p))(q)\) is the \( m \)-dimensional array (of dimensions \( d \)) which represents the derivative of the \( p^{th} \) dependent variable with respect to the \( q^{th} \) independent variable.

**Keywords**

\( x \) — \( m \)-element list of vectors defining the independent variables; by default, \( x(i) = \text{findgen}(d(i)) \).

**Example**

\[
\begin{align*}
  f &= \text{list}( \text{randomu}(s,10,20), \text{randomu}(s,10,20), \text{randomu}(s,10,20) ) \\
  j &= \text{jacobian}( f ) \\
  \text{for} \ p=0,2 \ \text{do for} \ q=0,1 \ \text{do pm, same}( (j(p))(q), \text{derivn}(f(p),q) )
\end{align*}
\]

**See Also**

CURVATURES, DERIVN, EUCLIDEAN, NORMALS
**LCM Function**

Standard Library function that returns the least common multiple of some integers greater than 1.

**Usage**

\[ result = \text{LCM}(i) \]

**Input Parameters**

\( i \) — An array of integers greater than 1.

**Returned Value**

\( result \) — An integer, the least common multiple of the integers \( i \).

**Keywords**

None.

**Examples**

\[ \text{pm, LCM([3,2,4])} \]

**See Also**

FACTOR, GCD, PRIME
**LISTARR Function**

Returns a list.

**Usage**

\[
\text{result} = \text{LISTARR}(\text{number\_elements},[\text{value}])
\]

**Input Parameters**

- **number\_elements** — The number of elements the created list should contain. Must be a scalar expression.
- **value** — (optional) The value with which to initialize each element of the list. May be any data type.

**Returned Value**

- **result** — A list with the requested number of elements.

**Keywords**

None.

**Discussion**

Values in the list are initialized to 0L unless **value** is specified.

**Examples**

INFO, LISTARR(2, 10), /Full
MINIMIZE Function

Standard Library function that minimizes a real valued function of n real variables.

Usage

\[ x = \text{MINIMIZE}(f, l, u, g, i, y) \]

Input Parameters

- **f** — A string specifying a user supplied function to be minimized. Input is a (m,n) array of m points in n-space (m variable); output is a (m,p+1) array \( b \), where \( p \) is the number of constraints, \( b({}^*,0) \) contains the objective function values at each of the m input points, and \( b({}^*,j) \) contains the corresponding values of the j'th constraint. All constraints must be of the form \( c(x) \leq 0 \).
- **l** — n-element vector of lower bounds for the independent variables
- **u** — n-element vector of upper bounds for the independent variables
- **g** — n-element vector giving an initial guess for the solution
- **i** — An integer limit on the number of iterations

Returned Value

- **x** — The n-element solution vector
- **y** — (optional) A (p+1)-element vector containing the objective function value at \( x \) followed by the constraint values at \( x \).

Keywords

- **d** — A string specifying a user-supplied gradient function. Input is the n-element vector at which to calculate the gradient(s). Output is a (n,p+1) array that contains the objective function gradient followed by the constraint gradients.
- **s** — An (n,2) array where \( s({}^*,0) \) is the maximum allowable step and \( s({}^*,1) \) is the minimum allowable step. The default is \[ [(u-l)/100], [(u-l)/100] \].

Examples

See wave/lib/user/examples/minimize_ex*.pro.
**MOLEC Function**

Standard Library function that creates an image of a ball and stick molecular model.

**Usage**

\[ \text{result} = \text{MOLEC}(\text{filename}) \]

**Input Parameters**

- **filename** — The name of an ASCII file describing the molecular model. Line 1 in the file consists of a single integer designating the number (m) of atoms. Each of the lines 2-(m+1) contains 7 floats describing an atom in terms of centroid, normalized RGB color components, and diameter. Line m+2 consists of a single integer equal to the number (n) of bonds; each of the lines (m+3)-(m+n+2) contains 6 floats describing a bond as endpoint1 followed by endpoint2.

**Returned Value**

- **result** — A 24-bit image of the molecular model.

**Keywords**

- \( h \) — A scale factor for adjusting atom size. The default is \( h=1.0 \).
- \( s \) — A 2-element vector specifying image size. The default is \( s=[500,500] \).
- \( v \) — A 3-by-4 double-precision floating-point array used to override the autogeneration of the view to that specified as: [viewpoint, top_left_viewplane, bottom_left_viewplane, bottom_right_viewplane]. \( v \) and \( !P.T \) control the 3d view.
- \( k \) — (output) A 3-by-4 double-precision floating-point array used to return the automatically calculated view as: [viewpoint, top_left_viewplane, bottom_left_viewplane, bottom_right_viewplane].

**Examples**

\[ \text{T3D, /Reset & TV, MOLEC(!data_dir+'molec.dat',h=0.6), true=3} \]
**MOMENT Function**

Standard Library function that computes moments of an array.

**Usage**

\[ result = \text{MOMENT}(a, i) \]

**Input Parameters**

- \( a \) — An array of \( n \) dimensions.
- \( i \) — A vector of \( n \) non-negative real numbers defining moment order.

**Returned Value**

\( result \) — A scalar double equal to:

\[ \sum_{J_{n-1}} \ldots \sum_{J_0} a(J_0 \ldots J_{n-1}) J_0^{i(0)} \ldots J_{n-1}^{i(n-1)} \]

**Keywords**

None.

**Examples**

\[ a = \text{BYTARR}(600, 500) \]
\[ x = \text{INDEX_CONV}(a, \text{LINDGEN(N_ELEMENTS(a)))} \]
\[ a(\text{WHERE}(x(*,0)-200)^2+(x(*,1)-300)^2 \text{LT 100^2})) = 255 \]
\[ \text{TV, a} \]
\[ \text{CENTROID} = \frac{\text{MOMENT}(a, [1,0]),\text{MOMENT}(a, [0,1])}{\text{MOMENT}(a, [0,0])} \]
\[ \text{PLOTS, CENTROID, /DEVICE, COLOR=0, PSYM=2} \]
**NEIGHBORS Function**

Standard Library function that finds the neighbors of specified array elements.

**Usage**

\[ result = \text{NEIGHBORS}(a, i) \]

**Input Parameters**

- **a** — An array of \( n \) dimensions.
- **i** — An \( m \)-element vector of \( m \) one-dimensional indices into \( a \).

**Returned Value**

- **result** — An \((m,*)\) array of one-dimensional indices into \( a \): \( \text{result}(j,*) \) contains \( i(j) \) and its neighbors.

**Keywords**

- **\( k \)** — A positive integer (less than or equal to \( n \)) defining connectivity. Two array cells are neighbors if they share a common boundary point, and if their centroids are within \( \sqrt{k} \) of each other. The default is \( k = 1 \), which implies neighbors share a common face.

**Examples**

\[
\begin{align*}
a &= \text{INDGEN}(8, 9) & \text{pm}, a \\
\text{pm}, \text{fix(NEIGHBORS}(a,[0,4,26,47,71])) \\
\text{pm}, \text{fix(NEIGHBORS}(a,[0,4,26,47,71],k=2))
\end{align*}
\]
**NORMALS Function**

Standard Library function that computes unit normals on a parametrically defined surface.

**Usage**

\[ n = \text{normals}(j) \]

**Input Parameters**

\( j \) — The Jacobian (computed by the JACOBIAN function) on the surface.

**Returned Value**

\( n \) — A 3-element list of 2-dimensional arrays of the same size as those in \( j \): \( n(i) \) is the array describing the distribution of the \( i^{th} \) component of the unit normal.

**Keywords**

None.

**Example**

See `wave/lib/user/examples/normals_ex.pro`.

**See Also**

CURVATURES, EUCLIDEAN, JACOBIAN
**PRIME Function**

Standard Library function that returns all positive primes less than or equal to a scalar input.

**Usage**

\[ result = \text{PRIME}(value) \]

**Input Parameters**

value — The scalar input.

**Returned Value**

result — A vector containing all positive primes less than or equal to value.

**Keywords**

None.

**Examples**

PRINT, PRIME(10)

**See Also**

FACTOR, GCD, LCM
PRODUCT Function

Returns the product of all elements in an array.

Usage

\[ \text{result} = \text{PRODUCT}(\text{array}) \]

Input Parameters

array — An array.

Returned Value

result — A scalar value equal to the product of all the elements in array.

Keywords

None.

Discussion

If array is of type single- or double-precision floating point, or single- or double-precision complex, the result will be of the same type. If array is of any other type, PRODUCT returns a single-precision floating-point result.

Examples

\[ \text{PM, PRODUCT( [2, 3, 4] )} \]
**RENDER24 Function**

Standard Library function that generates a ray-traced rendered 24-bit image of m objects.

**Usage**

\[result = \text{RENDER24}(b)\]

**Input Parameters**

\(b\) — A m-element list containing the m objects to render; objects are created using the CONE, CYLINDER, MESH, or SPHERE functions. The objects must be created with default material properties since these properties are controlled with keywords (see below).

**Returned Value**

\(result\) — (n,p,3) byte array containing a 24-bit image of the objects.

**Keywords**

\(c\) — (m,3) array of normalized RGB color components for the objects; by default, \(c(*,*) = 1.0\)

\(k\) — (m,3,3) array of normalized shade components for the objects: \(k(i,j,*)\) contains the ambient, reflective, and transmissive components for \(c(i,j)\). The sum of the three components must not exceed one. The default is \(k(i,j,*) = [0.0, 1.0, 0.0]\)

\(v\) — (3,4) array used to override the view automatically generated from \!p.t. If defined, \(v\) works like RENDER’s View keyword; if undefined, \(v\) works like RENDER’s Info keyword.

\(g\) — (4,q) array giving position and intensity for q light sources; the sum of the source intensities \(g(3,*)\) must equal one. The default is a single light source at the viewer’s eye.

\(s\) — A 2-element vector specifying image size. The default is \([256, 256]\)

**Examples**

\(b = \text{LIST}(\text{SPHERE}(), \text{CYLINDER}()) \& b(1).\text{transform}(3,2) = 2\)
REPLV Function

Standard Library function that replicates a vector into an array.

Usage

\[
result = \text{REPLV}(\text{vector}, \text{dim}_\text{vector}, \text{dim})
\]

Input Parameters

\text{vector} — The vector to be replicated.

\text{dim}_\text{vector} — A vector specifying the dimensions of the output array.

\text{dim} — An integer (\geq 0) designating the dimension to replicate.

Returned Value

\text{result} — An array of dimensions \text{dim}_\text{vector}.

Keywords

None.

Examples

\begin{verbatim}
PM, REPLV( [0,1], [2,4], 0 )
PM, REPLV( [0,1], [4,2], 1 )
PM, REPLV( [0,1], [4,2], 0 )
\end{verbatim}

See Also

EXPAND, REBIN, REPLICATE
**RESAMP Function**

Standard Library function that resamples an array to new dimensions.

**Usage**

\[
\text{result} = \text{RESAMP}(\text{array}, \text{dim}_1, \ldots, \text{dim}_n)
\]

**Input Parameters**

- **array** — An array of n dimensions.
- **dim\textsubscript{i}** — Integers (>0) specifying the new dimensions.

**Returned Value**

- **result** — The resampled version of \text{array}.

**Keywords**

- **Interp** — If set, n-linear interpolation is used instead of the default nearest-neighbor interpolation.

**Examples**

PM, RESAMP( [0,1,2,3], 3 )
PM, RESAMP( [0,1,2,3], 3, /i )
PM, RESAMP( [0,1,2,3], 6, /i )
PM, RESAMP( [[0,1,2,3],[4,5,6,7]], 6, 3, /i )

**See Also**

INTRP, REBIN
SAME Function

Standard Library function that tests if two variables are the same.

Usage

\[\text{result} = \text{SAME}(x, y)\]

Input Parameters

\[x\] — A variable.

\[y\] — A variable.

Returned Value

\[\text{result}\] — 1 if the two variables are the same (within keyword settings), 0 if not.

NOTE For Named Structures, the name is not tested, so \{a,x:1\} is same as \{b,x:1\}.

Keywords

\textit{NoType} — If set, the types of \(x\) and \(y\) are ignored. (FINDGEN(5) is same as INDGEN(5).)

\textit{NoDim} — If set, the dimensions of the arrays are ignored. ([1,2,3,4] is same as [[1,2],[3,4]] - same as SAME(a(*),b(*).)

\textit{NoVal} — If set, the values in the arrays are ignored. ([1,2,3] is same as [3,4,5].)

Examples

To test for exact match:

\[\text{result} = \text{SAME}(a, b)\]

To test for compatible sizes:

\[\text{result} = \text{SAME}(a, b, /\text{NoType}, /\text{NoVal})\]

(Replaces multiple SIZE calls.)

See Also

SIZE
SGN Function

Standard Library function that returns the sign of passed values.

Usage

\[ \text{result} = \text{SGN}(x) \]

Input Parameters

\( x \) — A scalar or array.

Returned Value

\( \text{result} \) — An integer array of the same size of \( x \) where each element is:

- 1 where \( x > 0 \)
- -1 where \( x < 0 \)
- 0 where \( x = 0 \)

Keywords

None.

Examples

PM, SGN([-0.5,0,0.5])
**SHIF Function**

Standard Library function that shifts an array along one of its dimensions.

**Usage**

\[ result = SHIF(array, dimension, shift_amount) \]

**Input Parameters**

- **array** — An array.
- **dimension** — An integer (\( \geq 0 \)) designating the shift dimension.
- **shift_amount** — An integer specifying the shift amount.

**Returned Value**

- **result** — An array of the same dimensions as **array**. **result** is obtained from the input array by a shift of **shift_amount** elements along dimension **dimension**. The shift is not cyclic; elements behind the shift are unaltered.

**Keywords**

- **y** — If set, the shift is cyclic.

**Examples**

```plaintext
PM, SHIF( [0,1,2,3,4,5], 0, 1 )
PM, SHIF( [0,1,2,3,4,5], 0, -2 )
PM, SHIF( [0,1,2,3,4,5], 0, -2, /y )
PM, SHIF( [[0,1,2],[3,4,5],[6,7,8]], 0, 1 )
PM, SHIF( [[0,1,2],[3,4,5],[6,7,8]], 1, 1 )
```

**See Also**

SHIFT
SLICE Function

Standard Library function that subsets an array along one of its dimensions.

Usage

\[ result = \text{SLICE}(\text{array}, \text{dimension}, \text{indices}) \]

Input Parameters

array — An array.

dimension — An integer (\( \geq 0 \)) designating the dimension to subset.

indices — A vector of indices into dimension dimension.

Returned Value

result — An array of slices perpendicular to dimension dimension.

Keywords

None.

Examples

\[
a = \text{INDGEN}(6, 5, 2) \quad \& \quad \text{pm, a, ['',''], \text{SLICE}(a, 0, [1,3,5])}
\]
**SMALL_INT Function**

Smallest Integer Function. Standard Library function that returns the smallest integer greater than or equal to the passed value. Also known as Ceiling Function.

**Usage**

\[ \text{result} = \text{SMALL\_INT}(x) \]

**Input Parameters**

\( x \) — An array (scalar).

**Returned Value**

\( \text{result} \) — A long array (scalar) of the same dimensions as \( x \): \( \text{result}(i) \) is the smallest integer greater than or equal to \( x(i) \).

**Keywords**

None.

**Examples**

PM, SMALL\_INT( [-0.5, 0, 0.5] )

\[x=\{-2.1, -1.9, 0, 1.9, 2.1\}\]

PRINT, SMALL\_INT(x)

; -2 -1 0 2 3

**See Also**

FIX, GREAT\_INT, NINT
**SORTN Function**

Standard Library function that sorts an array of n-tuples.

**Usage**

\[ result = \text{SORTN}(a) \]

**Input Parameters**

\( a \) — An (m,n) array of m n-tuples.

**Returned Value**

\( result \) — An array of indices, such that \( a(result,*) \) is the sorted version of \( a \).

**Keywords**

None.

**Examples**

\[ a = \text{BYTSCL}(\text{RANDOMU(seed,9,2), Top=5}) \]
\[ PM, a \]
\[ b = \text{SORTN}(a) \]
\[ PM, b \]
\[ PM, a(b,*) \]

**See Also**

SORT, UNIQUE, UNIQN
vtkADDATTRIBUTE Procedure

Collects point attributes for VTK datasets.

Usage

vtkADDATTRIBUTE, attributes

Input Parameters

attributes — A list variable containing all of the attributes for a dataset. Passing an undefined variable on the first call to creates the initial list. Using the variable on subsequent calls will add elements to the list. You should never have to create or modify the contents of this variable manually.

Keywords

Name — A scalar string specifying a name for this attribute. The default name is the attribute name in lower case.

Lookup_table_name — Only used with scalar attributes. A scalar string specifying the name of the lookup table to be associated with a scalar attribute. The default table is “default.”

One and only one of the following keywords can be used to add an attribute of the selected type:

Scalars — A vector of floating point numbers containing scalar values for each entry in points.

Lookup_table — An array of floating point numbers of size (3, m) containing normalized RGB values.

Vectors — An array of floating point numbers of size (3, n), where n equals the number of Points, containing the x, y, and z components for each vector.

Normals — An array of floating point numbers of size (3, n), where n equals the number of points, containing the x, y, and z components for each normal, where the x, y, and z values are normalized to a unit length of 1.

Color_scalars — An array of floating point numbers of size (m, n), where n equals the number of Points and m is the number of values per color scalar. Values are between 0.0 and 1.0.
**Texture_coordinates** — An array of floating point numbers of size \((m, n)\), where \(m\) is 1, 2, or 3 and \(n\) equals the number of points.

**Tensors** — An array of floating point numbers of size \((3, 3, n)\), where \(n\) equals the number of points.

**Discussion**

This procedure allows a set of attributes to be collected and passed to one of the dataset creation routines: `vtkPOLYDATA`, `vtkSTRUCTUREDPOINTS`, `vtkSTRUCTUREDGRID`, `vtkRECTILINIARGRID`, or `vtkUNSTRUCTUREDGRID`. Datasets can have one or more attributes associated with their points, and even more than one attribute of the same type, with the *Name* assigned to the attribute used to distinguish them. For *Scalars*, *Normals*, *Color_scalars*, *Texture_coordinates*, and *Tensors*, the number of supplied attributes must equal the number of points in the dataset to which they will be assigned.

---

**vtkAXES Procedure**

Creates a set of axes.

**Usage**

`vtkAXES`

**Input Parameters**

None.

**Keywords**

*Charsize* — A floating point scalar or three-element array, the size of the text for tickmark labels and axes labels. (Default: \(0.4\times \text{Lengths}\))

*Format* — A FORTRAN style format string to use for the tick mark labels. (Default: ‘(g10.2)’)

*Name* — Specify a name to be used to create this object. If an undefined variable is used or no name specified, then a random name is used. This name can be used in calls to `vtkCOMMAND` to modify this object.
**Position** — An array of three floating point numbers in data coordinates describing the origin for the axis. (Default: [0, 0, 0])

**Lengths** — An array of three floating point numbers describing the length of the x, y, and z axes, respectively, specified in data coordinates. (Default: [1, 1, 1])

**Labels** — If non-zero, any tick marks drawn are labeled with default values.

**LOD** — If nonzero, the tickmarks are created as level-of-detail actors to aid in keeping a high frame-rate during frequent render requests due to user mouse interaction. If set to a value greater than 1, the number of points to use in the random cloud.

**Sigfig** — An integer, the number of significant figures to use for the tick mark labels. (Default: 2).

**TextColor** — The color to use for text used for [XYZ]Title. See vtkWINDOW (page 85) for possible ways to specify the color. (Default: 'white')

**Tickscale** — A float, a scaling value passed to vtkSCATTER for the tick mark glyphs. (Default: 0.33)

**Ticksymbol** — An integer, passed to vtkSCATTER to set the glyph to use for the tick marks. (Default: 0, a sphere)

Other keywords are listed below. For a description of each keyword, see Chapter 3, Graphics and Plotting Keywords in the **PV-WAVE Reference**.

*[XYZ]Tickname*  *[XYZ]Ticks*  *[XYZ]Tickv*  *[XYZ]Title*

**Discussion**

This procedure creates three axes displayed as lines in the x, y, and z direction, with an optional label at the top of each axis.

**Example**

```vtk
vtkAxes, lengths = [1,1.5,2]
```

**See Also**

AXIS
**vtkCAMERA Procedure**

Changes the camera’s parameters.

**Usage**

vtkCAMERA

**Input Parameters**

None.

**Keywords**

*Name* — Specify a name to be used to create this object. If an undefined variable is used or no name is specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.

*Position* — An array of three floating point numbers describing the x, y, and z position for the camera in data coordinates.

*FocalPoint* — An array of three floating point numbers describing the x, y, and z position for the camera’s focal point in data coordinates.

*ClippingRange* — An array of two floating point numbers describing the distance from the camera to the front and back clipping planes (in data coordinates).

*ViewUp* — An array of three floating point numbers describing a vector that represents the up direction for the view.

*Distance* — A floating point number describing the distance from the focal point to the camera (which will modify the FocalPoint value) in data coordinates.

*ViewAngle* — A floating point number that sets the view angle of the camera in degrees.

*Azimuth* — A scalar value describing the angle in degrees to rotate the camera about the view up vector centered at the focal point. This moves the camera from side to side.

*Elevation* — A scalar value describing the angle in degrees to rotate the camera about the cross product of the direction of projection and the view up vector centered on the focal point. This moves the camera up and down.
Roll — A scalar value describing the angle in degrees to rotate the camera about the direction of projection. This rolls the camera about the direction of projection.

**Discussion**

A default camera is created for each vtkWINDOW with these properties: position and focal point such that all objects are visible, with the camera centered on the entire scene; view up along the Y axis; view angle set to 30 degrees; and a clipping range set to 0.1, 1000.0.

**NOTE**  vtkSURFACE, vtkSCATTER and vtkPOLYSHADE change this default and set the up vector to be along the z axis.

**Example**

```python
pyramid_list = [[0, 0, 0], [0, 1, 0], [1, 0, 0], [1, 1, 0], [.5, .5, 1]]
vertex_list = [3, 0, 2, 4, 3, 2, 3, 4, 3, 1, 4, 3, 1, 0, 4, 4, 0, 1, 3, 2]  
vtkPolyshade, pyramid_list, vertex_list  
vtkCamera, Azimuth=25, Elevation=45, ViewAngle=120
```

**vtkCLOSE Procedure**

Closes the VTK process.

**Usage**

vtkCLOSE

**Input Parameters**

None.

**Keywords**

None.
Discussion

This procedure closes all VTK windows and shuts down the Tcl/Tk-spawned process. It should be called before exiting PV-WAVE.

Example

A standard close call.

`vtkCLOSE`

See Also

`vtkINIT`, `vtkWINDOW`, `vtkERASE`, `vtkWDELETE`

---

**vtkCOLORBAR Procedure**

Adds a color bar legend to a VTK scene using the current PV-WAVE color table.

**Usage**

`vtkCOLORBAR`

**Keywords**

- **Vertical** — If set, the color bar is aligned vertically. Default alignment is horizontal.
- **Title** — The title of the legend. (Default: none)
- **Position** — A three-element array, the position of the lower left corner of the color bar. (Default: [0,0,0])
- **NumLabels** — The number of labels to draw. (Default: 5)
- **Width** — The width of the legend in device coordinates. (Default: 0.8, or 0.15 with `/Vertical`)
- **Height** — The height of the legend in device coordinates. (Default: 0.15, or 0.9 with `/Vertical`)
- **CRange** — A two-element vector, the range of colors (Default: [0,255])
- **LRange** — A two-element vector, the label range (Default: `CRange`)
**Sigfig** — An integer, the number of significant figures to use for the labels. (Default: 3).

**Name** — Specifies a name to be used to create this object. If an undefined variable is used or no name is specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.

**NoShadow** — If set, labels are drawn without shadows.

---

**vtkCOMMAND Procedure**

Sends Tcl and VTK commands to the Tcl process.

**Usage**

vtkCOMMAND, *command*

**Input Parameters**

*command* — A string representing the VTK command to invoke.

**Keywords**

**Result** — A string or string array containing any results from the execution of the command in the Tcl shell.

**Discussion**

The Basic interface to send raw Tcl or VTK commands to the spawned Tcl process.

**Example**

Use vtkCOMMAND to set the background blue.

```tcl
vtkwindow, 1
vtkcommand, ’renderer1 SetBackground 0.0 0.0 1.0’
vtkrenderwindow
```

**See Also**

vtkRENDERWINDOW
**vtkERASE Procedure**

Erases the contents of the current VTK window.

**Usage**

vtkERASE [, background_color]

**Input Parameters**

`background_color` — (optional) The background color to be used for the window, specified as a 24-bit color. See vtkWINDOW (page 85) for possible ways to specify the color.

**Keywords**

None.

**Discussion**

This procedure works like ERASE for PV-WAVE windows. It removes all actors, cameras, and lights from the current window.

**Example**

This example shows vtkERASE removing the axes from the window.

vtkwindow, 1
vtkaxes
vtkerase

**See Also**

ERASE, VtkCLOSE, vtkWINDOW
**vtkGRID Procedure**

Adds 3D grid lines to a VTK scene.

**Usage**

vtkGRID [, Number=n]

**Keywords**

*Number* — A scalar or a three element array, the number of segments with grid lines in each (x, y, z) direction. (Default: [1,1,1], a box)

*Lengths* — A scalar or a three-element array, the extent of the grid. (Default: [1,1,1])

*Position* — A three-element array, the position of the origin of the grid. (Default: [0,0,0])

*Color* — The color to use for the polylines (passed to vtkPLOTS). See vtkWINDOW for possible ways to specify the color. (Default: ‘white’)

*Thick* — A float, the thickness of the grid lines (passed to vtkPLOTS). (Default: 1.0)

*Name* — A string, the name to be used to create this object. If an undefined variable is used or no name is specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.

*UseAxes* — If nonzero, the most recently created vtkAXES scale is used to define the Lengths array, which then does not need to be defined explicitly.

*LOD* — If nonzero, use level-of-detail actors for the grid lines (passed to vtkPLOTS).

**Example**

vtkSURFACE, DIST(20)

vtkGRID, /useAxes, Thick=6

vtkGRID, Number=[2,2,6], Color=’red’, Thick=4, /useAxes
**vtkHEDGEHOG Procedure**

Creates a HedgeHog (vector) plot.

**Usage**

vtkHEDGEHOG, points, vectors, scalars

**Input Parameters**

points — A 3, n array of points (location of the lines).

deprecated vectors — A 3, n array of vectors (orientation and length of the lines).

scalars — (optional) An n-element array of scalars (colors of the lines).

**Keywords**

Scalefactor — A float, a scaling factor to control the size of the oriented lines of the HedgeHog object (Default: 1.0).

SRRange — A two-element integer array, the scalar range (Default: [0,255]).

LOD — If nonzero, a level-of-detail actor is created to aid in keeping a high frame-rate during frequent render requests due to user mouse interaction. If set to a value greater than 1, the number of points to use in the random cloud.

NoRotate — Does not perform any camera rotations. Used when a previous call to vtkSURFACE, vtkSCATTER or vtkPOLYSHADE has already set the camera angle.

NoAxes — If set, no axes are created.

NoErase — If nonzero, prevents the window from being erased to the background color before drawing the new scene. If not set, then all lights, cameras, and objects are removed from the scene before objects for the new scene are added.

These keywords are passed to vtkAXES:

Charsize TextColor [XYZ]Title [XYZ]Ticks


Labels Sigfig Format

See vtkAXES for descriptions.
Example

This example plots a Hanning surface and its normals.

```plaintext
LOADCT, 2
n = 50
x = REBIN(INDEXGEN(n),n,n)
y = TRANSPOSE(x)
z = (n/2)*(HANNING(n,n))
norm = NORMALS(JACOBIAN(LIST(x,y,z)));

points = FLTARR(3,n*n,/NoZero)
points(0,*) = x(*)
points(1,*) = y(*)
points(2,*) = z(*)
vectors = FLTARR(3,n*n,/NoZero)
vectors(0,*) = (norm(0))(*)*SQRT(z(*))
vectors(1,*) = (norm(1))(*)*SQRT(z(*))
vectors(2,*) = (norm(2))(*)*SQRT(z(*))

scalars = z(*);

vtkHedgeHog, points, vectors, scalars, /NoAxes, scalef=0.75
vtkSurface, REFORM(points(2,*),n,n), $
    Shades=REBIN([0.8,0.8,0.8,.75],4,n,n), $
    /NoErase, /NoRotate, /NoAxes
```

See Also

vtkAXES
vtkINIT Procedure

Initializes the VTK system.

Usage

tvtkINIT

Input Parameters

None.

Keywords

File — If set, a temporary file is used to communicate data sets to VTK instead of a socket connection. For very large data sets with many floating-point values, this method is considerably faster; however, read/write permissions are required. If set to one (/File), any data set greater than 1024 bytes is written to file. If set to a value (File=fbytes), data sets larger than fbytes are written to file; smaller data sets are sent via sockets. Setting /File is equivalent to setting File=1024. This keyword affects only data: commands are always sent by the socket connection.

Noshell — If set, the keyword is passed along to the SPAWN procedure that initiates the VTK Tcl process. This keyword is required when calling VTK routines from a JWA VE wrapper and should not be used otherwise.

Path — Used in conjunction with the File keyword, a string indicating the file path to the directory where the temporary file(s) are to be created.

Print — If present and nonzero, causes the output from the spawned Tcl/Tk shell to be sent back to PV-WAVE and displayed in the console. This keyword is useful for debugging low-level VTK calls.

Timeout — A floating point scalar specifying a time interval in seconds which vtkINIT will wait before giving up on establishing a socket connection to the spawned Tcl shell. (Default: 20)

Discussion

This procedure must be performed before any other VTK commands. It causes a Tcl/Tk shell to be spawned and sets up communication with it. It also initialized various internal VTK parameters. The following routines will automatically call
vtkINIT if it has not already been called: vtkWINDOW, vtkPOLYSHADE, vtkSURFACE, and vtkSCATTER.

Example
This example uses vtkINIT to initialize VTK with a timeout of 10 seconds.

```
vtkINIT, timeout=10
```

See Also
VtkCLOSE

---

**vtkLIGHT Procedure**

Adds a light to a VTK window.

**Usage**

```
vtkLIGHT
```

**Input Parameters**

None.

**Keywords**

- **Name** — Specifies a name to be used to create this object. If an undefined variable is used or no name specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.
- **Color** — The color for the light. See vtkWINDOW (page 85) for possible ways to specify the color. (Default: ‘white’)
- **Position** — An array of three floating point numbers describing the x, y, and z position for the light. The default behavior is to have the light follow the camera position.
- **FocalPoint** — An array of three floating point numbers describing the x, y, and z position for the light’s focal point.
- **Intensity** — A float value between 0.0 and 1.0 specifying the intensity of the light.
**DirectionAngle** — An array of two floating point numbers that set the position and focal point of a light based on elevation and azimuth. The light is moved to shine from the given angle. Angles are given in degrees.

**Discussion**

A white light, which follows the camera position, is created by default for a VTK Window.

**NOTE** Other light parameters not supported in this wrapper can be set using the assigned *Name* and *vtkCOMMAND*.

**Example**

```python
pyramid_list = [[0,0,0],[0,1,0],[1,0,0],[1,1,0],[.5,.5,1]]
vertex_list=[3,0,2,4,3,2,3,4,3,1,4,3,1,0,4,4,0,1,3,2]
vtkPolyshade, pyramid_list, vertex_list
  vtkLight, Color='0000FF'XL, Position=[10,10,10]
  vtkLight, Color='00FF00'XL, Position=[10,10,-10]
  vtkLight, Color='FF0000'XL, Position=[-10,-10,-10]
```

**vtkPLOTS Procedure**

Adds a polyline.

**Usage**

`vtkPLOTS, points`

**Input Parameters**

*points* — An array of floating point numbers of size (3, n) where n is the number of points. `points(0, *)` is taken as an x value, `points(1, *)` is taken as a y value, and `points(2, *)` is taken as a z value.
Keywords

**Name** — Specifies a name to be used to create this object. If an undefined variable is used or no name specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.

**Color** — The color to use for the polyline. See vtkWINDOW (page 85) for possible ways to specify the color. (Default: ‘white’)

**Thick** — A float describing the thickness of the polyline. The default is 1.0, which is scaled to correspond to a radius of 0.001 in data coordinates.

**LOD** — If nonzero, a level-of-detail actor is created to aid in keeping a high frame-rate during frequent render requests due to user mouse interaction. If set to a value greater than 1, the number of points to use in the random cloud.

**Nolines** — If nonzero, causes a cloud of points to be displayed rather than a polyline.

Discussion

This procedure is similar to the PLOTS procedure for PV-WAVE windows. Polylines are drawn as connected cylinders.

Example

```plaintext
z = findgen(1000)/999
x = z*sin(50*z)
y = z*cos(50*z)
vtkplots, transpose([[x],[y],[z]]), thick=5, color='blue'
vtkwindow,2
vtkplots, transpose([[x],[y],[z]]), color='red',/nolines
```

See Also

PLOTS, vtkSCATTER
vtkPOLYDATA Procedure

Passes vertex/polygon lists, lines, points, and triangles to VTK.

Usage

vtkPOLYDATA, points

Input Parameters

points — A two-dimensional array of floating point numbers of size (3, n) describing x, y, and z points.

Keywords

Restore — An associative array containing all of the data and attributes for a polydata dataset, usually created using the Save keyword. If this parameter is passed, then only the keywords Name and Filename can be used.

Name — Specifies a name to be used to create this data source. This name can be used in calls to vtkCOMMAND.

Filename — File to store data using standard VTK ASCII format.

Save — Returns the data in the specified variable stored in an associative array. Data is not sent to VTK if this parameter is specified.

Polygons — A vector of integers describing polygons, organized as vertex count followed by indices into points, repeated for all polygons. This is the same format as a polygon list used in POLYSHADE.

Vertices — A vector of integers describing vertices, organized as vertex count followed by indices into points, repeated for all vertices.

Lines — A vector of integers describing polylines, organized as vertex count followed by indices into points, repeated for all polylines.

Triangle_Strips — A vector of integers describing triangle strips, organized as vertex count followed by indices into points, repeated for all triangle strips.

Attributes — A list created using vtkADDATTRIBUTE containing one or more attributes associated with the points in the dataset.
**Discussion**

Contains points and polygons (like the polygon vertex list in PV-WAVE) as well as vertices, lines, and triangle strips. See the VTK documentation, which can be downloaded from [http://public.kitware.com](http://public.kitware.com), for more details on the data and attributes for the PolyData dataset format.

---

**vtkPOLYSHADE Procedure**

Renders a polygon object.

**Usage**

`vtkPOLYSHADE, vertices, polygons`

**Input Parameters**

*Vertices* — A \((3, n)\) array containing the \(x\), \(y\), and \(z\)-coordinates of each vertex in data coordinates.

*Polygons* — An integer or longword array containing the indices of the vertices of each polygon. The vertices of each polygon should be listed either clockwise or counterclockwise order when observed from outside the surface. The vertex description of each polygon is a vector of the form \([n, i_0, i_1, \ldots, i_{n-1}]\), and the array polygons is the concatenation of the lists of each polygon.

**Keywords**

*Name* — Specify a name to be used to create this object. If an undefined variable is used or no name specified then a random name is used. This name can be used in calls to `vtkCOMMAND` to modify this object.

*Color* — An array expression, of the same dimensions as the number of vertices passes (the value “\(n\)” above), containing the color index at each vertex. Alternately an expression describing one color to be used for the entire polygonal surface or wireframe. If this keyword is omitted, a white surface is displayed.

To specify a single color, see `vtkWINDOW` (page 85) for possible ways to specify the color. If a two-dimensional array of colors is specified (for an image overlay) the shades variable can be in any of these formats:
Wireframe — When present and nonzero, a wire-frame mesh is drawn rather than a shaded surface.

LOD — If nonzero, a level-of-detail actor is created to aid in keeping a high frame-rate during frequent render requests due to user mouse interaction. If set to a value greater than 1, the number of points to use in the random cloud.

NoAxes — If set, no axes are created.

NoRotate — Does not perform any camera rotations. Used when a previous call to vtkSURFACE, vtkSCATTER or vtkPOLYSHADE has already set the camera angle.

NoErase — If nonzero prevents the window from being erased to the background color before drawing the new scene. If not set then all lights, cameras, and objects are removed from the scene before objects for the new scene are added.

These keywords are passed to vtkAXES:

- FIX(n) A one-dimensional array of short integers or bytes specifying an index into the current PV-WAVE color table for each point. The RGB color for each point is obtained from the corresponding entry in the current PV-WAVE color table.
- LONG(n) A one-dimensional array of long integers specifying the 24-bit color at each point.
- FLOAT(3, n) A floating point array of size (3, n) containing the normalized values specifying the red, green, and blue components of the color at each point.
- FLOAT(4, n) A floating point array of size (4, n) containing the normalized values specifying the red, green, blue, and alpha components of the color at each vertex. The alpha component is the transparency where 0.0 is completely transparent and 1.0 is opaque.

Other keywords are listed below. For a description of each keyword, see Chapter 3, Graphics and Plotting Keywords in the PV-WAVE Reference.

Ax Az
Discussion

This procedure is similar to the POLYSHADE procedure for PV-WAVE windows. Wireframes can be produced as well as surfaces shaded in one color or overlaid with an image. Transparency is also supported.

Example

```python
pyramid_list = [[0, 0, 0], [0, 1, 0], [1, 0, 0], [1, 1, 0], [.5, .5, 1]]
vertex_list=[3, 0, 2, 4, 3, 2, 3, 4, 3, 3, 1, 4, 3, 1, 0, 4, 4, 0, 1, 3, 2]
vtkPolyshade, pyramid_list, vertex_list, color='blue'
```

See Also

AXIS, POLYSHADE

---

**vtkPPMREAD Function**

Reads a PPM file.

**Usage**

```python
image = vtkPPMREAD (filename)
```

**Input Parameters**

- `filename` — File path of PPM file.

**Returned Value**

- `image` — An array (3, width, height) of bytes containing the 24-bit image.

**Keywords**

None.
Discussion

This function is used to read the rudimentary PPM binary files created by VTK and containing images stored as RGB values. The images can be displayed with \texttt{TV}, image, True=1 or converted to an 8-bit image using \texttt{ipcolor\_24\_8}.

Example

Example of reading the file \texttt{wave.ppm} and storing an image.

\begin{verbatim}
Image=vtkppmread('wave.ppm')
\end{verbatim}

See Also

\texttt{vtkPPMWRITE}, \texttt{vtkTVRD}

\underline{vtkPPMWRITE Procedure}

Writes the contents of a VTK window to a PPM file.

Usage

\begin{verbatim}
vtkPPMWRITE [, window_index]
\end{verbatim}

Input Parameters

\textit{window_index} — (optional) An integer specifying the index of an existing VTK window. If omitted, the current window is used.

Keywords

\textit{Filename} — File path to store PPM file. (Default: \texttt{‘wave.ppm’})

Discussion

This procedure saves a snapshot of the selected window as a PPM file. It is important to make sure the VTK window fully visible when this routine is called because an obscuring portion of another window will be captured as part of the image. This is a limitation of VTK.
Example 1
Writing a PPM file.
vtkwindow, 1
vtkaxes
vtkPPMWRITE, 1

Example 2
Writing the PPM file to the file name of PV.ppm.
vtkwindow, 2, background='blue'
vtkPPMWRITE, 2, filename='PV.ppm'

See Also
vtkPPMREAD, vtkTVRD

**vtkRECTILINEARGRID Procedure**

Passes data describing a rectilinear grid to VTK.

**Usage**
vtkRECTILINEARGRID, *Dimensions*

**Input Parameters**

*Dimensions* — A 3-element vector of integers describing dimensions in x, y, and z. Use 1 for the third dimension if only a two-dimensional array is described.

**Keywords**

*Restore* — An associative array containing all of the data and attributes for a poly-data dataset, usually created using the *Save* keyword. If this parameter is passed, then only the keywords *Name* and *Filename* can be used.

*Name* — Specifies a name to be used to create this data source. This name can be used in calls to vtkCOMMAND.

*Filename* — File to store data using standard VTK ASCII format.
**Save** — Returns the data in the specified variable stored in an associative array. Data is *not* sent to VTK if this parameter is specified.

**X_coordinates** — A vector of floating point numbers of the same length as the first dimension in *Dimensions* describing monotonically increasing coordinate values. Increasing integers starting with 0 are used as a default.

**Y_coordinates** — A vector of floating point numbers of the same length as the second dimension in *Dimensions* describing monotonically increasing coordinate values. Increasing integers starting with 0 are used as a default.

**Z_coordinates** — A vector of floating point numbers of the same length as the third dimension in *Dimensions* describing monotonically increasing coordinate values. Increasing integers starting with 0 are used as a default.

**Attributes** — A list created using *vtkADDATTRIBUTE* containing one or more attributes associated with the points in the dataset.

**Discussion**

This procedure creates a dataset with a regular topology and semiregular geometry aligned along the x, y, and z axes. See the VTK documentation, which can be downloaded from [http://public.kitware.com](http://public.kitware.com), for more details on the data and attributes for the RectilinearGrid dataset format.

---

**vtkRENDERWINDOW Procedure**

Renders a VTK window.

**Usage**

`vtkRENDERWINDOW [, window_index]`

**Input Parameters**

*window_index* — (optional) An integer specifying the index of an existing VTK window. If omitted, the current window is used.

**Keywords**

None.
Discussion

Call this procedure after all objects (lights, cameras, surfaces, polygon meshes, etc.) have been added to the window. This routine starts the rendering process and creates the initial rendered scene. You need to call this procedure only if you used the *Norender* keyword with vtkWINDOW, or if you are making low-level calls to VTK using vtkCOMMAND.

Example 1

This example shows the essence of vtkRENDERWINDOW.

```plaintext
vtkwindow, 1, /norender
vtkaxes
vtkrenderwindow
```

Example 2

A more complicated example:

```plaintext
vtkwindow, 2, /norender
V=[[0,0,0],[1,0,0],[1,1,0],[0,1,0]]
p=[4,0,1,2,3]
vtkpolyshade, v, p
vtkaxes
vtktext, 'This is VTK', charsize=[1.0,1.0,1.0], color='blue'
vtkrenderwindow, 2
```

See Also

vtkWINDOW, vtkCOMMAND
vtkSCATTER Procedure

Renders 3D points.

Usage

vtkSCATTER, points

Input Parameters

points — A float array of size (3, n) describing x, y, and z points in data coordinates.

Keywords

Name — Specifies a name to be used to create this object. If an undefined variable is used or no name specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.

Symbol — A scalar integer describing the type of marker or glyph to be displayed for each point. Supported values are 0-4 where:

0 = Sphere
1 = Cube
2 = Cone
3 = Cylinder
4 = Earth

Color — An array expression of the same dimensions as the number of points (the value n above), containing the color index at each point. Alternately an expression describing one color to be used for all points. If this keyword is omitted, white points are displayed.

To specify a single color, see vtkWINDOW (page 85). If a vector of colors is specified, the color variable can be in any of these formats:

FIX(n) A one-dimensional array of short integers or bytes specifying an index into the current PV-WAVE color table for each point. The RGB color for each point is obtained from the corresponding entry in the current PV-WAVE color table.
LONG(n) A one-dimensional array of long integers specifying the 24-bit color at each point.

FLOAT(3, n) A floating point array of size (3, n) containing the normalized values specifying the red, green, and blue components of the color at each point.

Scale — A float value specifying the scaling factor for the size of each glyph. (Default: 1.0)

LOD — If nonzero, the glyphs are created as level-of-detail actors to aid in keeping a high frame-rate during frequent render requests due to user mouse interaction. If set to a value greater than 1, the number of points to use in the random cloud.

NoRotate — Does not perform any camera rotations. Used when a previous call to vtkSURFACE, vtkSCATTER or vtkPOLYSHADE has already set the camera angle.

NoErase — If nonzero, prevents the window from being erased to the background color before drawing the new scene. If not set, then all lights, cameras, and objects are removed from the scene before objects for the new scene are added.

NoAxes — If present and non zero, then no x, y, and z axes will be drawn.

TextColor — The color for text used for the axes titles. See vtkWINDOW (page 85) for possible ways to specify the color. (Default: ‘white’)

These keywords are passed to vtkAXES:

TickSymbol Labels Sigfig Format

Other keywords are listed below. For a description of each keyword, see Chapter 3, Graphics and Plotting Keywords in the PV-WAVE Reference.

Ax Az Charsize

[XYZ]Title [XYZ]Range

Discussion
This procedure plots points in space using three-dimensional markers (glyphs) with optional axes.
Example

data=FLTARR(3,200)
    ; Create the data array.
s=DC_READ_FREE(!Data_Dir+'scattered.dat',data,/Column,/Resize)
    ; Read in the data
vtkSCATTER,data
    ; A quick look at the data.
TEK_COLOR
    ; Set up a color table.
s=SIZE(data) & c=INDGEN(s(2))
    ; Create an array of color values for the points.
vtkSCATTER,data,Color=c,Xtitle='X Values',Ytitle='Y Values',Ztitle='Z Values',TextColor=16,ax=10,az=100,Scale=.6
    ; A better look at the data.

See Also

AXIS  vtkPLOTS

vtkSLICEVOL Procedure

Creates a sliced 3D volume at specific x, y, z locations.

Usage

vtkSLICEVOL, v, [sx=sx, sy=sy, sz=sz, xc=xc, yc=yc, zc=zc]

Input Parameters

v — A 3D array, the volume to slice.

Keywords

sx — A 1D array, the x coordinate(s) at which to slice the volume.
sy — A 1D array, the y coordinate(s) at which to slice the volume.
sz — A 1D array, the z coordinate(s) at which to slice the volume.
$xc$ — A 1D array with the same number of elements as the first dimension of $v$, the $x$ coordinates of the volume.

$yc$ — A 1D array with the same number of elements as the second dimension of $v$, the $y$ coordinates of the volume.

$zc$ — A 1D array with the same number of elements as the third dimension of $v$, the $z$ coordinates of the volume.

$Interp$ — If set, the shading is interpolated (passed to RESAMP).

$Dim$ — An integer, the number of vertices on each side of each plane. (Default: 25)

$Name$ — Specifies a name to be used to create this object. If an undefined variable is used or no name specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.

$Wireframe$ — When present and nonzero, a wire-frame mesh is drawn rather than a shaded surface.

$LOD$ — If nonzero, a level-of-detail actor is created to aid in keeping a high frame-rate during frequent render requests due to user mouse interaction. If set to a value greater than 1, the number of points to use in the random cloud.

$NoRotate$ — Does not perform any camera rotations. Used when a previous call to vtkSURFACE, vtkSCATTER or vtkPOLYSHADE has already set the camera angle.

$NoErase$ — If nonzero, prevents the window from being erased to the background color before drawing the new scene. If not set, then all lights, cameras, and objects are removed from the scene before objects for the new scene are added.

Other keywords are listed below. For a description of each keyword, see Chapter 3, Graphics and Plotting Keywords in the PV-WAVE Reference.

Ax

Az

Discussion

If no slices are requested through the $sx$, $sy$, and $sz$ keywords, the volume is sliced at the midpoints of each index. If $xc$, $yc$, or $zc$ are not provided, indices into $v$ are used.

Example

\[
x = \text{genvect}(-5, 5, .25) \\
y = \text{genvect}(-4, 4, .2)
\]
z = genvec(-3,3,.15)
v = sqrt(TENSOR_ADD(TENSOR_ADD(x^2,0.3*y^2),1.5*z^2))
vtkSliceVol, v, sx=[-4,.4,2.6], sy=-.15, sz=[-3,1], xc=x, yc=y, zc=z, dim=15

See Also
SLICE

vtkSTRUCTUREDGRID Procedure
Passes data describing a structured grid to VTK.

Usage
vtkSTRUCTUREDGRID, dimensions, points

Input Parameters
dimensions — A 3-element vector of integers describing dimensions in x, y, and z. Use “1” for the third dimension if only a two-dimensional array is described.
points — A two-dimensional array of floating point numbers of size (3, n) describing x, y, and z points.

Keywords
Restore — An associative array containing all of the data and attributes for a polydata dataset, usually created using the Save keyword. If this parameter is passed, then only the keywords Name and Filename can be used.
Name — Specifies a name to be used to create this data source. This name can be used in calls to vtkCOMMAND.
Filename — File to store data using standard VTK ASCII format.
Save — Returns the data in the specified variable stored in an associative array. Data is not sent to VTK if this parameter is specified.
Attributes — A list created using vtkADDATTRIBUTE containing one or more attributes associated with the points in the dataset.
Discussion
1, 2, or 3D point data on a topological grid, where the actual points are specified as $x$, $y$, and $z$ values in Cartesian coordinates. See the VTK documentation, which can be downloaded from http://public.kitware.com, for more details on the data and attributes for the StructuredGrid dataset format.

vtkSTRUCTUREDPOINTS Procedure
Passes data describing structured points to VTK.

Usage
vtkSTRUCTUREDPOINTS, dimensions

Input Parameters
dimensions — A 3-element vector of integers describing dimensions in $x$, $y$, and $z$. Use 1 for the third dimension if only a two-dimensional array is described.

Keywords
Restore — An associative array containing all of the data and attributes for a poly-data dataset, usually created using the Save keyword. If this parameter is passed then only the keywords Name and Filename can be used.

Name — Specifies a name to be used to create this data source. This name can be used in calls to vtkCOMMAND.

Filename — File to store data using standard VTK ASCII format.

Save — Returns the data in the specified variable stored in an associative array. Data is NOT sent to VTK if this parameter is specified.

Origin — A 3-element vector of floating point numbers containing the $x$, $y$, and $z$ origin point for the data.

Spacing — A 3-element vector of floating point numbers containing the spacing (width, height, length) of the cubical cells that compose the data set.

Attributes — A list created using vtkADDATTRIBUTE containing one or more attributes associated with the points in the dataset.
Discussion
Definition of a 1, 2, or 3D arrays (describing lines, grids and voxels), their origin, and spacing. See the VTK documentation, which can be downloaded from http://public.kitware.com, for more details on the data and attributes for the Structured-Points dataset format.

vtkSURFACE Procedure
Renders a surface.

Usage
vtkSURFACE, z [,x] [,y]

Input Parameters
z — A two-dimensional array containing the values that describe the surface. If x and y are supplied, the surface is plotted as a function of the x and y locations specified by their contents. Otherwise, the surface is generated as a function of the array index of each element of z.

x — (optional) A vector or two-dimensional array specifying the x-coordinates for the surface.

If x is a vector, each element of x specifies the x-coordinate for a column of z. For example, x(0) specifies the x-coordinate for z(0, *).

If x is a two-dimensional array, each element of x specifies the x-coordinate of the corresponding point in z (xij specifies the x-coordinate for zij).

y — (optional) A vector or two-dimensional array specifying the y-coordinates for the surface.

If y is a vector, each element of y specifies the y coordinate for a row of z. For example, y(0) specifies the y-coordinate for z (*, 0).

If y is a two-dimensional array, each element of y specifies the y-coordinate of the corresponding point in z (yij specifies the y-coordinate for zij).
Keywords

Name — Specifies a name to be used to create this object. If an undefined variable is used or no name specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.

Shades — An array expression of the same dimensions as z, containing the color index at each point. Alternately, an expression describing one color to be used for the entire surface or wireframe. If this keyword is omitted, a white surface is displayed.

To specify a single color, see vtkWINDOW (page 85). If a two-dimensional array of colors is specified (for an image overlay), the shades variable can be in any of these formats:

Fix(x, y) — A two-dimensional array of short integers or bytes specifying an index into the current PV-WAVE color table for each point. The RGB color for each point is obtained from the corresponding entry in the current PV-WAVE color table.

Long(x, y) — A two-dimensional array of long integers specifying the 24-bit color.

Float(3, x, y) — A floating point array of size (3, x, y) containing the normalized values specifying the red, green, and blue components of the color at each point.

Float(4,x) — A floating point array of size (4, x, y) containing the normalized values specifying the red, green, blue, and alpha components of the color at each point. The alpha component is the transparency where 0.0 is completely transparent and 1.0 is opaque.

Wireframe — When present and nonzero, a wire-frame mesh is drawn rather than a shaded surface.

LOD — If nonzero, a level-of-detail actor is created to aid in keeping a high frame-rate during frequent render requests due to user mouse interaction. If set to a value greater than 1, the number of points to use in the random cloud.

NoRotate — Does not perform any camera rotations. Used when a previous call to vtkSURFACE, vtkSCATTER or vtkPOLY SHADE has already set the camera angle.

NoAxes — If present and non zero, then no x, y, or z axes will be drawn.

NoErase — If nonzero, prevents the window from being erased to the background color before drawing the new scene. If not set, then all lights, cameras, and objects are removed from the scene before objects for the new scene are added.

TextColor — The color to use for text used for the axes titles. See vtkWINDOW for possible ways to specify the color. (Default: ‘white’)

These keywords are passed to vtkAXES:

TickSymbol  Labels  Sigfig  Format

Other keywords are listed below. For a description of each keyword, see Chapter 3, *Graphics and Plotting Keywords* in the *PV-WAVE Reference*.

**Discussion**

This procedure is similar to the SURFACE and SHADE_SURF procedures for PV-WAVE windows. Wireframes can be produced as well as surfaces shaded in one color or overlaid with an image. Transparency is also supported.

**Example**

This example demonstrates a surface created in a regular PV-WAVE window compared to one using vtkSURFACE.

```pikes=FLTARR(60,40)
s=DC_READ_FREE(!Data_Dir+'pikes elev.dat',pikes);
; Read in the data values for elevation.
snow=FLTARR(60,40)
s=DC_READ_FREE(!Data_Dir+'snowpack.dat',snow);
; Read in the data for snowpack
loadct,5
; Load a color table.
surface,pikes
; Create a wiremesh surface using a regular PV-WAVE window.
vtksurface,pikes/250,/wireframe
; Create a wiremesh surface using the VTK toolkit. Notice that the toolkit doesn't scale
; the data for you so in order to make sense out of the resulting graphic you need to
; scale the data yourself, in this example it was done by dividing by 250.
shade_surf,pikes,shades=bytscl(snow)
; Create a shaded surface using a regular PV-WAVE window.
vtksurface,pikes/250,shades=bytscl(snow)```
vtkSURFGEN Procedure

Generates a 3D surface from sampled points assumed to lie on a surface.

Usage

vtkSURFGEN, points

Input Parameters

points — A 3, n array of points that lie on a surface.

Keywords

Reverse — By default, the normals of the computed surface are inward facing. If outward normals are required, set this keyword.

Neighbors — An integer, the number of neighbors each points has. Use a larger value if the spread of points is not even. (Default: 20)

Spacing — A float, the spacing of the 3D sampling grid. If not set, the VTK class makes a reasonable guess.

Filename — An ASCII VTK file to create containing the dataset generated by the VTK filter.

Data — (Output). A returned associative array containing two keys: data("POINTS") is a 3, n float array containing the points generated by the VTK filter and data("VERTICES") is an n+1 element long array containing topology information for the generated dataset. A filename must be defined to have data returned by this keyword.

Name — A string, the name to be used to create this object. If an undefined variable is used or no name is specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.
**Color** — The color to use for the generated surface. See vtkWINDOW for possible ways to specify the color. (Default: ‘white’)

**Wireframe** — When present and nonzero, a wire-frame mesh is drawn rather than a shaded surface.

**LOD** — If nonzero, a level-of-detail actor is created to aid in keeping a high framerate during frequent render requests due to user mouse interaction. If set to a value greater than 1, the number of points to use in the random cloud.

This routine also accepts these keywords to control the initial camera and the axes (see vtkSURFACE and vtkAXES):

```
[XYZ]Range      Ax    Az    NoRotate
NoAxes          NoErase Charsize TextColor
Tickscale       Ticksymbol Labels Format
Sigfig
```

**Example**

```math
@math_startup
s = TRANSPOSE(random(100, /Sphere, Parameter=3))
vtkSURFGEN, s, /NoAxes
vtkSCATTER, s, /NoAxes, Symb=1, Color='red', Scale=0.5, $
   /NoRotate, /NoErase
```

**See Also**

vtkAXES, vtkSURFACE
**vtkTEXT Procedure**

Adds a text string.

**Usage**

vtkTEXT, string

**Input Parameters**

*string* — The scalar string containing the text that is to be output to the display surface. If not of string type, it is converted prior to use.

**Keywords**

*Name* — Specifies a name to be used to create this object. If an undefined variable is used or no name specified, then a random name is used. This name can be used in calls to vtkCOMMAND to modify this object.

*Position* — An array of three floating point numbers specifying the $x$, $y$, and $z$ position for the beginning of text. (Default: [0,0,0])

*Color* — The color to use for text. See vtkWINDOW (page 85) for possible ways to specify the color. (Default: ‘white’)

*Follow* — If nonzero, forces the text to always be facing the camera.

*Orientation* — An array of three floating point numbers specifying the $x$, $y$, and $z$ rotations of text in degrees. The actual rotations are performed in this order: $z$ then $x$ and finally $y$. This keyword has no effect if *Follow* is specified.

Keyword Charsize is also supported. For a description, see Chapter 3, *Graphics and Plotting Keywords* in the PV-WAVE Reference.

**Discussion**

This procedure is similar to the XYOUTS procedure for PV-WAVE windows.

**Example**

vtkText, "This is vtkText", color="red", charsize=10
**vtkTVRD Function**

Returns the contents of a VTK window as a bitmapped image.

**Usage**

\[image = \text{vtkTVRD}([\text{window\_index}])\]

**Input Parameters**

`window_index` — (optional) An integer specifying the index of an existing VTK window. If omitted, the current window is used.

**Keywords**

*Filename* — File path to store a temporary PPM file. Default is “wave.ppm.”

**Returned Value**

`image` — An array (3, width, height) of bytes containing the 24-bit image.

**Discussion**

This function works like TVRD for PV-WAVE windows. It uses vtkPPMWRITE and vtkPPMREAD to save and then read the contents of a window. The temporary file created is deleted when done.

**Example**

\[
vtkWINDOW, 7
\]
\[
vtkAXES
\]
\[
tv, \text{vtkTVRD}(7), /\text{TRUE}
\]

**See Also**

TVRD, vtkPPMWRITE, vtkPPMREAD
vtkUNSTRUCTUREDGRID Procedure

Passes data describing an unstructured grid to VTK.

Usage

vtkUNSTRUCTUREDGRID, points, cells, cell_types

Input Parameters

points — A two-dimensional array of floating point numbers of size (3, n) describing x, y, and z points.

cells — A Vector of integers describing cells, organized as vertex count followed by indices into Points, repeated for all cells.

Cell_types — A Vector of integers describing the cell type for each cell. Valid types are values between 1-12.

Keywords

Restore — An associative array containing all of the data and attributes for a poly-data dataset, usually created using the Save keyword. If this parameter is passed, then only the keywords Name and Filename can be used.

Name — Specifies a name to be used to create this data source. This name can be used in calls to vtkCOMMAND.

Filename — File to store data using standard VTK ASCII format.

Save — Returns the data in the specified variable stored in an associative array. Data is NOT sent to VTK if this parameter is specified.

Attributes — A list created using vtkADDATTRIBUTE containing one or more attributes associated with the points in the dataset.

Discussion

Arbitrary combinations of twelve (12) cell types, ranging from points, lines, polygons to voxels. See the VTK documentation, which can be downloaded from http://public.kitware.com, for more details on the data and attributes for the UnStructuredGrid dataset format.
**vtkWDELETE Procedure**

Closes a VTK window without shutting down the Tcl process.

**Usage**

vtkWDELETE [, window_index]

**Input Parameters**

*window_index* — (optional) An integer specifying the index of an existing VTK window. If omitted, the current window is used.

**Keywords**

All — If nonzero, causes all VTK windows to be closed.

**Discussion**

This procedure works like WDELETE for PV-WAVE windows. It closes an individual VTK window but does not shut down the Tcl process. Use vtkCLOSE to close all windows and shut down the spawned Tcl process.

**Example 1**

Deleting a window.

vtkwindow, 1
vtkwdelete

**Example 2**

vtkwindow, 1
vtkwindow, 2
vtkwindow, 3
vtkwdelete, /all

**See Also**

vtkCLOSE, vtkWINDOW
vtkWINDOW Procedure

Creates a VTK window.

Usage

vtkWINDOW [,window_index]

Input Parameters

window_index — (optional) An integer specifying the index of the newly created window.

If window_index is omitted, 0 is used as the index of the new window.

If the value of window_index specifies an existing window, the existing window is deleted and a new window is created.

Keywords

Free — If nonzero, creates a window using an unused window index. This keyword can be used instead of specifying the window_index parameter.

NoRender — If nonzero, prevents individual objects (vtkLIGHT, vtkAXES, vtkPOLYSHADE, etc.) from being rendered as they are added to the window. Specifying NoRender can speed up the initial display of a scene if you have multiple objects in it. If specified, you must manually call vtkRENDERWINDOW after you have added all of your objects to the window.

Background — Background color for the window. The color can be specified in any of the following ways (the color ‘red’ is used here as an example):

‘red’ See the file <vni>/vtk-3_2/lib/vtkcolornames.pro for a complete list of supported color names, where <vni> is the path to the PV-WAVE installation.

‘FF0000’XL A long integer hexadecimal value specifying the 24-bit color.

[1.0, 0.0, 0.0] A three-element vector of normalized floating point values specifying the red, green, and blue components of the color.
If a short byte or short integer value is passed, the RGB color is obtained from the corresponding entry in the current PV-WAVE color table. In this case, when TEK_COLOR has been called, color index 2 is red.

/NoInteract — If nonzero, indicates that you do not wish to provide the standard set of mouse controls for viewing the 3D scene. The resulting scene can be manipulated only by programmatically setting the positional parameters for objects or cameras.

XPos, YPos — The x and y positions of the lower-left corner of the new window, specified in device coordinates.

If no position is specified, a position of (0,0) is used.

XSize — The width of the window, in pixels. (Default: 400)

YSize — The height of the window, in pixels. (Default: 400)

Discussion
This procedure is similar to the PV-WAVE WINDOW command. It allows the created window to have built-in interaction associated with it.

Example 1
This example shows how to bring up a VTK window.

vtkWINDOW, 1

Example 2
This example shows how to bring up a VTK window with a blue background and with the mouse controls disabled. windownum is the number of the free window.

vtkWINDOW, windownum, /Free, background=’blue’, /noInteract

See Also
vtkRENDERWINDOW, vtkCLOSE, vtkDELETE, vtkERASE, vtkWSET
vtkWRITEVRML Procedure

Creates a Virtual Reality Modeling Language file (VRML .wrl file) from a scene in a VTK window.

Usage

vtkWRITEVRML, filename [, WindowID=id, Speed=s]

Input Parameters

Filename — A string, the file to write (should end in “.wrl”).

Keywords

Windowid — An integer, the VTK window to use as the source. (Default: the currently active VTK window)

Speed — A float, the navigation speed. (Default: 4.0)

Discussion

To view a VRML .wrl file in a browser, you need a plug-in. Consult the FAQ at http://www.vrml.org for current information and to obtain a plug-in.

See Also

VRML Routines
**vtkWSET Procedure**

Sets the active VTK window.

**Usage**

vtkWSET [, window_index]

**Input Parameters**

`window_index` — (optional) An integer specifying the index of an existing VTK window. If omitted, the current window is used.

**Keywords**

None.

**Discussion**

This procedure works like WSET for PV-WAVE windows, which is used to select the current, or "active" window to be used by the VTK routines.

**Example**

Setting the window to the first window opened.

```
vtkwindow, 1
vtkwindow, 2
vtkwindow, 3
vtkaxes
vtkwset, 1
vtkaxes
```

**See Also**

vtkWINDOW, WSET
WgOrbit Procedure

Creates an interactive window for viewing objects.

Usage

WgOrbit, vertices, polygons, parent, shell

Input Parameters

vertices — A (3,n) array of points on the surfaces of the objects.

polygons — A vector defining polygons which describe the surfaces: it is a concatenation of vectors of the form \([m,i_1,\ldots,i_m]\) where \(m\) is the number of vertices defining a polygon and where \(vertices(*,i_1),\ldots,vertices(*,i_m)\) are those vertices arranged in counter-clockwise order when viewed from outside the object.

parent — (optional) The widget ID of the parent widget.

Output Parameters

shell — (optional) The ID of the newly created widget.

Keywords

position — A two-element vector positioning the widget’s upper-left corner (measured in pixels from the upper-left corner of the screen).

shades — A vector specifying the color for each vertex.

size — Two-element vector specifying window size. The default is [500,500].

title — A string specifying the title for the widget.

wid — (output) The window ID of the graphics window.

Examples

POLY_SPHERE, 1, 10, 10, v, p

\[v(1:2,*\) = [ 2*v(1,*), 3*v(2,*) ]\]

WgOrbit, v, p
WIN32_PICK_PRINTER Function

Displays a Windows printer dialog.

Usage

\[ \text{printer\_name} = \text{WIN32\_PICK\_PRINTER}( ) \]

Input Parameters

None.

Returned Value

\[ \text{printername} \] — The name of the printer.

Keywords

None.

See Also

WIN32_PICK_FONT
New PV-WAVE:Database Connection Functions

This section lists the new functions that have been added to PV-WAVE:Database Connection for version 7.5. For complete descriptions, see the PV-WAVE:Database Connection User’s Guide.

DB_GET_BINARY Function

Returns binary large objects (BLOBS) from a DBMS (database management system) server.

Usage

\[
\text{list}_\text{var} = \text{DB_GET_BINARY}(\text{handle}, \text{sql}_\text{query})
\]

Input Parameters

- \text{handle} — DBMS connection handle (returned by DB_CONNECT).
- \text{sql}_\text{query} — A string containing an SQL statement to execute on the DBMS server. It must be a query (SELECT) statement.

Returned Value

- \text{list}_\text{var} — A PV-WAVE LIST variable, one for each row in the query. Each element in the LIST is a PV-WAVE array of type BYTE.

Keywords

None.

Discussion

Since binary large objects (BLOBS) are transmitted from most DBMS systems in a different way from other data types, using DB_SQL to handle BLOBS would compromise performance.

For queries that return more than one row, specify the order of the rows with the \text{ORDER BY} clause in the \text{sql}_\text{query}.

\underline{NOTE} One column will cause an error.

\underline{CAUTION} The value of \text{sql}_\text{query} is subject to the following restrictions:

- It must be a query. UPDATE, INSERT, and/or DELETE will cause an error.
- It must only return one column. Queries that return more than one column will cause an error.
**NULL_PROCESSOR Function**

Facilitates the use of the Null_Info keyword for the DB_SQL function by extracting the list of rows containing missing data for one or more columns.

**Usage**

```
table = NULL_PROCESSOR(null_info_object,['col1','col2',...,coln'],Comp=comp)
```

**Input Parameters**

- `null_info_object` — The object returned by the Null_Info keyword in the DB_SQL call.
- `col` — The list of column names.

**Keywords**

- `Comp=comp` — Produces the complement to the result, that is, the result contains a list of rows with missing data. `comp` contains a list of rows with no missing data.

**Discussion**

Assuming the following use of the DB_SQL Null_Info keyword:

```
table = db_sql(db_connect('oracle', 'user_id/user_pw'), 'select * from blanktest', null_info=foo)
```

where blanktest contains the data given below, which has missing data for ID_NO in the 4th, 9th, and 11th rows and missing data for ANIMAL_NAME in the 3rd, 8th, and 10th rows.
Then,

\[
\text{jjj} = \text{NULL\_PROCESSOR(\text{foo}, ['ID\_NO', 'ANIMAL\_NAME'], \text{Comp} = \text{comp})}
\]

produces the results

\[
\begin{align*}
\text{jjj} & = 2 \quad 3 \quad 7 \quad 8 \quad 9 \quad 10 \\
\text{comp} & = 0 \quad 1 \quad 4 \quad 5 \quad 6
\end{align*}
\]

This output can be utilized as in the following examples.

\[\text{Table2} = \text{table(\text{comp})}\]

produces a table with only rows and no missing values or as in the table given above.

<table>
<thead>
<tr>
<th>ID_NO</th>
<th>ANIMAL_NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>golden</td>
</tr>
<tr>
<td>2</td>
<td>chirpy</td>
</tr>
<tr>
<td>5</td>
<td>KC</td>
</tr>
<tr>
<td>6</td>
<td>skip</td>
</tr>
<tr>
<td>7</td>
<td>sparky</td>
</tr>
<tr>
<td>10</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ID_NO</th>
<th>ANIMAL_NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>golden</td>
</tr>
<tr>
<td>2</td>
<td>chirpy</td>
</tr>
<tr>
<td>5</td>
<td>KC</td>
</tr>
<tr>
<td>6</td>
<td>skip</td>
</tr>
<tr>
<td>7</td>
<td>sparky</td>
</tr>
</tbody>
</table>
Then, 

\[ \text{Table3=table(jjj)} \]

produces a table containing only rows with missing data (note how zeros have been substituted for values of ID_NO that are missing).

<table>
<thead>
<tr>
<th>ID_NO</th>
<th>ANIMAL_ NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>harry</td>
</tr>
<tr>
<td>8</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>sneakers</td>
</tr>
<tr>
<td>10</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>harvey</td>
</tr>
</tbody>
</table>

Instead, if you want only the locations where one field is missing, a different db_sql call, \[ jji=foopro(fooo,['ID_NO'],Comp=comp) \], returns an array, \[ jji \], with the rows where ID_NO is missing (3 8 10).

Remember that rows are counted beginning with 0.
New PV-WAVE:IMSL Mathematics Commands

This section lists the new functions and procedures that have been added to PV-WAVE:IMSL Mathematics for version 7.5.

Chapter 4: Quadrature

INTFCN_QMC Function

Integrates a function on a hyper-rectangle using a quasi-Monte Carlo method.

Usage

\[
\text{result} = \text{INTFCN\_QMC}(f, a, b)
\]

Input Parameters

- \(f\) — Scalar string specifying the user-supplied function to be integrated. Function \(f\) accepts as input an array of data points at which the function is to be evaluated and returns the scalar value of the function.
- \(a\) — One-dimensional array containing the lower limits of integration.
- \(b\) — One-dimensional array containing the upper limits of integration.

Returned Value

The value of

\[
\int_{a_n}^{b_n-1} \cdots \int_{a_0}^{b_0-1} f(x_0, \ldots, x_{n-1}) \, dx_{n-1} \cdots
\]

is returned. If no value can be computed, then NaN is returned.

Input Keywords

- **Err_Abs** — Absolute accuracy desired.
  
  Default: \(\text{Err\_Abs} = 1.e-4\).
- **Err_Rel** — Relative accuracy desired.
Default: Err_rel = 1.e-4.

**Max_Evals** — Number of evaluations allowed.

Default: No limit

**Base** — The value of BASE used to compute the Faure sequence.

**Skip** — The value of SKIP used to compute the Faure sequence.

**Double** — If present and nonzero, double precision is used.

**Output Keywords**

**Err_est** — Named variable into which an estimate of the absolute value of the error is stored.

**Discussion**

Integration of functions over hypercubes by direct methods, such as INTFCN-HYPER, is practical only for fairly low dimensional hypercubes. This is because the amount of work required increases exponential as the dimension increases.

An alternative to direct methods is Monte Carlo, in which the integral is evaluated as the value of the function averaged over a sequence of randomly chosen points. Under mild assumptions on the function, this method will converge like $1/n^{1/2}$, where $n$ is the number of points at which the function is evaluated.

It is possible to improve on the performance of Monte Carlo by carefully choosing the points at which the function is to be evaluated. Randomly distributed points tend to be non-uniformly distributed. The alternative to a sequence of random points is a *low-discrepancy* sequence. A low-discrepancy sequence is one that is highly uniform.

This function is based on the low-discrepancy Faure sequence, as computed by FAURE_NEXT_PT.

**Example**

```fortran
FUNCTION F, x
    S = 0.0
    sign = -1.0
    FOR i = 0, N_ELEMENTS(x)-1 DO BEGIN
        prod = 1.0
        FOR j = 0, i DO BEGIN
```

96 Chapter 2: New Commands

PV-WAVE 7.5 New Features Guide
prod = prod*x(j)
END
S = S + sign*prod
sign = -sign
END
RETURN, s
END

ndim = 10
a = FLTARR(ndim)
a(*) = 0
b = FLTARR(ndim)
b(*) = 1
result = intfcn_qmc( 'f', a, b)
PM, result
-0.333010

Chapter 9: Special Functions

CUM_INTR Function

Evaluates the cumulative interest paid between two periods.

Usage

result = CUM_INTR (rate, n_periods, present_value, start, end_per, when)

Input Parameters

rate — Interest rate.
n_periods — Total number of payment periods. n_periods cannot be less than or equal to 0.
present_value — The current value of a stream of future payments, after discounting the payments using some interest rate.
**start** — Starting period in the calculation. *start* cannot be less than 1; or greater than *end_per*.

**end_per** — Ending period in the calculation.

**when** — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.

**Returned Value**

**result** — The cumulative interest paid between the first period and the last period. If no result can be computed, NaN is returned.

**Input Keywords**

**Double** — If present and nonzero, double precision is used.

**Discussion**

Function CUM_INTR evaluates the cumulative interest paid between the first period and the last period.

It is computed using the following:

\[
\sum_{i=start}^{end\_per} interest_i
\]

where *interest_i* is computed from the function INT_PAYMENT for the *i*th period.

**Example**

In this example, CUM_INTR computes the total interest paid for the first year of a 30-year $200,000 loan with an annual interest rate of 7.25%. The payment is made at the end of each month.

```
PRINT, CUM_INTR(0.0725 / 12, 12*30, 200000., 1, 12, 0)
-14436.5
```
**CUM_PRINC Function**

Evaluates the cumulative principal paid between two periods.

**Usage**

\[ result = \text{CUM_PRINC} \left( rate, n\_periods, \text{present\_value}, \text{start}, \text{end\_per}, \text{when} \right) \]

**Input Parameters**

- **rate** — Interest rate.
- **n\_periods** — Total number of payment periods. \( n\_periods \) cannot be less than or equal to 0.
- **present\_value** — The current value of a stream of future payments, after discounting the payments using some interest rate.
- **start** — Starting period in the calculation. \( start \) cannot be less than 1; or greater than \( \text{end\_per} \).
- **end\_per** — Ending period in the calculation.
- **when** — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.

**Returned Value**

- **result** — The cumulative principal paid between the first period and the last period. If no result can be computed, NaN is returned.

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.

**Discussion**

Function CUM_PRINC evaluates the cumulative principal paid between the first period and the last period.
It is computed using the following:

\[
\sum_{i=\text{start}}^{\text{end} \text{ per}} \text{principal}_i
\]

where \(\text{principal}_i\) is computed from the function PRINC_PAYMENT for the \(i\)th period.

**Example**

In this example, CUM_PRINC computes the total principal paid for the first year of a 30-year $200,000 loan with an annual interest rate of 7.25%. The payment is made at the end of each month.

\[
\text{PRINT, CUM_PRINC}(0.0725 / 12, 12*30, 200000., 1, 12, 0)
\]

\[-1935.73\]

---

**DEPRECIATION_DB Function**

Evaluates the depreciation of an asset using the fixed-declining balance method.

**Usage**

\[
\text{result} = \text{DEPRECIATION_DB}(\text{cost, salvage, life, period, month})
\]

**Input Parameters**

- **cost** — Initial value of the asset.
- **salvage** — The value of an asset at the end of its depreciation period.
- **life** — Number of periods over which the asset is being depreciated.
- **period** — Period for which the depreciation is to be computed. \(\text{period}\) cannot be less than or equal to 0, and cannot be greater than \(\text{life} + 1\).
- **month** — Number of months in the first year. \(\text{month}\) cannot be greater than 12 or less than 1.
Returned Value

result — The depreciation of an asset for a specified period using the fixed-declining balance method. If no result can be computed, NaN is returned.

Input Keywords

Double — If present and nonzero, double precision is used.

Discussion

Function DEPRECIATION_DB computes the depreciation of an asset for a specified period using the fixed-declining balance method. Function DEPRECIATION_DB varies depending on the specified value for the argument period, see table below.

<table>
<thead>
<tr>
<th>Period</th>
<th>Formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>period = 1</td>
<td>( cost \times rate \times \frac{month}{12} )</td>
</tr>
<tr>
<td>period = life</td>
<td>((cost - \text{total depreciation from periods}) \times rate \times \frac{12 - month}{12})</td>
</tr>
<tr>
<td>period other than 1 or life</td>
<td>((cost - \text{total depreciation from prior periods}) \times rate)</td>
</tr>
</tbody>
</table>

where

\[
rate = 1 - \left( \frac{\text{salvage}}{\text{cost}} \right)^{\frac{1}{\text{life}}}
\]

NOTE: rate is rounded to three decimal places.

Example

In this example, DEPRECIATION_DB computes the depreciation of an asset, which costs $2,500 initially, a useful life of 3 periods and a salvage value of $500, for each period.

ans = fltarr(4)
life = 3
cost = 2500
salvage = 500
life = 3
month = 6
for period = 1, life+1 DO $
ans(period-1) = depreciation_db(cost, salvage, life, $ 
   period, month)
PM, ans
   518.750
   822.219
   480.998
   140.692

**DEPRECIATION_DDB Function**

Evaluates the depreciation of an asset using the double-declining balance method.

**Usage**

\[ result = \text{DEPRECIATION_DDB} \left( \text{cost}, \text{salvage}, \text{life}, \text{period}, \text{factor} \right) \]

**Input Parameters**

- **cost** — Initial value of the asset.
- **salvage** — The value of an asset at the end of its depreciation period.
- **life** — Number of periods over which the asset is being depreciated.
- **period** — Period for which the depreciation is to be computed. **period** cannot be greater than **life**.
- **factor** — Rate at which the balance declines. **factor** must be positive.

**Returned Value**

**result** — The depreciation of an asset using the double-declining balance method for a period specified by the user. If no result can be computed, NaN is returned.
Input Keywords

Double — If present and nonzero, double precision is used.

Discussion

Function DEPRECIATION_DDB computes the depreciation of an asset using the double-declining balance method for a specified period. It is computed using the following:

\[
\left(\text{cost} - \text{salvage} \times \text{total depreciation from prior periods}\right) \times \left(\frac{\text{factor}}{\text{life}}\right)
\]

Example

In this example, DEPRECIATION_DDB computes the depreciation of an asset, which costs $2,500 initially, lasts 24 periods and a salvage value of $500, for each period.

\[
\begin{align*}
\text{ans} &= \text{fltarr}(24) \\
\text{life} &= 24 \\
\text{cost} &= 2500 \\
\text{salvage} &= 500 \\
\text{factor} &= 2 \\
\text{FOR period} &= 1, \text{life DO} \$
\end{align*}
\]

\[
\begin{align*}
\text{ans}(\text{period} - 1) &= \text{depreciation_ddb(cost, salvage, life, period, factor)} \\
\text{PM, ans}
\end{align*}
\]

\[
\begin{align*}
208.333 & \\
190.972 & \\
175.058 & \\
160.470 & \\
147.097 & \\
134.839 & \\
123.603 & \\
113.302 & \\
103.860 & \\
95.2054 & \\
\end{align*}
\]
DEPRECIATION_SLN Function

Evaluates the depreciation of an asset using the straight-line method.

Usage

\[ result = \text{DEPRECIATION\_SLN}(\text{cost, salvage, life}) \]

Input Parameters

\textit{cost} — Initial value of the asset.
\textit{salvage} — The value of an asset at the end of its depreciation period.
\textit{life} — Number of periods over which the asset is being depreciated.

Returned Value

\textit{result} — The straight line depreciation of an asset for its life. If no result can be computed, NaN is returned.
Input Keywords

*Double* — If present and nonzero, double precision is used.

Discussion

Function DEPRECIATION_SLN computes the straight line depreciation of an asset for its life.

It is computed using the following:

\[
\frac{\text{cost}-\text{salvage}}{\text{life}}
\]

Example

In this example, DEPRECIATION_SLN computes the depreciation of an asset, which costs $2,500 initially, lasts 24 periods and a salvage value of $500.

```
PRINT, DEPRECIATION_SLN(2500, 500, 24)
83.3333
```

**DEPRECIATION_SYD Function**

Evaluates the depreciation of an asset using the sum-of-years digits method.

Usage

\[
\text{result} = \text{DEPRECIATION_SYD} (\text{cost}, \text{salvage}, \text{life}, \text{period})
\]

Input Parameters

*cost* — Initial value of the asset.

*salvage* — The value of an asset at the end of its depreciation period.

*life* — Number of periods over which the asset is being depreciated.

*period* — Period for which the depreciation is to be computed. *period* cannot be greater than *life*. 
Returned Value

`result` — The sum-of-years digits depreciation of an asset for a specified period. If no result can be computed, NaN is returned.

Input Keywords

`Double` — If present and nonzero, double precision is used.

Discussion

Function `DEPRECIATION_SYD` computes the sum-of-years digits depreciation of an asset for a specified period.

It is computed using the following:

\[(cost - salvage)(period) \frac{(life + 1)(life)}{2}\]

Example

In this example, `DEPRECIATION_SYD` computes the depreciation of an asset, which costs $25,000 initially, lasts 15 years, and a salvage value of $5,000, for the 14th year.

```plaintext
PRINT, DEPRECIATION_SYD(25000, 5000, 15, 14)
333.333
```

**DEPRECIATION_VDB Function**

Evaluates the depreciation of an asset for any given period using the variable-declining balance method.

**Usage**

`result = DEPRECIATION_VDB (cost, salvage, life, start, end_per, factor, sln)`

**Input Parameters**

`cost` — Initial value of the asset.

`salvage` — The value of an asset at the end of its depreciation period.
**life**—Number of periods over which the asset is being depreciated.

**start**—Starting period in the calculation. *start* cannot be less than 1; or greater than *end_per*.

**end_per**—Final period for the calculation. *end_per* cannot be greater than *life*.

**factor**—Rate at which the balance declines. *factor* must be positive.

**sln**—If equal to zero, do not switch to straight-line depreciation even when the depreciation is greater than the declining balance calculation.

**Returned Value**

**result**—The depreciation of an asset for any given period, including partial periods, using the variable-declining balance method. If no result can be computed, NaN is returned.

**Input Keywords**

**Double**—If present and nonzero, double precision is used.

**Discussion**

Function DEPRECIATION_VDB computes the depreciation of an asset for any given period using the variable-declining balance method using the following:

If \( sln = 0 \)

\[
\sum_{i=\text{start}+1}^{\text{end}_\text{per}} \text{ddbi}_i
\]

If \( sln \neq 0 \)

\[
A + \sum_{k=1}^{\text{end}_\text{per}} \frac{\text{cost} - \text{salvage}}{\text{end} - k + 1}
\]

where \( \text{ddbi}_i \) is computed from the function DEPRECIATION_DDB for the \( i \)th period. \( k \) = the first period where straight-line depreciation is greater than

\[
A = \sum_{i=\text{start}+1}^{i-1} \text{ddbi}_i
\]
the depreciation using the double-declining balance method.

**Example**

In this example, DEPRECIATION_VDB computes the depreciation of an asset between the 10th and 15th year, which costs $25,000 initially, lasts 15 years, and has a salvage value of $5,000.

```
PRINT, DEPRECIATION_VDB(25000., 5000., 15, 10, 15, 2, 0)
```

976.69

---

**DOLLAR_DECIMAL Function**

Converts a fractional price to a decimal price.

**Usage**

```
result = DOLLAR_DECIMAL (fractional_num, fraction)
```

**Input Parameters**

- `fractional_num` — Whole number of dollars plus the numerator, as the fractional part.
- `fraction` — Denominator of the fractional dollar. `fraction` must be positive.

**Returned Value**

- `result` — The dollar price expressed as a decimal number. The dollar price is the whole number part of fractional-dollar plus its decimal part divided by fraction. If no result can be computed, NaN is returned.

**Input Keywords**

- `Double` — If present and nonzero, double precision is used.

**Discussion**

Function DOLLAR_DECIMAL converts a dollar price, expressed as a fraction, into a dollar price, expressed as a decimal number.
It is computed using the following:

\[ idollar + \left\lfloor \frac{\text{fractional}_\text{num} - idollar}{\text{fraction}} \right\rfloor \times 10^{(ifrac+1)} \]

where \( idollar \) is the integer part of \( \text{fractional}_\text{num} \), and \( ifrac \) is the integer part of \( \log(\text{fraction}) \).

**Example**

In this example, \text{DOLLAR\_DECIMAL} converts \$1 \frac{1}{4} to \$1.25.

\begin{verbatim}
PRINT, DOLLAR\_DECIMAL(1.1, 4)
1.25000
\end{verbatim}
Discussion

Function DOLLAR_FRACTION converts a dollar price, expressed as a decimal number, into a dollar price, expressed as a fractional price. If no result can be computed, NaN is returned.

It can be found by solving the following

$$idollar + \left[ \frac{\text{decimal}_\text{dollar} - idollar}{10^\text{ifrac}} \right] / \text{fraction}$$

where $idollar$ is the integer part of the $\text{decimal}_\text{dollar}$, and $ifrac$ is the integer part of $\log(\text{fraction})$.

Example

In this example, DOLLAR_FRACTION converts $1.25 to $1 1/4.

```plaintext
PRINT, DOLLAR_FRACTION(1.25, 4)
1.10000
```

---

**EFFECTIVE_RATE Function**

Evaluates the effective annual interest rate.

**Usage**

$$result = \text{EFFECTIVE_RATE} (\text{nominal}_\text{rate}, n_\text{periods})$$

**Input Parameters**

- $\text{nominal}_\text{rate}$ — The interest rate as stated on the face of a security.
- $n_\text{periods}$ — Number of compounding periods per year.

**Returned Value**

- $result$ — The effective annual interest rate. If no result can be computed, NaN is returned.
Input Keywords

*Double* — If present and nonzero, double precision is used.

Discussion

Function **EFFECTIVE_RATE** computes the continuously-compounded interest rate equivalent to a given periodically-compounded interest rate. The nominal interest rate is the periodically-compounded interest rate as stated on the face of a security.

It can found by solving the following:

\[
\left(1 + \frac{\text{nominal\_rate}}{\text{n\_periods}}\right)^{\text{n\_periods}} - 1
\]

Example

In this example, **EFFECTIVE_RATE** computes the effective annual interest rate of the nominal interest rate, 6\%, compounded quarterly.

```plaintext
PRINT, EFFECTIVE_RATE(0.06, 4)
0.0613635
```

**FUTURE_VALUE** Function

Evaluates the future value of an investment.

Usage

```plaintext
result = FUTURE_VALUE (rate, n\_periods, payment, present\_value, when)
```

Input Parameters

*rate* — Interest rate.

*n\_periods*— Total number of payment periods.

*payment* — Payment made in each period.

*present\_value* — The current value of a stream of future payments, after discounting the payments using some interest rate.
**when** — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.

**Returned Value**

**result** — The future value of an investment. If no result can be computed, NaN is returned.

**Input Keywords**

**Double** — If present and nonzero, double precision is used.

**Discussion**

Function FUTURE_VALUE computes the future value of an investment. The future value is the value, at some time in the future, of a current amount and a stream of payments.

It can be found by solving the following:

If $rate=0$

$$present\_value + (payment)(n\_periods) + future\_value=0$$

If $rate \neq 0$

$$present\_value(1+rate)^{n\_periods} + payment\left[1+rate\left(\frac{n\_periods - 1}{rate}\right)\right] + future\_value=0$$

**Example**

In this example, FUTURE_VALUE computes the value of $30,000 payment made annually at the beginning of each year for the next 20 years with an annual interest rate of 5%.

```plaintext
PRINT, FUTURE_VALUE(0.05, 20, -30000.00, -30000.00, 1)
1.12118e+06
```
**FUTURE_VAL_SCHD Function**

Evaluates the future value of an initial principal taking into consideration a schedule of compound interest rates.

**Usage**

\[ result = \text{FUTURE\_VAL\_SCHD}(\text{principal}, \text{schedule}) \]

**Input Parameters**

- **principal** — Principal or present value.
- **schedule** — One-dimensional array of interest rates to apply.

**Returned Value**

- **result** — The future value of an initial principal after applying a schedule of compound interest rates. If no result can be computed, NaN is returned.

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.

**Discussion**

Function FUTURE\_VAL\_SCHD computes the future value of an initial principal after applying a schedule of compound interest rates.

It is computed using the following with \( count = \text{N\_ELEMENTS}(\text{schedule}) \):

\[ \sum_{i=1}^{count} (\text{principal} \times \text{schedule}_i) \]

where \( \text{schedule}_i \) = interest rate at the \( i \)th period.

**Example**

In this example, FUTURE\_VAL\_SCHD computes the value of a $10,000 investment after 5 years with interest rates of 5%, 5.1%, 5.2%, 5.3% and 5.4%, respectively.
principal = 10000.0
schedule = [ .050, .051, .052, .053, .054 ]
PRINT, FUTURE_VAL_SCHD(principal, schedule)
       12884.8

**INT_PAYMENT Function**

Evaluates the interest payment for an investment for a given period.

**Usage**

\[
result = \text{INT\_PAYMENT} (rate, period, n\_periods, present\_value, future\_value, when)
\]

**Input Parameters**

*rate* — Interest rate.

*period* — Payment period.

*n\_periods* — Total number of periods.

*present\_value* — The current value of a stream of future payments, after discounting the payments using some interest rate.

*future\_value* — The value, at some time in the future, of a current amount and a stream of payments.

*when* — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.

**Returned Value**

*result* — The interest payment for an investment for a given period. If no result can be computed, NaN is returned.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.
Discussion

Function INT_PAYMENT computes the interest payment for an investment for a given period.

It is computed using the following:

\[
present\_value \left(1 + rate\right)^{n\_periods-1} + payment \left(1 + rate\right)^{n\_periods-1} \times \frac{\left(1 + rate\right)^{n\_periods-1} - rate}{rate}
\]

Example

In this example, INT_PAYMENT computes the interest payment for the second year of a 25-year $100,000 loan with an annual interest rate of 8%. The payment is made at the end of each period.

\[
\text{PRINT, INT\_PAYMENT(0.08, 2, 25, 100000.00, 0.0, 0)}
\]

-7890.57

INT_RATE_ANNUITY Function

Evaluates the interest rate per period of an annuity.

Usage

\[
result = \text{INT\_RATE\_ANNUITY (n\_periods, payment, present\_value, future\_value, when)}
\]

Input Parameters

- \(n\_periods\) — Total number of periods.
- \(payment\) — Payment made each period.
- \(present\_value\) — The current value of a stream of future payments, after discounting the payments using some interest rate.
- \(future\_value\) — The value, at some time in the future, of a current amount and a stream of payments.
when — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.

Returned Value

result — The interest rate per period of an annuity. If no result can be computed, NaN is returned.

Input Keywords

Double — If present and nonzero, double precision is used.
Xguess — If present, the value is used as the initial guess at the interest rate.
Highest — If present, the value is used as the maximum value of the interest rate allowed.

Discussion

Function INT_RATE_ANNUITY computes the interest rate per period of an annuity. An annuity is a security that pays a fixed amount at equally spaced intervals.
It can be found by solving the following:

If $rate = 0$
\[
\text{present}_\text{value} + (\text{payment})(n\_\text{periods}) + \text{future}_\text{value} = 0
\]

If $rate \neq 0$
\[
\text{present}_\text{value}(1 + rate)^{n\_\text{periods}} + \text{payment}[1 + rate(\text{when})] \left(\frac{(1 + rate)^{n\_\text{periods}} - 1}{rate}\right) + \text{future}_\text{value} = 0
\]

**Example**

In this example, INT_RATE_ANNUITY computes the interest rate of a $20,000 loan that requires 70 payments of $350 to pay off the loan.

\[
\text{PRINT, 12*INT_RATE_ANNUITY(70, \ -350, 20000, 0, 1)}
\]

\[
0.0734513
\]

**INT_RATE_RETURN Function**

Evaluates the internal rate of return for a schedule of cash flows.

**Usage**

\[
result = \text{INT_RATE_RETURN (values)}
\]

**Input Parameters**

values — One-dimensional array of cash flows which occur at regular intervals, which includes the initial investment.

**Returned Value**

result — The internal rate of return for a schedule of cash flows. If no result can be computed, NaN is returned.
Input Keywords

*Double* — If present and nonzero, double precision is used.

*Xguess* — If present, the value is used as the initial guess at the internal rate of return.

*Highest* — If present, the value is used as the maximum value of the internal rate of return allowed.

**Discussion**

Function INT_RATE_RETURN computes the internal rate of return for a schedule of cash flows. The internal rate of return is the interest rate such that a stream of payments has a net present value of zero.

It is found by solving the following with \(\text{count} = \text{N\_ELEMENTS (values)}\):

\[
0 = \sum_{i=1}^{\text{count}} \frac{\text{value}_i}{(1+\text{rate})^i}
\]

where \(\text{value}_i = \text{the } i\text{th cash flow}, \text{rate} \text{ is the internal rate of return.}

**Example**

In this example, INT_RATE_RETURN computes the internal rate of return for nine cash flows, $-800, $800, $800, $600, $600, $800, $800, $700 and $3,000, with an initial investment of $4,500.

\[
\text{values} = [\begin{array}{c}
-4500., -800., 800., 800., 600., 600., 800., 800., 700., 3000.
\end{array} ]
\]

\[
\text{PRINT, INT\_RATE\_RETURN(values)}
\]

\[
0.0720820
\]

**INT_RATE_SCHD Function**

Evaluates the internal rate of return for a schedule of cash flows. It is not necessary that the cash flows be periodic.

**Usage**

\[
\text{result} = \text{INT\_RATE\_SCHD (values, dates)}
\]
Input Parameters

$values$ — One-dimensional array of cash flows, which includes the initial investment.

dates — One-dimensional array of dates cash flows are made. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

Returned Value

$result$ — The internal rate of return for a schedule of cash flows that is not necessarily periodic. If no result can be computed, NaN is returned.

Input Keywords

$Double$ — If present and nonzero, double precision is used.

$Xguess$ — If present, the value is used as the initial guess at the internal rate of return.

$Highest$ — If present, the value is used as the maximum value of the internal rate of return allowed.

Discussion

Function INT_RATE_SCHD computes the internal rate of return for a schedule of cash flows that is not necessarily periodic. The internal rate such that the stream of payments has a net present value of zero.

It can be found by solving the following with $count = N\_ELEMENTS (values)$:

\[
0 = \sum_{i=1}^{count} \frac{value_i}{(1+rate)^{\frac{d_i-d_1}{365}}}
\]

In the equation above, $d_i$ represents the $i$th payment date. $d_1$ represents the 1st payment date. $value_i$ represents the $i$th cash flow. $rate$ is the internal rate of return.
Example

In this example, INT_RATE_SCHD computes the internal rate of return for nine cash flows, $-800, $800, $800, $600, $600, $800, $800, $700 and $3,000, with an initial investment of $4,500.

```pawave
months = [1, 10, 5, 5, 6, 7, 8, 9, 10, 11]
days = [1, 1, 5, 5, 1, 1, 30, 15, 15, 1]
dates = VAR_TO_DT(years, months, days)
v = [-4500., -800, 800, 800., 600., 600, 800, 800, 700, 3000]
PRINT, INT_RATE_SCHD(v, dates)
   0.0769003
```

**MOD_INTERN_RATE Function**

Evaluates the modified internal rate of return for a schedule of periodic cash flows.

**Usage**

```pawave
result = MOD_INTERN_RATE (values, finance_rate, reinvest_rate)
```

**Input Parameters**

`values` — One-dimensional array of cash flows.

`finance_rate` — Interest paid on the money borrowed.

`reinvest_rate` — Interest rate received on the cash flows.

**Returned Value**

`result` — The modified internal rate of return for a schedule of periodic cash flows. If no result can be computed, NaN is returned.

**Input Keywords**

`Double` — If present and nonzero, double precision is used.
Discussion

Function MOD_INTERN_RATE computes the modified internal rate of return for a schedule of periodic cash flows. The modified internal rate of return differs from the ordinary internal rate of return in assuming that the cash flows are reinvested at the cost of capital, not at the internal rate of return.

It also eliminates the multiple rates of return problem.

It is computed using the following:

\[
\left\{ \frac{- (pnpv)(1 + reinvest\_rate)^{n\_periods}}{(nnpv)(1 + finance\_rate)} \right\}^{\frac{1}{n\_periods-1}} - 1
\]

where \( pnpv \) is calculated from the function NET_PRES_VALUE for positive values in \( values \) using \( reinvest\_rate \), and where \( nnpv \) is calculated from the function NET_PRES_VALUE for negative values in \( values \) using \( finance\_rate \).

Example

In this example, MOD_INTERN_RATE computes the modified internal rate of return for an investment of $4,500 with cash flows of $-800, $800, $800, $600, $600, $800, $800, $700 and $3,000 for 9 years.

```plaintext
value = [-4500., -800., 800., 800., 600., 600., 800., 800., 700., 3000.]
finance_rate = .08
reinvest_rate = .055
PRINT, MOD_INTERN_RATE(value, finance_rate, reinvest_rate)
0.0665972
```

NET_PRES_VALUE Function

Evaluates the net present value of a stream of unequal periodic cash flows, which are subject to a given discount rate.

Usage

```plaintext
result = NET_PRES_VALUE (rate, values)
```
**Input Parameters**

*rate* — Interest rate per period.

*values* — One-dimensional array of equally-spaced cash flows.

**Returned Value**

*result* — The net present value of an investment. If no result can be computed, NaN is returned.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.

**Discussion**

Function NET_PRES_VALUE computes the net present value of an investment. Net present value is the current value of a stream of payments, after discounting the payments using some interest rate.

It is found by solving the following with \( count = N\_ELEMENTS(values) \):

\[
\sum_{i=1}^{\text{count}} \frac{\text{value}_i}{(1 + \text{rate})^i}
\]

where \( \text{value}_i \) = the \( i \)th cash flow.

**Example**

In this example, NET_PRES_VALUE computes the net present value of a $10 million prize paid in 20 years ($50,000 per year) with an annual interest rate of 6%.

```plaintext
rate = 0.06
value = FLTARR(20)
value(*) = 500000.
PRINT, NET_PRES_VALUE(rate, value)
5.73496e+06
```
**NOMINAL_RATE Function**

Evaluates the nominal annual interest rate.

**Usage**

\[
\text{result} = \text{NOMINAL_RATE} (\text{effective_rate}, \text{n_periods})
\]

**Input Parameters**

- **effective_rate** — The amount of interest that would be charged if the interest was paid in a single lump sum at the end of the loan.
- **n_periods** — Number of compounding periods per year.

**Returned Value**

- **result** — The nominal annual interest rate. If no result can be computed, NaN is returned.

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.

**Discussion**

Function NOMINAL_RATE computes the nominal annual interest rate. The nominal interest rate is the interest rate as stated on the face of a security. It is computed using the following:

\[
\left[ \left(1 + \text{effective\_rate} \right)^{\frac{1}{\text{n\_periods}}} - 1 \right] * \text{n\_periods}
\]

**Example**

In this example, NOMINAL_RATE computes the nominal annual interest rate of the effective interest rate, 6.14\%, compounded quarterly.

\[
\text{PRINT, NOMINAL\_RATE(0.0614, 4)}
\]
**NUM_PERIODS Function**

Evaluates the number of periods for an investment for which periodic and constant payments are made and the interest rate is constant.

**Usage**

\[
\text{result} = \text{NUM_PERIODS} (\text{rate}, \text{payment}, \text{present\_value}, \text{future\_value}, \text{when})
\]

**Input Parameters**

- **rate** — Interest rate on the investment.
- **payment** — Payment made on the investment.
- **present\_value** — The current value of a stream of future payments, after discounting the payments using some interest rate.
- **future\_value** — The value, at some time in the future, of a current amount and a stream of payments.
- **when** — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.

**Returned Value**

- **result** — The number of periods for an investment.

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.

**Discussion**

Function NUM_PERIODS computes the number of periods for an investment based on periodic, constant payment and a constant interest rate.
It can be found by solving the following:

If \( rate = 0 \)
\[
present\_value + (payment)(n\_periods) + future\_value = 0
\]

If \( rate \neq 0 \)
\[
present\_value(1 + rate)^{n\_periods} + payment[(1 + rate)(when)]^{(1 + rate)^{n\_periods} - 1}\]
\[
+ future\_value = 0
\]

**Example**

In this example, NUM_PERIODS computes the number of periods needed to pay off a $20,000 loan with a monthly payment of $350 and an annual interest rate of 7.25%. The payment is made at the beginning of each period.

\[
PRINT, \ \text{NUM\_PERIODS}(0.0725 / 12, -350., 20000., 0., 1)
\]

\[
70
\]

**PAYMENT Function**

Evaluates the periodic payment for an investment.

**Usage**

\[
result = \text{PAYMENT}(rate, n\_periods, present\_value, future\_value, when)
\]

**Input Parameters**

- \( rate \) — Interest rate.
- \( n\_periods \) — Total number of periods.
- \( present\_value \) — The current value of a stream of future payments, after discounting the payments using some interest rate.
- \( future\_value \) — The value, at some time in the future, of a current amount and a stream of payments.
- \( when \) — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.
Returned Value

*result* — The periodic payment for an investment. If no result can be computed, NaN is returned.

Input Keywords

*Double* — If present and nonzero, double precision is used.

Discussion

Function PAYMENT computes the periodic payment for an investment. It can be found by solving the following:

If $rate = 0$

$$present_value + (payment)(n\_periods) + future\_value = 0$$

If $rate \neq 0$

$$present_value(1 + rate)^{n\_periods} + payment\{1 + rate\(when\)\}\(1 + rate\)^{n\_periods} - 1$$

$$rate$$

+ $future\_value = 0$

Example

In this example, PAYMENT computes the periodic payment of a 25-year $100,000 loan with an annual interest rate of 8%. The payment is made at the end of each period.

```
PRINT, PAYMENT(0.08, 25, 100000., 0., 0)
-9367.88
```
**PRESENT_VALUE Function**

Evaluates the net present value of a stream of equal periodic cash flows, which are subject to a given discount rate.

**Usage**

\[ result = \text{PRESENT\_VALUE} (rate, n\_periods, payment, future\_value, when) \]

**Input Parameters**

- **rate** — Interest rate.
- **n\_periods** — Total number of periods.
- **payment** — Payment made in each period.
- **future\_value** — The value, at some time in the future, of a current amount and a stream of payments.
- **when** — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.

**Returned Value**

- **result** — The present value of an investment. If no result can be computed, NaN is returned.

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.
Discussion

Function PRESENT_VALUE computes the present value of an investment.

If \( rate = 0 \)

\[
\text{present\_value} + (\text{payment}) (n\_periods) + \text{future\_value} = 0
\]

If \( rate \neq 0 \)

\[
\text{present\_value}(1+rate)^{n\_periods} + \text{payment}[1+rate(\text{when})] \frac{(1+rate)^{n\_periods} - 1}{rate} + \text{future\_value} = 0
\]

Example

In this example, PRESENT_VALUE computes the present value of 20 payments of $500,000 per payment ($10 million) with an annual interest rate of 6%. The payment is made at the end of each period.

\[
\text{PRINT, PRESENT\_VALUE(0.06, 20, 500000., 0., 0)}
\]

\[-5.73496e+06\]

\textbf{PRES\_VAL\_SCHD Function}

Evaluates the present value for a schedule of cash flows. It is not necessary that the cash flows be periodic.

\textbf{Usage}

\[
\text{result} = \text{PRES\_VAL\_SCHD} (\text{rate}, \text{values}, \text{dates})
\]

\textbf{Input Parameters}

\textit{rate} — Interest rate.

\textit{values} — One-dimensional array of cash flows.
**dates** — One-dimensional array of dates cash flows are made. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

**Returned Value**

**result** — The present value for a schedule of cash flows that is not necessarily periodic. If no result can be computed, NaN is returned.

**Input Keywords**

**Double** — If present and nonzero, double precision is used.

**Discussion**

Function PRES_VAL_SCHD computes the present value for a schedule of cash flows that is not necessarily periodic.

It can be found by solving the following with \( count = \text{N\_ELEMENTS}(values) \):

\[
\sum_{i=1}^{count} \frac{value_i}{(1 + rate)^{d_i - d_1/365}}
\]

In the equation above, \( d_i \) represents the \( i \)th payment date, \( d_1 \) represents the 1st payment date, and \( value_i \) represents the \( i \)th cash flow.

**Example**

In this example, PRES_VAL_SCHD computes the present value of 3 payments, $1,000, $2,000 and $1,000, with an interest rate of 5% made on January 3, 1997, January 3, 1999 and January 3, 2000.

\[
\begin{align*}
rate &= 0.05 \\
values &= [1000.0, 2000.0, 1000.0] \\
dates &= \text{VAR\_TO\_DT}([1997, 1999, 2000], [1, 1, 1], [3, 3, 3])
\end{align*}
\]

PRINT, PRES_VAL_SCHD(rate, values, dates)

\[
3677.90
\]
**PRINC PAYMENT Function**

Evaluates the payment on the principal for a specified period.

**Usage**

\[
\text{result} = \text{PRINC PAYMENT} (\text{rate} , \text{period} , \text{n_periods} , \text{present_value} , \text{future_value} , \text{when})
\]

**Input Parameters**

- **rate** — Interest rate.
- **period** — Payment period.
- **n_periods** — Total number of periods.
- **present_value** — The current value of a stream of future payments, after discounting the payments using some interest rate.
- **future_value** — The value, at some time in the future, of a current amount and a stream of payments.
- **when** — Time in each period when the payment is made, either 0 for at the end of period or 1 for at the beginning of period.

**Returned Value**

\[
\text{result} — \text{The payment on the principal for a given period. If no result can be computed, NaN is returned.}
\]

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.

**Discussion**

Function PRINC PAYMENT computes the payment on the principal for a given period.

It is computed using the following:

\[
\text{payment}_i - \text{interest}_i
\]
where $payment_i$ is computed from the function PAYMENT for the $i$th period, $interest_i$ is calculated from the function INT_PAYMENT for the $i$th period.

**Example**

In this example, PRINC_PAYMENT computes the principal paid for the first year on a 30-year $100,000 loan with an annual interest rate of 8%. The payment is made at the end of each year.

```
PRINT, PRINC_PAYMENT(0.08, 1, 30, 100000., 0., 0)
-882.742
```

**ACCR_INT_MAT Function**

Evaluator the interest which has accrued on a security that pays interest at maturity.

**Usage**

```
result = ACCR_INT_MAT (issue, maturity, coupon_rate, par_value, basis)
```

**Input Parameters**

- **issue** — The date on which interest starts accruing. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.
- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.
- **coupon_rate** — Annual interest rate set forth on the face of the security; the coupon rate.
- **par_value** — Nominal or face value of the security used to calculate interest payments.
- **basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.
**Returned Value**

**result** — The interest which has accrued on a security that pays interest at maturity. If no result can be computed, NaN is returned.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.

**Discussion**

Function ACCR_INT_MAT computes the accrued interest for a security that pays interest at maturity:

\[
(par\_value)(rate)\left(\frac{A}{D}\right)
\]

In the above equation, \(A\) represents the number of days starting at issue date to maturity date and \(D\) represents the annual basis.

**Example**

In this example, ACCR_INT_MAT computes the accrued interest for a security that pays interest at maturity using the US (NASD) 30/360 day count method. The security has a par value of $1,000, the issue date of October 1, 2000, the maturity date of November 3, 2000, and a coupon rate of 6%.

<table>
<thead>
<tr>
<th>basis</th>
<th>PDay count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>
issue = VAR_TO_DT(2000, 10, 1)
maturity = VAR_TO_DT(2000, 11, 3)
rate = .06
par = 1000.
basis = 1
PRINT, ACCR_INT_MAT(issue, maturity, rate, par, basis)
      5.33333

**ACCR_INT_PER Function**

Evaluates the interest which has accrued on a security that pays interest periodically.

**Usage**

\[
result = \text{ACCR_INT_PER}(\text{issue}, \text{first_coupon}, \text{settlement}, \text{coupon_rate}, \\
\text{par_value}, \text{frequency}, \text{basis})
\]

**Input Parameters**

- **issue** — The date on which interest starts accruing. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **first_coupon** — First date on which an interest payment is due on the security (e.g. coupon date). For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **coupon_rate** — Annual interest rate set forth on the face of the security; the coupon rate.

- **par_value** — Nominal or face value of the security used to calculate interest payments.

- **frequency** — Frequency of the interest payments. It should be either 1, 2 or 4.
The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>frequency</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>One payment per year (Annual payment)</td>
</tr>
<tr>
<td>2</td>
<td>Two payments per year (Semi-annual payment)</td>
</tr>
<tr>
<td>4</td>
<td>Four payments per year (Quarterly payment)</td>
</tr>
</tbody>
</table>

*basis* — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>basis</th>
<th>PDay count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

Returned Value

*result* — The accrued interest for a security that pays periodic interest. If no result can be computed, NaN is returned.

Input Keywords

*Double* — If present and nonzero, double precision is used.

Discussion

Function ACCR_INT_PER computes the accrued interest for a security that pays periodic interest.

In the equation below, $A_i$ represents the number days which have accrued for the $i$th quasi-coupon period within the odd period. (The quasi-coupon periods are periods obtained by extending the series of equal payment periods to before or after the actual payment periods.) $NC$ represents the number of quasi-coupon
periods within the odd period, rounded to the next highest integer. (The odd period is a period between payments that differs from the usual equally spaced periods at which payments are made.) \( NL_i \) represents the length of the normal \( i \)th quasi-coupon period within the odd period. \( NL_i \) is expressed in days.

Function ACCR_INT_PER can be found by solving the following:

\[
\left( \frac{par\_value}{frequency} \right) \left( \frac{\sum_{i=1}^{NC} \left( \frac{A_i}{NL_i} \right)}{rate} \right)
\]

**Example**

In this example, ACCR_INT_PER computes the accrued interest for a security that pays periodic interest using the US (NASD) 30/360 day count method. The security has a par value of $1,000, the issue date of October 1, 1999, the settlement date of November 3, 1999, the first coupon date of March 31, 2000, and a coupon rate of 6%.

```plaintext
issue = VAR_TO_DT(1999, 10, 1)
first_coupon = VAR_TO_DT(2000, 3, 31)
settlement = VAR_TO_DT(1999, 11, 3)
rate = .06
par = 1000.
frequency = 2
basis = 1
PRINT, ACCR_INT_PER(issue, first_coupon, settlement, $
rate, par, frequency, basis)
 5.33333
```

**BOND_EQV_YIELD Function**

Evaluates the bond-equivalent yield of a Treasury bill.

**Usage**

\[
result = \text{BOND_EQV_YIELD} (settlement, maturity, discount\_rate)
\]
Input Parameters

settlement — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.

maturity — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.

discount_rate — The interest rate implied when a security is sold for less than its value at maturity in lieu of interest payments.

Returned Value

result — The bond-equivalent yield of a Treasury bill. If no result can be computed, NaN is returned.

Input Keywords

Double — If present and nonzero, double precision is used.

Discussion

Function BOND_EQV_YIELD computes the bond-equivalent yield for a Treasury bill.

It is computed using the following:

\[ \text{if } DSM \leq 182 \]

\[ \frac{365 \times \text{discount_rate}}{360 - \text{discount_rate} \times DSM} \]

otherwise,

\[ -\frac{DSM}{365} + \sqrt{\left(\frac{DSM}{365}\right)^2 - \left(\frac{2 \times DSM}{365} - 1\right) \times \frac{\text{discount_rate} \times DSM}{\text{discount_rate} \times DSM - 360}} \frac{DSM}{365} - 0.5 \]

In the above equation, DSM represents the number of days starting at settlement date to maturity date.
Example
In this example, BOND_EQV_YIELD computes the bond-equivalent yield for a Treasury bill with the settlement date of July 1, 1999, the maturity date of July 1, 2000, and discount rate of 5% at the issue date.
PRINT, BOND_EQV_YIELD(settlement, maturity, discount)
0.052857

CONVEXITY Function
Evaluates the convexity for a security.

Usage
result = CONVEXITY (settlement, maturity, coupon_rate, yield, frequency, basis)

Input Parameters
settlement — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.
maturity — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.
coupon_rate — Annual interest rate set forth on the face of the security; the coupon rate.
yield — Annual yield of the security.
frequency — Frequency of the interest payments. It should be either 1, 2 or 4.
The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

*basis* — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

### Input Keywords

*Double* — If present and nonzero, double precision is used.

### Discussion

Function CONVEXITY computes the convexity for a security. Convexity is the sensitivity of the duration of a security to changes in yield.
It is computed using the following:

\[
\frac{1}{(q^{*} \text{ frequency})^{2}} \left\{ \sum_{t=1}^{n} t(t+1) \left( \frac{\text{coupon\_rate}}{\text{frequency}} q^{-t} + n(n+1)q^{-n} \right) \right. \\
\left. \sum_{t=1}^{n} \left( \frac{\text{coupon\_rate}}{\text{frequency}} q^{-t} + q^{-n} \right) \right\}
\]

where \( n \) is calculated from the function COUPON_NUM and

\[
q = 1 + \frac{\text{yield}}{\text{frequency}}.
\]

**Example**

In this example, CONVEXITY computes the convexity for a security with the settlement date of July 1, 1990, and maturity date of July 1, 2000, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(1990, 7, 1)  
maturity = VAR_TO_DT(2000, 7, 1)  
coupon = .075  
yield = .09  
frequency = 2  
basis = 3

PRINT, CONVEXITY(settlement, maturity, $  
coupon, yield, frequency, basis)  
59.4050
```

**COUPON_DAYS Function**

Evaluates the number of days in the coupon period containing the settlement date.

**Usage**

\[
result = \text{COUPON\_DAYS} (\text{settlement}, \text{maturity}, \text{frequency}, \text{basis})
\]
Input Parameters

settlement — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.

maturity — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.

frequency — Frequency of the interest payments. It should be either 1, 2 or 4.

<table>
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<tr>
<th>frequency</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>One payment per year</td>
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<td></td>
<td>(Annual payment)</td>
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<td>Four payments per year</td>
</tr>
<tr>
<td></td>
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</table>

basis — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

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<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

Returned Value

result — The number of days in the coupon period which contains the settlement date. If no result can be computed, NaN is returned.
Input Keywords

*Double* — If present and nonzero, double precision is used.

Discussion

Function COUPON_DAYS computes the number of days in the coupon period that contains the settlement date. For a good discussion on day count basis, see *SIA Standard Securities Calculation Methods* 1993, vol. 1, pages 17-35.

Example

In this example, COUPON_DAYS computes the number of days in the coupon period of a bond with the settlement date of November 11, 1996, and the maturity date of March 1, 2009, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(1996, 11, 11)
maturity = VAR_TO_DT(2009, 3, 1)
frequency = 2
basis = 3
PRINT, COUPON_DAYS(settlement, maturity, frequency, basis)
182.500
```

**COUPON_NUM Function**

Evaluates the number of coupons payable between the settlement date and the maturity date.

Usage

```
result = COUPON_NUM (settlement, maturity, frequency, basis)
```

Input Parameters

*settlement* — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*maturity* — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.
**frequency** — Frequency of the interest payments. It should be either 1, 2 or 4.

**basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

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</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

**Returned Value**

**result** — The number of coupons payable between the settlement date and the maturity date.

**Input Keywords**

**Double** — If present and nonzero, double precision is used.

**Discussion**

Function COUPON_NUM computes the number of coupons payable between the settlement date and the maturity date. For a good discussion on day count basis, see *SIA Standard Securities Calculation Methods 1993*, vol. 1, pages 17-35.

**Example**

In this example, COUPON_NUM computes the number of coupons payable with the settlement date of November 11, 1996, and the maturity date of March 1, 2009, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(1996, 11, 11)
maturity = VAR_TO_DT(2009, 3, 1)
frequency = 2
basis = 3
```
PRINT, COUPON_NUM(settlement, maturity, frequency, basis)

**SETTLEMENT_DB Function**

Evaluates the number of days starting with the beginning of the coupon period and ending with the settlement date.

**Usage**

\[ \text{result} = \text{SETTLEMENT_DB (settlement, maturity, frequency, basis)} \]

**Input Parameters**

*settlement* — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*maturity* — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*frequency* — Frequency of the interest payments. It should be either 1, 2 or 4.

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<td>Four payments per year</td>
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<tr>
<td></td>
<td>(Quarterly payment)</td>
</tr>
</tbody>
</table>

*basis* — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

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144 Chapter 2: New Commands

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<thead>
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<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

Returned Value

`result` — The number of days in the period starting with the beginning of the coupon period and ending with the settlement date.

Input Keywords

`double` — If present and nonzero, double precision is used.

Discussion

Function SETTLEMENT_DB computes the number of days from the beginning of the coupon period to the settlement date. For a good discussion on day count basis, see *SIA Standard Securities Calculation Methods* 1993, vol. 1, pages 17-35.

Example

In this example, SETTLEMENT_DB computes the number of days from the beginning of the coupon period to November 11, 1996, of a bond with the maturity date of March 1, 2009, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(1996, 11, 11)
maturity = VAR_TO_DT(2009, 3, 1)
frequency = 2
basis = 3
PRINT, SETTLEMENT_DB(settlement, maturity, frequency, basis)
71
```
**COUPON_DNC Function**

Evaluates the number of days starting with the settlement date and ending with the next coupon date.

**Usage**

\[
result = \text{COUPON}_D\text{NC} \left( \text{settlement}, \text{maturity}, \text{frequency}, \text{basis} \right)
\]

**Input Parameters**

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **frequency** — Frequency of the interest payments. It should be either 1, 2 or 4.

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</tr>
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<td>(Quarterly payment)</td>
</tr>
</tbody>
</table>

- **basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

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<tr>
<td>2</td>
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</tr>
</tbody>
</table>
Returned Value

\textit{result} — The number of days starting with the settlement date and ending with the next coupon date.

Input Keywords

\textit{Double} — If present and nonzero, double precision is used.

Discussion

Function COUPON_DNC computes the number of days from the settlement date to the next coupon date. For a good discussion on day count basis, see \textit{SIA Standard Securities Calculation Methods} 1993, vol. 1, pp. 17-35.

Example

In this example, COUPON_DNC computes the number of days from November 11, 1996, to the next coupon date of a bond with the maturity date of March 1, 2009, using the Actual/365 day count method.

\begin{verbatim}
settlement = VAR_TO_DT(1996, 11, 11)
maturity = VAR_TO_DT(2009, 3, 1)
frequency = 2
basis = 3
PRINT, COUPON_DNC(settlement, maturity, frequency, basis)
\end{verbatim}

\begin{verbatim}
110
\end{verbatim}
DEPREC_AMORDEGRC Function

Evaluates the depreciation for each accounting period. During the evaluation of the function a depreciation coefficient based on the asset life is applied.

Usage

\[
result = \text{DEPREC\_AMORDEGRC}(\text{cost}, \text{issue}, \text{first\_period}, \text{salvage}, \text{period}, \text{rate}, \text{basis})
\]

Input Parameters

- **cost**— Initial value of the asset.
- **issue**— The date on which interest starts accruing. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.
- **first\_period**— Date of the end of the first period. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.
- **salvage** — The value of an asset at the end of its depreciation period.
- **period** — Depreciation for the accounting period to be computed.
- **rate** — Depreciation rate.
- **basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

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<tr>
<td>4</td>
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</tr>
</tbody>
</table>
Returned Value

.result — The depreciation for each accounting period. If no result can be computed, NaN is returned.

Input Keywords

Double — If present and nonzero, double precision is used.

Discussion

Function DEPREC_AMORDEGRC computes the depreciation for each accounting period. This function is similar to DEPREC_AMORLINC. However, in this function a depreciation coefficient based on the asset life is applied during the evaluation of the function.

Example

In this example, DEPREC_AMORDEGRC computes the depreciation for the second accounting period using the US (NASD) 30/360 day count method. The security has the issue date of November 1, 1999, end of first period of November 30, 2000, cost of $2,400, salvage value of $300, depreciation rate of 15%.

```plaintext
issue = VAR_TO_DT(1999, 11, 1)
first_period = VAR_TO_DT(2000, 11, 30)
cost = 2400.
salvage = 300.
period = 2
rate = .15
basis = 1
PRINT, DEPREC_AMORDEGRC(cost, issue, first_period, $
salvage, period, rate, basis
335.000
```

DEPREC_AMORLINC Function

Evaluates the depreciation for each accounting period. This function is similar to DEPREC_AMORDEGRC, except that DEPREC_AMORDEGRC has a
depreciation coefficient that is applied during the evaluation that is based on the asset life.

**Usage**

\[
result = \text{DEPREC\_AMORLINC} (cost, issue, first\_period, salvage, period, rate, basis)
\]

**Input Parameters**

- **cost** — Initial value of the asset.
- **issue** — The date on which interest starts accruing. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.
- **first\_period** — Date of the end of the first period. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.
- **salvage** — The value of an asset at the end of its depreciation period.
- **period** — Depreciation for the accounting period to be computed.
- **rate** — Depreciation rate.
- **basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

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<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

**Returned Value**

- **result** — The depreciation for each accounting period. If no result can be computed, NaN is returned.
Input Keywords

*Double* — If present and nonzero, double precision is used.

Discussion

Function DEPREC_AMORLINC computes the depreciation for each accounting period.

Example

In this example, DEPREC_AMORLINC computes the depreciation for the second accounting period using the US (NASD) 30/360 day count method. The security has the issue date of November 1, 1999, end of first period of November 30, 2000, cost of $2,400, salvage value of $300, depreciation rate of 15%.

```plaintext
issue = VAR_TO_DT(1999, 11, 1)
first_period = VAR_TO_DT(2000, 11, 30)
cost = 2400.
salvage = 300.
period = 2
rate = .15
basis = 1
PRINT, DEPREC_AMORLINC(cost, issue, first_period, $
    salvage, period, rate, basis)
360.000
```

**DISCOUNT_PR Function**

Evaluates the price of a security sold for less than its face value.

Usage

```plaintext
result = DISCOUNT_PR (settlement, maturity, discount_rate, redemption, $
    basis)
```
Input Parameters

**settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

**maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

**discount_rate** — The interest rate implied when a security is sold for less than its value at maturity in lieu of interest payments.

**redemption** — Redemption value per $100 face value of the security.

**basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
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<td>European 30/360</td>
</tr>
</tbody>
</table>

Returned Value

**result** — The price per face value for a discounted security. If no result can be computed, NaN is returned.

Input Keywords

**Double** — If present and nonzero, double precision is used.

Discussion

Function DISCOUNT_PR computes the price per $100 face value of a discounted security.
It is computed using the following:

\[
\text{redemption} = \left( \frac{\text{discount rate}}{\text{redemption}} \right) \left( \frac{\text{DSM}}{\text{B}} \right)
\]

In the equation above, DSM represents the number of days starting at the settlement date and ending with the maturity date. B represents the number of days in a year based on the annual basis.

**Example**

In this example, DISCOUNT_PR computes the price of the discounted bond with the settlement date of July 1, 2000, and maturity date of July 1, 2001, at the discount rate of 5% using the US (NASD) 30/360 day count method.

```plaintext
settlement = VAR_TO_DT(2000, 7, 1)
maturity = VAR_TO_DT(2001, 7, 1)
discount = .05
redemption = 100.
basis = 1
PRINT, DISCOUNT_PR(settlement, maturity, discount, $
redemption, basis
95.0000
```

**DISCOUNT_RT Function**

Evaluates the interest rate implied when a security is sold for less than its value at maturity in lieu of interest payments.

**Usage**

\[
\text{result} = \text{DISCOUNT_RT}(\text{settlement, maturity, price, redemption, basis})
\]

**Input Parameters**

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.
**maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

**price** — Price per $100 face value of the security.

**redemption** — Redemption value per $100 face value of the security.

**basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

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<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

**Returned Value**

**result** — The discount rate for a security. If no result can be computed, NaN is returned.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.

**Discussion**

Function DISCOUNT_RT computes the discount rate for a security. The discount rate is the interest rate implied when a security is sold for less than its value at maturity in lieu of interest payments.

It is computed using the following:

\[
\left( \frac{\text{redemption} - \text{price}}{\text{price}} \right) \times \frac{B}{\text{DSM}}
\]
In the equation above, $B$ represents the number of days in a year based on the annual basis and $DSM$ represents the number of days starting with the settlement date and ending with the maturity date.

**Example**

In this example, `DISCOUNT_RT` computes the discount rate of a security which is selling at $97.975 with the settlement date of February 15, 2000, and maturity date of June 10, 2000, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(2000, 2, 15)
maturity = VAR_TO_DT(2000, 6, 10)
price = 97.975
redemption = 100.
basis = 3
PRINT, DISCOUNT_RT(settlement, maturity, price, redemption, basis)
0.0637177
```

**DISCOUNT_YLD Function**

Evaluates the annual yield of a discounted security.

**Usage**

```
result = DISCOUNT_YLD (settlement, maturity, price, redemption, basis)
```

**Input Parameters**

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **price** — Price per $100 face value of the security.

- **redemption** — Redemption value per $100 face value of the security.
**basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

**Returned Value**

**result** — The annual yield for a discounted security. If no result can be computed, NaN is returned.

**Input Keywords**

**Double** — If present and nonzero, double precision is used.

**Discussion**

Function DISCOUNT_YLD computes the annual yield for a discounted security.

It is computed using the following:

\[
\left( \frac{\text{redemption} - \text{price}}{\text{price}} \right) \left( \frac{B}{DSM} \right)
\]

In the equation above, \( B \) represents the number of days in a year based on the annual basis, and \( DSM \) represents the number of days starting with the settlement date and ending with the maturity date.

**Example**

In this example, DISCOUNT_YLD computes the annual yield for a discounted security which is selling at $95.40663 with the settlement date of July 1, 1995,
and maturity date of July 1, 2005, using the US (NASD) 30/360 day count method.

```plaintext
calendar = VAR_TO_DT(1995, 7, 1)
maturity = VAR_TO_DT(2005, 7, 1)
price = 95.40663
redemption = 105.
basis = 1
PRINT, DISCOUNT_YLD(settlement, maturity, price, redemption$, basis)
0.0100552
```

---

**DURATION Function**

Evaluates the annual duration of a security where the security has periodic interest payments.

**Usage**

```plaintext
result = DURATION (settlement, maturity, coupon_rate, yield, frequency, basis)
```

**Input Parameters**

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **coupon_rate** — Annual interest rate set forth on the face of the security; the coupon rate.

- **yield** — Annual yield of the security.

- **frequency** — Frequency of the interest payments. It should be either 1, 2 or 4.
### DURATION Function

The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

**Returned Value**

*result* — The annual duration of a security with periodic interest payments. If no result can be computed, NaN is returned.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.

**Discussion**

Function DURATION computes the Macaulay’s duration of a security with periodic interest payments. The Macaulay’s duration is the weighted-average time to the payments, where the weights are the present value of the payments.

<table>
<thead>
<tr>
<th>frequency</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>One payment per year (Annual payment)</td>
</tr>
<tr>
<td>2</td>
<td>Two payments per year (Semi-annual payment)</td>
</tr>
<tr>
<td>4</td>
<td>Four payments per year (Quarterly payment)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
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</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
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</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

*basis* — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.
It is computed using the following:

\[
\frac{DSC}{E}^{100} \frac{1}{1 + \frac{yield}{freq}}^{N-1 + \frac{DSC}{E}} + \sum_{k=1}^{N} \left( \frac{100 \cdot \text{coupon}_\text{rate}}{freq} \left( 1 + \frac{yield}{freq} \right)^{k-1 + \frac{DSC}{E}} \right) \times \left( k - 1 + \frac{DSC}{E} \right) \times \frac{1}{freq}
\]

In the equation above, \(DSC\) represents the number of days starting with the settlement date and ending with the next coupon date. \(E\) represents the number of days within the coupon period. \(N\) represents the number of coupons payable from the settlement date to the maturity date. \(freq\) represents the frequency of the coupon payments annually.

**Example**

In this example, DURATION computes the annual duration of a security with the settlement date of July 1, 1995, and maturity date of July 1, 2005, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(1995, 7, 1)
maturity = VAR_TO_DT(2005, 7, 1)
coupon = .075
yield = .09
frequency = 2
basis = 3
PRINT, DURATION(settlement, maturity, coupon, $
                   yield, frequency, basis)
```

```
7.04195
```
**INT_RATE_SEC Function**

Evaluates the interest rate of a fully invested security.

**Usage**

\[
result = \text{INT RATE SEC} (\text{settlement, maturity, investment, redemption, basis})
\]

**Input Parameters**

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **investment** — The total amount one has invested in the security.

- **redemption** — Amount to be received at maturity.

- **basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
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</tr>
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<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

**Returned Value**

- **result** — The interest rate for a fully invested security. If no result can be computed, NaN is returned.
Input Keywords

**Double** — If present and nonzero, double precision is used.

Discussion

Function INT_RATE_SEC computes the interest rate for a fully invested security.

It is computed using the following:

\[
\left( \frac{\text{redemption} - \text{investment}}{\text{investment}} \right) \left( \frac{B}{\text{DSM}} \right)
\]

In the equation above, \( B \) represents the number of days in a year based on the annual basis, and \( DSM \) represents the number of days in the period starting with the settlement date and ending with the maturity date.

Example

In this example, INT_RATE_SEC computes the interest rate of a $7,000 investment with the settlement date of July 1, 1995, and maturity date of July 1, 2005, using the Actual/365 day count method. The total amount received at the end of the investment is $10,000.

```plaintext
settlement = VAR_TO_DT(1995, 7, 1)
maturity = VAR_TO_DT(2005, 7, 1)
investment = 7000.
redemption = 10000.
basis = 3
PRINT, INT_RATE_SEC(settlement, maturity, investment,$
    redemption, basis)
0.0428219
```
**DURATION_MAC Function**

Evaluates the modified Macauley duration of a security.

**Usage**

\[
result = \text{DURATION_MAC} (\text{settlement}, \text{maturity}, \text{coupon\_rate}, \text{yield}, \\
\text{frequency}, \text{basis})
\]

**Input Parameters**

*settlement* — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*maturity* — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*coupon\_rate* — Annual interest rate set forth on the face of the security; the coupon rate.

*yield* — Annual yield of the security.

*frequency* — Frequency of the interest payments. It should be either 1, 2 or 4.

<table>
<thead>
<tr>
<th>frequency</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>One payment per year (Annual payment)</td>
</tr>
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<td>Two payments per year (Semi-annual payment)</td>
</tr>
<tr>
<td>4</td>
<td>Four payments per year (Quarterly payment)</td>
</tr>
</tbody>
</table>

*basis* — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
</tbody>
</table>
Returned Value

**result** — The modified Macauley duration of a security is returned. The security has an assumed par value of $100. If no result can be computed, NaN is returned.

Input Keywords

**Double** — If present and nonzero, double precision is used.

Discussion

Function DURATION_MAC computes the modified Macauley duration for a security with an assumed par value of $100.

It is computed using the following:

\[
\frac{duration}{yield} = \frac{1}{frequency}
\]

where \( duration \) is calculated from the function DURATION.

Example

In this example, DURATION_MAC computes the modified Macauley duration of a security with the settlement date of July 1, 1995, and maturity date of July 1, 2005, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(1995, 7, 1)
maturity = VAR_TO_DT(2005, 7, 1)
coupon = 0.075
yield = 0.09
```

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
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<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>
frequency = 2
basis = 3
PRINT, DURATION_MAC(settlement, maturity, $
coupon, yield, frequency, basis)
    6.73871

**COUPON_NCD Function**

Evaluates the first coupon date which follows the settlement date.

**Usage**

\[
result = \text{COUPON\_NCD} (settlement, maturity, frequency, basis)
\]

**Input Parameters**

*settlement* — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*maturity* — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*frequency* — Frequency of the interest payments. It should be either 1, 2 or 4.

<table>
<thead>
<tr>
<th>frequency</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>One payment per year</td>
</tr>
<tr>
<td></td>
<td>(Annual payment)</td>
</tr>
<tr>
<td>2</td>
<td>Two payments per year</td>
</tr>
<tr>
<td></td>
<td>(Semi-annual payment)</td>
</tr>
<tr>
<td>4</td>
<td>Four payments per year</td>
</tr>
<tr>
<td></td>
<td>(Quarterly payment)</td>
</tr>
</tbody>
</table>

*basis* — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.
 Returned Value

*result* — The first coupon date which follows the settlement date.

 Input Keywords

*Double* — If present and nonzero, double precision is used.

 Discussion

Function COUPON_NCD computes the next coupon date after the settlement date. For a good discussion on day count basis, see *SIA Standard Securities Calculation Methods* 1993, vol 1, pages 17-35.

 Example

In this example, COUPON_NCD computes the next coupon date of a bond with the settlement date of November 11, 1996, and the maturity date of March 1, 2009, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(1996, 11, 11)
maturity = VAR_TO_DT(2009, 3, 1)
frequency = 2
basis = 3
ans = COUPON_NCD(settlement, maturity, frequency, basis)
DT_TO_STR, ans, d, DateFmt=4
01/March/1997
```

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
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<td>Actual/360</td>
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<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>
**COUPON_PCD Function**

Evaluates the coupon date which immediately precedes the settlement date.

**Usage**

\[ \text{result} = \text{COUPON}_P\text{CD (settlement, maturity, frequency, basis)} \]

**Input Parameters**

*settlement* — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*maturity* — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

*frequency* — Frequency of the interest payments. It should be either 1, 2 or 4.

<table>
<thead>
<tr>
<th>frequency</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>One payment per year</td>
</tr>
<tr>
<td></td>
<td>(Annual payment)</td>
</tr>
<tr>
<td>2</td>
<td>Two payments per year</td>
</tr>
<tr>
<td></td>
<td>(Semi-annual payment)</td>
</tr>
<tr>
<td>4</td>
<td>Four payments per year</td>
</tr>
<tr>
<td></td>
<td>(Quarterly payment)</td>
</tr>
</tbody>
</table>

*basis* — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
</tbody>
</table>
 Returned Value

result — The coupon date which immediately precedes the settlement date.

 Input Keywords

Double — If present and nonzero, double precision is used.

 Discussion

Function COUPON_PCD computes the coupon date which immediately precedes the settlement date. For a good discussion on day count basis, see *SIA Standard Securities Calculation Methods* 1993, vol 1, pages 17-35.

 Example

In this example, COUPON_PCD computes the previous coupon date of a bond with the settlement date of November 11, 1986, and the maturity date of March 1, 1999, using the Actual/365 day count method.

```matlab
settlement = VAR_TO_DT(1996, 11, 11)
maturity = VAR_TO_DT(2009, 3, 1)
frequency = 2
basis = 3
ans = COUPON_PCD(settlement, maturity, frequency, basis)
DT_TO_STR, ans, d, DateFmt=4
PRINT, d
01/September/1996
```

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>
**PRICE_PERIODIC Function**

Evaluates the price, per $100 face value, of a security that pays periodic interest.

**Usage**

\[
\text{result} = \text{PRICE\_PERIODIC}(\text{settlement}, \text{maturity}, \text{rate}, \text{yield}, \text{redemption}, \text{frequency}, \text{basis})
\]

**Input Parameters**

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid.

- **rate** — Annual interest rate set forth on the face of the security; the coupon rate.

- **yield** — Annual yield of the security.

- **redemption** — Redemption value per $100 face value of the security.

- **frequency** — Frequency of the interest payments. It should be either 1, 2 or 4.

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
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<td></td>
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</tr>
<tr>
<td>4</td>
<td>Four payments per year</td>
</tr>
<tr>
<td></td>
<td>(Quarterly payment)</td>
</tr>
</tbody>
</table>

- **basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.
Returned Value

**result** — The price per $100 face value of a security that pays periodic interest. If no result can be computed, NaN is returned.

Input Keywords

**Double** — If present and nonzero, double precision is used.

Discussion

Function PRICE_PERIODIC computes the price per $100 face value of a security that pays periodic interest.

It is computed using the following:

\[
\frac{\text{redemption}}{1 + \frac{\text{yield}}{\text{frequency}}} \left( \sum_{k=1}^{N} \frac{100^a \text{rate}}{\text{frequency}} \left( \frac{DSC}{E} \right) \right) - \frac{100^a \text{rate}}{\text{frequency}} \left( \frac{A}{E} \right)
\]

In the above equation, \(DSC\) represents the number of days in the period starting with the settlement date and ending with the next coupon date. \(E\) represents the number of days within the coupon period. \(N\) represents the number of coupons payable in the timeframe from the settlement date to the redemption date. \(A\) represents the number of days in the timeframe starting with the beginning of coupon period and ending with the settlement date.
Example

In this example, PRICE_PERIODIC computes the price of a bond that pays coupon every six months with the settlement of July 1, 1995, the maturity date of July 1, 2005, a annual rate of 6%, annual yield of 7% and redemption value of $105 using the US (NASD) 30/360 day count method.

```plaintext
settlement = VAR_TO_DT(1995, 7, 1)
maturity = VAR_TO_DT(2005, 7, 1)
rate = .06
yield = .07
redemption = 105.
frequency = 2
basis = 1
PRINT, PRICE_PERIODIC(settlement, maturity, rate, yield, $redemption, frequency, basis)

95.4067
```

**PRICE_MATURITY Function**

Evaluates the price, per $100 face value, of a security that pays interest at maturity.

Usage

```plaintext
result = PRICE_MATURITY (settlement, maturity, issue, rate, yield, basis)
```

Input Parameters

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **issue** — The date on which interest starts accruing. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.
rate — Annual interest rate set forth on the face of the security; the coupon rate.

yield — Annual yield of the security.

basis — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
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<th>Day count basis</th>
</tr>
</thead>
<tbody>
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<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

Returned Value

result — The price per $100 face value of a security that pays interest at maturity. If no result can be computed, NaN is returned.

Input Keywords

Double — If present and nonzero, double precision is used.

Discussion

Function PRICE_MATURITY computes the price per $100 face value of a security that pays interest at maturity.

It is computed using the following:

\[
\left[\frac{100 + \left(\frac{DIM}{B} \cdot rate \cdot 100\right)}{1 + \left(\frac{DSM}{B} \cdot yield\right)}\right] \cdot \left(\frac{A}{B} \cdot rate \cdot 100\right)
\]

In the equation above, \(B\) represents the number of days in a year based on the annual basis. \(DSM\) represents the number of days in the period starting with the settlement date and ending with the maturity date. \(DIM\) represents the number
of days in the period starting with the issue date and ending with the maturity date. \( A \) represents the number of days in the period starting with the issue date and ending with the settlement date.

**Example**

In this example, \texttt{PRICE_MATURITY} computes the price at maturity of a security with the settlement date of August 1, 2000, maturity date of July 1, 2001 and issue date of July 1, 2000, using the US (NASD) 30/360 day count method. The security has 5\% annual yield and 5\% interest rate at the date of issue.

```plaintext
settlement = VAR_TO_DT(2000, 8, 1)
maturity = VAR_TO_DT(2001, 7, 1)
issue = VAR_TO_DT(2000, 7, 1)
rate = .05
yield = .05
basis = 1
PRINT, PRICE_MATURITY(settlement, maturity, issue, $
                      rate, yield, basis)

         99.9817
```

**MATURITY_REC Function**

Evaluates the amount one receives when a fully invested security reaches the maturity date.

**Usage**

```plaintext
result = MATURITY_REC(settlement, maturity, investment, discount_rate, basis)
```

**Input Parameters**

- \texttt{settlement} — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, \textit{Working with Date/Time Data} in the PV-WAVE User’s Guide.

- \texttt{maturity} — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, \textit{Working with Date/Time Data} in the PV-WAVE User’s Guide.
**investment** — The total amount one has invested in the security.

**discount_rate** — The interest rate implied when a security is sold for less than its value at maturity in lieu of interest payments.

**basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

**Returned Value**

**result** — The amount one receives when a fully invested security reaches its maturity date. If no result can be computed, NaN is returned.

**Input Keywords**

**Double** — If present and nonzero, double precision is used.

**Discussion**

Function MATURITY_REC computes the amount received at maturity for a fully invested security.

It is computed using the following:

\[
\text{result} = \frac{\text{investment} \times \text{DIM}}{B} \times (1 - \left(\frac{\text{discount_rate} \times \text{DIM}}{B}\right))
\]

In the equation above, \(B\) represents the number of days in a year based on the annual basis, and \(\text{DIM}\) represents the number of days in the period starting with the issue date and ending with the maturity date.
Example

In this example, MATURITY_REC computes the amount received of a $7,000 investment with the settlement date of July 1, 1995, maturity date of July 1, 2005 and discount rate of 6%, using the Actual/365 day count method.

```plaintext
settlement = VAR_TO_DT(1995, 7, 1)
maturity = VAR_TO_DT(2005, 7, 1)
investment = 7000.
discount = .06
basis = 3
PRINT, MATURITY_REC(settlement, maturity, investment,$
  discount, basis)

17521.6
```

**TBILL_PRICE Function**

Evaluates the price per $100 face value of a Treasury bill.

**Usage**

```
result = TBILL_PRICE (settlement, maturity, discount_rate)
```

**Input Parameters**

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **discount_rate** — The interest rate implied when a security is sold for less than its value at maturity in lieu of interest payments.

**Returned Value**

- **result** — The price per $100 face value of a Treasury bill. If no result can be computed, NaN is returned.
Input Keywords

Double — If present and nonzero, double precision is used.

Discussion

Function TBILL_PRICE computes the price per $100 face value for a Treasury bill.

It is computed using the following:

\[
100 \left( 1 - \frac{\text{discount}_\text{rate} \times DSM}{360} \right)
\]

In the equation above, DSM represents the number of days in the period starting with the settlement date and ending with the maturity date (any maturity date that is more than one calendar year after the settlement date is excluded).

Example

In this example, TBILL_PRICE computes the price for a Treasury bill with the settlement date of July 1, 2000, the maturity date of July 1, 2001, and a discount rate of 5% at the issue date.

```
settlement = VAR_TO_DT(2000, 7, 1)
maturity = VAR_TO_DT(2001, 7, 1)
discount = .05
PRINT, TBILL_PRICE(settlement, maturity, discount)
```

94.9306

TBILL_YIELD Function

Evaluates the yield of a Treasury bill.

Usage

\[
\text{result} = \text{TBILL\_YIELD} (\text{settlement}, \text{maturity}, \text{price})
\]
Input Parameters

settlement — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.

maturity — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.

price — Price per $100 face value of the Treasury bill.

Returned Value

result — The yield for a Treasury bill. If no result can be computed, NaN is returned.

Input Keywords

Double — If present and nonzero, double precision is used.

Discussion

Function TBILL_YIELD computes the yield for a Treasury bill.

It is computed using the following:

\[
\frac{100 - \frac{price}{DSM}}{\frac{360}{price}}
\]

In the equation above, DSM represents the number of days in the period starting with the settlement date and ending with the maturity date (any maturity date that is more than one calendar year after the settlement date is excluded).

Example

In this example, TBILL_YIELD computes the yield for a Treasury bill with the settlement date of July 1, 2000, the maturity date of July 1, 2001, and priced at $94.93.

settlement = VAR_TO_DT(2000, 7, 1)
maturity = VAR_TO_DT(2001, 7, 1)
price = 94.93
YEAR_FRACTION Function

Evaluates the fraction of a year represented by the number of whole days between two dates.

Usage

\[
\text{result} = \text{YEAR_FRACTION (date\_start, date\_end, basis)}
\]

Input Parameters

date\_start — Initial date. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.

date\_end — Ending date. For a more detailed discussion on dates see Chapter 8, Working with Date/Time Data in the PV-WAVE User’s Guide.

basis — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

Returned Value

result — The fraction of a year represented by the number of whole days between two dates. If no result can be computed, NaN is returned.
Input Keywords

*Double* — If present and nonzero, double precision is used.

Discussion

Function *YEAR_FRACTION* computes the fraction of the year. It is computed using the following:

\[
\frac{A}{D}
\]

where \( A \) = the number of days from *start* to *end*, \( D \) = annual basis.

Example

In this example, *YEAR_FRACTION* computes the year fraction between August 1, 2000, and July 1, 2001, using the US (NASD) 30/360 day count method.

```plaintext
date_start = VAR_TO_DT(2000, 8, 1)
date_end = VAR_TO_DT(2001, 7, 1)
basis = 1
PRINT, YEAR_FRACTION(date_start, date_end, basis)
```

0.916667

**YIELD_MATURITY Function**

Evaluates the annual yield of a security that pays interest at maturity.

Usage

\[\text{result} = \text{YIELD_MATURITY} (\text{settlement}, \text{maturity}, \text{issue}, \text{rate}, \text{price}, \text{basis})\]

Input Parameters

*settlement* — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.
**maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

**issue** — The date on which interest starts accruing. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

**rate** — Interest rate at date of issue of the security.

**price** — Price per $100 face value of the security.

**basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.

<table>
<thead>
<tr>
<th>basis</th>
<th>Day count basis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Actual/Actual</td>
</tr>
<tr>
<td>1</td>
<td>US (NASD) 30/360</td>
</tr>
<tr>
<td>2</td>
<td>Actual/360</td>
</tr>
<tr>
<td>3</td>
<td>Actual/365</td>
</tr>
<tr>
<td>4</td>
<td>European 30/360</td>
</tr>
</tbody>
</table>

**Returned Value**

**result** — The annual yield of a security that pays interest at maturity. If no result can be computed, NaN is returned.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.

**Discussion**

Function YIELD_MATURITY computes the annual yield of a security that pays interest at maturity.
It is computed using the following:

\[
\frac{\left\{1+\left(\frac{DIM}{B} \cdot rate\right)\right\} - \left\{\frac{price}{100} + \left(\frac{A}{B} \cdot rate\right)\right\} - \left\{\frac{B}{DSM}\right\}}{\left\{\frac{price}{100} + \left(\frac{A}{B} \cdot rate\right)\right\}}
\]

In the equation above, \(DIM\) represents the number of days in the period starting with the issue date and ending with the maturity date. \(DSM\) represents the number of days in the period starting with the settlement date and ending with the maturity date. \(A\) represents the number of days in the period starting with the issue date and ending with the settlement date. \(B\) represents the number of days in a year based on the annual basis.

**Example**

In this example, YIELD_MATURITY computes the annual yield of a security that pays interest at maturity which is selling at $95.40663 with the settlement date of August 1, 2000, the issue date of July 1, 2000, the maturity date of July 1, 2010, and the interest rate of 6% at the issue using the US (NASD) 30/360 day count method.

```plaintext
settlement = VAR_TO_DT(2000, 8, 1)
maturity = VAR_TO_DT(2010, 7, 1)
issue = VAR_TO_DT(2000, 7, 1)
rate = .06
price = 95.40663
basis = 1
PRINT, YIELD_MATURITY(settlement, maturity, issue, rate, price, basis)
0.0673905
```
**YIELD_PERIODIC Function**

Evaluates the yield of a security that pays periodic interest.

**Usage**

\[
\text{result} = \text{YIELD\_PERIODIC}(\text{settlement}, \text{maturity}, \text{coupon\_rate}, \text{price}, \\
\text{redemption}, \text{frequency}, \text{basis})
\]

**Input Parameters**

- **settlement** — The date on which payment is made to settle a trade. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **maturity** — The date on which the bond comes due, and principal and accrued interest are paid. For a more detailed discussion on dates see Chapter 8, *Working with Date/Time Data* in the PV-WAVE User’s Guide.

- **coupon\_rate** — Annual coupon rate.

- **price** — Price per $100 face value of the security.

- **redemption** — Redemption value per $100 face value of the security.

- **frequency** — Frequency of the interest payments. It should be either 1, 2 or 4.

<table>
<thead>
<tr>
<th>frequency</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>One payment per year (Annual payment)</td>
</tr>
<tr>
<td>2</td>
<td>Two payments per year (Semi-annual payment)</td>
</tr>
<tr>
<td>4</td>
<td>Four payments per year (Quarterly payment)</td>
</tr>
</tbody>
</table>

- **basis** — The method for computing the number of days between two dates. It should be either 0, 1, 2, 3 or 4.
**YIELD_PERIODIC Function**

**Returned Value**

*result* — The yield of a security that pays interest periodically. If no result can be computed, NaN is returned.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.

*Xguess* — If present, the value is used as the initial guess at the internal rate of return.

*Highest* — If present, the value is used as the maximum value of the internal rate of return allowed.

**Discussion**

Function YIELD_PERIODIC computes the yield of a security that pays periodic interest. If there is one coupon period use the following:

\[
\left( \frac{\text{redemption}}{100} + \frac{\text{coupon\_rate}}{\text{frequency}} \right) - \left[ \frac{\text{price}}{100} + \left( \frac{\text{A\_coupon\_rate}}{\text{E}} \right) \right] \right] \left( \frac{\text{frequency\_E}}{\text{DSR}} \right)
\]

In the equation above, *DSR* represents the number of days in the period starting with the settlement date and ending with the redemption date. *E* represents the number of days within the coupon period. *A* represents the number of days in the period starting with the beginning of coupon period and ending with the settlement date.
If there is more than one coupon period use the following:

\[
\text{price} = \left( \frac{\text{redemption}}{1 + \frac{\text{yield}}{\text{frequency}} \left( N - 1 + \frac{DSC}{E} \right)} \right) + \sum_{k=1}^{N} \frac{100^{\text{rate}}}{\text{frequency}} \left( \frac{\text{yield}}{1 + \frac{\text{yield}}{\text{frequency}} \left( k - 1 + \frac{DSC}{E} \right)} \right) - \left( \frac{100^{\text{rate}}}{\text{frequency}} \cdot A \right) = 0
\]

In the equation above, \( DSC \) represents the number of days in the period from the settlement to the next coupon date. \( E \) represents the number of days within the coupon period. \( N \) represents the number of coupons payable in the period starting with the settlement date and ending with the redemption date. \( A \) represents the number of days in the period starting with the beginning of the coupon period and ending with the settlement date.

**Example**

In this example, YIELD_PERIODIC computes yield of a security which is selling at $95.40663 with the settlement date of July 1, 1985, the maturity date of July 1, 1995, and the coupon rate of 6% at the issue using the US (NASD) 30/360 day count method.

```plaintext
settlement = VAR_TO_DT(2000, 7, 1)
maturity = VAR_TO_DT(2010, 7, 1)
coupon_rate = .06
price = 95.40663
redemption = 105.
frequency = 2
basis = 1
PRINT, YIELD_PERIODIC(settlement, maturity, coupon_rate, $ price, redemption, frequency, basis)
0.0700047
```
FAURE_INIT Function

Initializes the structure used for computing a shuffled Faure sequence.

Usage

\[ \text{result} = \text{FAURE_INIT}(ndim) \]

Input Parameters

\[ ndim \] — The dimension of the hyper-rectangle.

Returned Value

A structure that contains information about the sequence.

Input Keywords

\[ \text{Base} \] — The base of the Faure sequence.

Default: The smallest prime greater than or equal to \( ndim \).

\[ \text{Skip} \] — The number of points to be skipped at the beginning of the Faure sequence. Default:

\[ \left\lfloor \frac{\log B}{\log \text{base}} \right\rfloor \]

Discussion

Discrepancy measures the deviation from uniformity of a point set.
The discrepancy of the point set

\[ x_1, \ldots, x_n \in [0,1]^d, d \geq 1, \]

is

\[ D_n^{(d)} = \sup_{E} \left| \frac{\Lambda(E;n)}{n} - \lambda(E) \right|, \]

where the supremum is over all subsets of \([0,1]^d\) of the form

\[ E = \left[ 0, t_1 \right) \times \cdots \times \left[ 0, t_d \right), 0 \leq t_j \leq 1, 1 \leq j \leq d, \]

\( \lambda \) is the Lebesque measure, and

\( (E;n) \)

is the number of the \(x_j\) contained in \(E\).

The sequence \(x_1, x_2, \ldots\) of points \([0,1]^d\) is a low-discrepancy sequence if there exists a constant \(c(d)\), depending only on \(d\), such that

\[ D_n^{(d)} \leq c(d) \frac{(\log n)^d}{n} \]

for all \(n > 1\).

Generalized Faure sequences can be defined for any prime base \(b \geq d\). The lowest bound for the discrepancy is obtained for the smallest prime \(b \geq d\), so the keyword \textit{Base} defaults to the smallest prime greater than or equal to the dimension.

The generalized Faure sequence \(x_1, x_2, \ldots\), is computed as follows:

Write the positive integer \(n\) in its \(b\)-ary expansion,

\[ n = \sum_{i=0}^{\infty} a_i(n)b^i \]
where $a_i(n)$ are integers,

$$0 \leq a_i(n) < b$$

The $j$-th coordinate of $x_n$ is

$$x_n^{(j)} = \sum_{k=0}^{d-1} \sum_{d=0}^{d-1} c_{kd}^{(j)} a_d(n) b^{-k-1}, \quad 1 \leq j \leq d$$

The generator matrix for the series,

$$c_{kd}^{(j)}$$

is defined to be

$$c_{kd}^{(j)} = j^d - k^d$$

and

$$c_{kd}$$

is an element of the Pascal matrix,

$$c_{kd} = \begin{cases} 
\frac{d!}{k!(d-k)!} & k \leq d \\
0 & k > d 
\end{cases}$$

It is faster to compute a shuffled Faure sequence than to compute the Faure sequence itself. It can be shown that this shuffling preserves the low-discrepancy property.

The shuffling used is the $b$-ary Gray code. The function $G(n)$ maps the positive integer $n$ into the integer given by its $b$-ary expansion.

The sequence computed by this function is $x(G(n))$, where $x$ is the generalized Faure sequence.
Example

In this example, five points in the Faure sequence are computed. The points are in the three-dimensional unit cube.

Note that FAURE_INIT is used to create a structure that holds the state of the sequence. Each call to FAURE_NEXT_PT returns the next point in the sequence and updates the state structure.

```plaintext
state = FAURE_INIT(3)
p = FAURE_NEXT_PT(5, state)
FM, p
0.333689  0.492659  0.064065
0.667022  0.825992  0.397399
0.778133  0.270436  0.175177
0.111467  0.603770  0.508510
0.444800  0.937103  0.841843
```

**FAURE_NEXT_PT Function**

Computes a shuffled Faure sequence.

**Usage**

```plaintext
result = FAURE_NEXT_PT(npts, state)
```

**Input Parameters**

- **npts** — The number of points to generate in the hyper-rectangle.
- **state** — State structure created by a call to FAURE_INIT.

**Returned Value**

An array of size `npts` by `state.dim` containing the `npts` next points in the shuffled Faure sequence.

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.
Output Keywords

Skip — The current point in the sequence. The sequence can be restarted by initializing a new sequence using this value for Skip, and using the same dimension for ndim.

Discussion

Discrepancy measures the deviation from uniformity of a point set. The discrepancy of the point set

\[ x_1, \ldots, x_n \in [0,1]^d, d \geq 1, \]

is

\[ D_n^{(d)} = \sup_E \left| \frac{A(E:n)}{n} - \lambda(E) \right|, \]

where the supremum is over all subsets of \([0,1]^d\) of the form

\[ E = [0,t_1) \times \ldots \times [0,t_j), 0 \leq t_j \leq 1, 1 \leq j \leq d, \]

\(\lambda\) is the Lebesque measure, and

\((E;n)\)

is the number of the \(x_j\) contained in \(E\).

The sequence \(x_1, x_2, \ldots\) of points \([0,1]^d\) is a low-discrepancy sequence if there exists a constant \(c(d)\), depending only on \(d\), such that

\[ D_n^{(d)} \leq c(d) \frac{\log n}{n} \]

for all \(n > 1\).

Generalized Faure sequences can be defined for any prime base \(b \geq d\). The lowest bound for the discrepancy is obtained for the smallest prime \(b \geq d\), so the keyword Base defaults to the smallest prime greater than or equal to the dimension.
The generalized Faure sequence \( x_1, x_2, \ldots \), is computed as follows:

Write the positive integer \( n \) in its \( b \)-ary expansion

\[
n = \sum_{i=0}^{\infty} a_i(n)b^i
\]

where \( a_i(n) \) are integers,

\[
0 \leq a_i(n) < b
\]

The \( j \)-th coordinate of \( x_n \) is

\[
x_n^{(j)} = \sum_{k=0}^{\infty} \sum_{d=0}^{\infty} c_{kd}^{(j)} a_d(n) b^{-k-1}, \quad 1 \leq j \leq d
\]

The generator matrix for the series,

\[
c_d^{(j)}
\]

is defined to be

\[
c_{kd}^{(j)} = j d - k c_{kd}
\]

and

\[
c_{kd}
\]

is an element of the Pascal matrix,

\[
c_{kd} = \begin{cases} 
  d! & k \leq d \\
  c!(d-c)! & k > d \\
  0 & k > d
\end{cases}
\]

It is faster to compute a shuffled Faure sequence than to compute the Faure sequence itself. It can be shown that this shuffling preserves the low-discrepancy property.
The shuffling used is the \textit{b-ary} Gray code. The function $G(n)$ maps the positive integer $n$ into the integer given by its \textit{b-ary} expansion.

The sequence computed by this function is $x(G(n))$, where $x$ is the generalized Faure sequence.

\textbf{Example}

In this example, five points in the Faure sequence are computed. The points are in the three-dimensional unit cube.

Note that FAURE\_INIT is used to create a structure that holds the state of the sequence. Each call to FAURE\_NEXT\_PT returns the next point in the sequence and updates the state structure.

\begin{verbatim}
state = FAURE\_INIT(3)
p = FAURE\_NEXT\_PT(5, state)
PM, p
\end{verbatim}

\begin{verbatim}
0.333689  0.492659  0.0640654
0.667022  0.825992  0.397399
0.778133  0.270436  0.175177
0.111467  0.603770  0.508510
0.444800  0.937103  0.841843
\end{verbatim}

\section*{New PV-WAVE:IMSL Statistics Commands}

This section lists the new functions and procedures have been added to PV-WAVE:IMSL Statistics for version 7.5.

\section*{Chapter 8: Time Series and Forecasting}

\textbf{KALMAN Procedure}

Performs Kalman filtering and evaluates the likelihood function for the state-space model.
Usage
KALMAN, b, covb, n, ss, alndet

Input/Output Parameters

b — One dimensional array of containing the estimated state vector. The input is the estimated state vector at time \( k \) given the observations through time \( k - 1 \). The output is the estimated state vector at time \( k + 1 \) given the observations through time \( k \). On the first call to KALMAN, the input \( b \) must be the prior mean of the state vector at time.

covb — Two dimensional array of size N_ELEMENTS(\( b \)) by N_ELEMENTS(\( b \)) such that \( \text{covb} \cdot \sigma^2 \) is the mean squared error matrix for \( b \). Before the first call to KALMAN, \( \text{covb} \cdot \sigma^2 \) must equal the variance-covariance matrix of the state vector.

n — Named variable containing the rank of the variance-covariance matrix for all the observations. \( n \) must be initialized to zero before the first call to KALMAN. In the usual case when the variance-covariance matrix is nonsingular, \( n \) equals the sum of the N_ELEMENTS(\( Y \)) from the invocations to KALMAN. See the keyword section below for the definition of \( Y \).

ss — Named variable containing the generalized sum of squares. \( ss \) must be initialized to zero before the first call to KALMAN. The estimate of \( \sigma^2 \) is given by

\[
\frac{ss}{n}
\]

alndet — Named variable containing the natural log of the product of the nonzero eigenvalues of \( P \) where \( P \cdot \sigma^2 \) is the variance-covariance matrix of the observations. Although \( \text{alndet} \) is computed, KALMAN avoids the explicit computation of \( P \). \( \text{alndet} \) must be initialized to zero before the first call to KALMAN. In the usual case when \( P \) is nonsingular, \( \text{alndet} \) is the natural log of the determinant of \( P \).

Input Keywords

\( Y \) — One dimensional array containing the observations. Keywords \( Y \), \( Z \) and \( R \) indicate an update step and must be used together.
**R** — Two dimensional array if size \( \text{N}_\text{ELEMENTS}(Y) \) by \( \text{N}_\text{ELEMENTS}(Y) \) containing the matrix such that \( R \ast \sigma^2 \) is the variance-covariance matrix of errors in the observation equation. Keywords \( Y, Z \) and \( R \) indicate an update step and must be used together.

**T_matrix** — Two dimensional array if size \( \text{N}_\text{ELEMENTS}(b) \) by \( \text{N}_\text{ELEMENTS}(b) \) containing the transition matrix in the state equation.

Default: \( T\text{matrix} = \text{identity matrix} \)

**Q_matrix** — Two dimensional array if size \( \text{N}_\text{ELEMENTS}(b) \) by \( \text{N}_\text{ELEMENTS}(b) \) matrix such that \( Q\text{matrix} \ast \sigma^2 \) is the variance-covariance matrix of the error vector in the state equation.

Default: There is no error term in the state equation

**Tolerance** — Tolerance used in determining linear dependence.

Default: Tolerance = 100*eps where eps is machine precision.

**Output Keywords**

**V** — One dimensional array of length \( \text{N}_\text{ELEMENTS}(Y) \) containing the one-step-ahead prediction error.

**Covv** — Two dimensional array if size \( \text{N}_\text{ELEMENTS}(Y) \) by \( \text{N}_\text{ELEMENTS}(Y) \) containing a matrix such that \( \text{Covv} \ast \sigma^2 \) is the variance-covariance matrix of \( v \).

**Discussion**

Routine KALMAN is based on a recursive algorithm given by Kalman (1960), which has come to be known as the Kalman filter. The underlying model is known as the state-space model. The model is specified stage by stage where the stages generally correspond to time points at which the observations become available. The routine KALMAN avoids many of the computations and storage requirements that would be necessary if one were to process all the data at the end of each stage in order to estimate the state vector. This is accomplished by using previous computations and retaining in storage only those items essential for processing of future observations.

The notation used here follows that of Sallas and Harville (1981). Let \( y_k \) (input in keyword \( Y \)) be the \( n_k \times 1 \) vector of observations that become available at time \( k \). The subscript \( k \) is used here rather than \( t \), which is more customary in time series, to emphasize that the model is expressed in stages \( k = 1, 2, \ldots \) and
that these stages need not correspond to equally spaced time points. In fact, they need not correspond to time points of any kind. The observation equation for the state-space model is

\[ y_k = Z_k b_k + e_k \quad k = 1, 2, \ldots \]

Here, \( Z_k \) is an \( n_k \times q \) known matrix and \( b_k \) is the \( q \times 1 \) state vector. The state vector \( b_k \) is allowed to change with time in accordance with the state equation

\[ b_{k+1} = T_{k+1} b_k + w_{k+1} \quad k = 1, 2, \ldots \]

starting with \( b_1 = \mu_1 + w_1 \).

The change in the state vector from time \( k \) to \( k + 1 \) is explained in part by the transition matrix \( T_{k+1} \) (the identity matrix by default, or optionally input using keyword \( T\text{\_MATRIX} \)), which is assumed known. It is assumed that the \( q \)-dimensional \( w_k \)s (\( k = 1, 2, \ldots, K \)) are independently distributed multivariate normal with mean vector 0 and variance-covariance matrix \( \sigma^2 Q_{k+1} \), that the \( n_k \)-dimensional \( e_k \)s (\( k = 1, 2, \ldots, K \)) are independently distributed multivariate normal with mean vector 0 and variance-covariance matrix \( \sigma^2 R_k \), and that the \( w_k \)s and \( e_k \)s are independent of each other. Here, \( \mu_1 \) is the mean of \( b_1 \) and is assumed known, \( \sigma^2 \) is an unknown positive scalar. \( Q_{k+1} \) (input in \( Q \)) and \( R_k \) (input in keyword \( R \)) are assumed known.

Denote the estimator of the realization of the state vector \( b_k \) given the observations \( y_1, y_2, \ldots, y_j \) by

\[ \hat{b}_{kj} \]

By definition, the mean squared error matrix for

\[ \hat{b}_{kl} \]

is

\[ \sigma^2 C_{kl} = E(\hat{b}_{kl} - b_k)(\hat{b}_{kl} - b_k)^T \]
At the time of the $k$-th invocation, we have

$$\hat{\beta}_{k|k-1}$$

and

$$C_{k|k-1},$$ which were computed from the $(k-1)$-st invocation, input in $b$ and $covb$, respectively. During the $k$-th invocation, routine KALMAN computes the filtered estimate

$$\hat{\beta}_{k|k}$$

along with $C_{k|k}$. These quantities are given by the update equations:

$$\hat{\beta}_{k|k} = \hat{\beta}_{k|k-1} + C_{k|k-1}Z_k^TH_k^{-1}v_k$$

$$C_{k|k} = C_{k|k-1} - C_{k|k-1}Z_k^TH_k^{-1}Z_kC_{k|k-1}$$

where

$$v_k = y_k - Z_k\hat{\beta}_{k|k-1}$$

and where

$$H_k = R_k + Z_kC_{k|k-1}Z_k^T$$

Here, $v_k$ (stored in $v$) is the one-step-ahead prediction error, and $\sigma^2H_k$ is the variance-covariance matrix for $v_k$. $H_k$ is stored in $cov$. The “start-up values” needed on the first invocation of KALMAN are

$$\hat{\beta}_{1|0} = \mu_1$$

and $C_{1|0} = Q_f$ input via $b$ and $covb$, respectively. Computations for the $k$-th invocation are completed by KALMAN computing the one-step-ahead estimate

$$\hat{\beta}_{k+1|k}$$
along with $C_{k+1|k}$ given by the prediction equations:

$$
\hat{\beta}_{k+1|k} = T_{k+1}\hat{\beta}_{k|k}
$$

$$
C_{k+1|k} = T_{k+1}C_{k|k}T_{k+1}^T + Q_{k+1}
$$

If both the filtered estimates and one-step-ahead estimates are needed by the user at each time point, KALMAN can be invoked twice for each time point—first without $T\_matrix$ and $Q\_matrix$ to produce

$$
\hat{\beta}_{k|k}
$$

and $C_{k|k}$, and second without keywords $Y$, $Z$, and $R$ to produce

$$
\hat{\beta}_{k+1|k}
$$

and $C_{k+1|k}$ (Without $T\_matrix$ and $Q\_matrix$, the prediction equations are skipped. Without keywords $Y$, $Z$, and $R$, the update equations are skipped.).

Often, one desires the estimate of the state vector more than one-step-ahead, i.e., an estimate of

$$
\hat{\beta}_{j|j}
$$

is needed where $k > j + 1$. At time $j$, KALMAN is invoked with keywords $Y$, $Z$, and $R$ to compute

$$
\hat{\beta}_{j+1|j}
$$

Subsequent invocations of KALMAN without keywords $Y$, $Z$, and $R$ can compute

$$
\hat{\beta}_{j+2|j}, \hat{\beta}_{j+3|j}, \ldots, \hat{\beta}_{k|j}
$$

Computations for

$$
\hat{\beta}_{k|j}
$$
and $C_{klj}$ assume the variance-covariance matrices of the errors in the observation equation and state equation are known up to an unknown positive scalar multiplier, $\sigma^2$. The maximum likelihood estimate of $\sigma^2$ based on the observations $y_1, y_2, \ldots, y_m$, is given by

$$\hat{\sigma}^2 = SS / N$$

where

$$N = \sum_{k=1}^{m} n_k \text{ and } SS = \sum_{k=1}^{m} v_k^T H_k^{-1} v_k$$

$N$ and $SS$ are the input/output arguments $n$ and $ss$.

If $\sigma^2$ is known, the $R_{k}$s and $Q_{k}$s can be input as the variance-covariance matrices exactly. The earlier discussion is then simplified by letting $\sigma^2 = 1$.

In practice, the matrices $T_k$, $Q_k$, and $R_k$ are generally not completely known. They may be known functions of an unknown parameter vector $\theta$. In this case, KALMAN can be used in conjunction with an optimization program (see routine FMINV, PV-WAVE: IMSL Mathematics Reference, Chapter 8, “Optimization”) to obtain a maximum likelihood estimate of $\theta$. The natural logarithm of the likelihood function for $y_1, y_2, \ldots, y_m$ differs by no more than an additive constant from

$$L(\theta, \sigma^2; y_1, y_2, \ldots, y_m) \propto \frac{1}{2} N \sigma^2$$

$$\frac{1}{2} \sum_{k=1}^{m} \left[ (H_k) \right] \frac{1}{2} \sigma^2 \sum_{k=1}^{m} v_k^T H_k^{-1} v_k$$

(Harvey 1981, page 14, equation 2.21).

Here,
(stored in $\text{alndet}$) is the natural logarithm of the determinant of $V$ where $\sigma^2 V$ is the variance-covariance matrix of the observations.

Minimization of $-2L(\theta, \sigma^2; y_1, y_2, \ldots, y_m)$ over all $\theta$ and $\sigma^2$ produces maximum likelihood estimates. Equivalently, minimization of $-2L_c(\theta; y_1, y_2, \ldots, y_m)$ where

$$L_c(\theta; y_1, y_2, \ldots, y_m) = -\frac{1}{2} N \ln \left( \frac{SS}{N} \right) - \frac{1}{2} \sum_{k=1}^{m} \ln \{\text{det}(H_k)\}$$

produces maximum likelihood estimates

$$\hat{\theta} \text{ and } \hat{\sigma}^2 = \frac{SS}{N}$$

The minimization of $-2L_c(\theta; y_1, y_2, \ldots, y_m)$ instead of $-2L(\theta, \sigma^2; y_1, y_2, \ldots, y_m)$, reduces the dimension of the minimization problem by one. The two optimization problems are equivalent since

$$\hat{\sigma}^2(\theta) = \frac{SS(\theta)}{N}$$

minimizes $-2L(\theta, \sigma^2; y_1, y_2, \ldots, y_m)$ for all $\theta$, consequently,

$$\hat{\sigma}^2(\theta)$$

can be substituted for $\sigma^2$ in $L(\theta, \sigma^2; y_1, y_2, \ldots, y_m)$ to give a function that differs by no more than an additive constant from $L_c(\theta; y_1, y_2, \ldots, y_m)$.

The earlier discussion assumed $H_k$ to be nonsingular. If $H_k$ is singular, a modification for singular distributions described by Rao (1973, pages 527–528) is used. The necessary changes in the preceding discussion are as follows:

1. Replace

$$H_k^{-1}$$

by a generalized inverse.

2. Replace $\text{det}(H_k)$ by the product of the nonzero eigenvalues of $H_k$. 
3. Replace \( N \) by

\[
\sum_{k=1}^{m} \text{rank}(H_k)
\]


**Example 1**

Routine KALMAN is used to compute the filtered estimates and one-step-ahead estimates for a scalar problem discussed by Harvey (1981, pages 116–117). The observation equation and state equation are given by

\[
\begin{align*}
y_k &= b_k + e_k \\
b_{k+1} &= b_k + w_{k+1} \\
&k = 1, 2, 3, 4
\end{align*}
\]

where the \( e_k \)'s are identically and independently distributed normal with mean 0 and variance \( \sigma^2 \), the \( w_k \)'s are identically and independently distributed normal with mean 0 and variance \( 4\sigma^2 \), and \( b_1 \) is distributed normal with mean 4 and variance \( 16\sigma^2 \). Two invocations of KALMAN are needed for each time point in order to compute the filtered estimate and the one-step-ahead estimate. The first invocation does not use the keywords \( T \_matrix \) and \( Q \_matrix \) so that the prediction equations are skipped in the computations. The update equations are skipped in the computations in the second invocation.

This example also computes the one-step-ahead prediction errors. Harvey (1981, page 117) contains a misprint for the value \( \nu_4 \) that he gives as 1.197. The correct value of \( \nu_4 = 1.003 \) is computed by KALMAN.

Note that this example is in the form of a WAVE procedure, with the output following the procedure.

PRO EX_KALMAN

\[
\begin{align*}
z &= 1 \\
r &= 1 \\
q &= 4 \\
t &= 1
\end{align*}
\]
b = 4
covb = 16

ydata = [4.4, 4, 3.5, 4.6]

n = 0
ss = 0
alndet = 0
format = "(2I4, 2F8.3, I4, 4F8.3)"
PRINT, " k j b covb n ss alndet v covv"
FOR i = 0, 3 DO BEGIN
  y = ydata(i)
  ; Update
  kalman, b, covb, n, ss, alndet, $
  Y = y, Z = Z, R = r, $
  v = v, covv = covv
  PRINT, i, i, b, covb, n, ss, alndet, v, covv, format = format

  ; Predict
  kalman, b, covb, n, ss, alndet, $
  t_matrix = t, q = q
  PRINT, i+1, i, b, covb, n, ss, alndet, v, covv, format = format
END
END

Output

<table>
<thead>
<tr>
<th>k</th>
<th>j</th>
<th>b</th>
<th>covb</th>
<th>n</th>
<th>ss</th>
<th>alndet</th>
<th>v</th>
<th>covv</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>4.376</td>
<td>0.941</td>
<td>1</td>
<td>0.009</td>
<td>2.833</td>
<td>0.400</td>
<td>17.000</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>4.376</td>
<td>4.941</td>
<td>1</td>
<td>0.009</td>
<td>2.833</td>
<td>0.400</td>
<td>17.000</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>4.063</td>
<td>0.832</td>
<td>2</td>
<td>0.033</td>
<td>4.615</td>
<td>-0.376</td>
<td>5.941</td>
</tr>
</tbody>
</table>
Chapter 11: Probability Distribution Functions and Inverses

**BINOMIALPDF Function**

Evaluates the binomial probability function.

**Usage**

\[ result = \text{BINOMIALPDF} (k, n, p) \]

**Input Parameters**

- \( k \) — Argument for which the binomial probability function is to be evaluated.
- \( n \) — Number of Bernoulli trials.
- \( p \) — Probability of success on each trial.

**Returned Value**

\( result \) — The probability that a binomial random variable takes a value equal to \( k \).

**Discussion**

The function BINOMIALPDF evaluates the probability that a binomial random variable with parameters \( n \) and \( p \) takes on the value \( k \). It does this by computing probabilities of the random variable taking on the values in its range less than
(or the values greater than) $k$. These probabilities are computed by the recursive relationship

$$\Pr(X = j) = \frac{(n+1-j)p}{j(1-p)} \Pr(X = j-1)$$

To avoid the possibility of underflow, the probabilities are computed forward from 0, if $k$ is not greater than $n$ times $p$, and are computed backward from $n$, otherwise. The smallest positive machine number, $\epsilon$, is used as the starting value for computing the probabilities, which are rescaled by $(1-p)^n\epsilon$ if forward computation is performed and by $p^n\epsilon$ if backward computation is done.

For the special case of $p = 0$, BINOMIALPDF returns 0 if $k$ is greater than 0 and to 1 otherwise; and for the case $p = 1$, BINOMIALPDF returns 0 if $k$ is less than $n$ and to 1 otherwise.

**Example**

Suppose $X$ is a binomial random variable with $n = 5$ and $p = 0.95$. In this example, we find the probability that $X$ is equal to 3.

```plaintext
PRINT, BINOMIALPDF(3, 5, .95)
0.0214344
```

---

**Chapter 12: Random Number Generation**

**RANDOM_TABLE Procedure**

Sets or retrieves the current table used in either the shuffled or GFSR random number generator.

**Usage**

```plaintext
RANDOM_TABLE, table, /Get
RANDOM_TABLE, table, /Set
```
Input/Output Parameters

`table` — One dimensional array used in the generators. For the shuffled generators table is length 128. For the GFSR generator table is length 1565. The argument `table` is input if the keyword `Set` is used, and output if the keyword `Get` is used.

Input Keywords

`Set` — If present and nonzero, then the specified table is being set.

`Get` — If present and nonzero, then the specified table is being retrieved.

`Gfsr` — If present and nonzero, then the specified GFSR table is being set or retrieved.

`Double` — If present and nonzero, double precision is used. This keyword is active only when the shuffled table is being set or retrieved.

Discussion

The values in `table` are initialized by the IMSL random number generators. The values are all positive except if the user wishes to reinitialize the array, in which case the first element of the array is input as a nonpositive value. (Usually, one should avoid reinitializing these arrays, but it might be necessary sometimes in restarting a simulation.) If the first element of `table` is set to a nonpositive value on the call to `RANDOM_TABLE` with the keyword `Set`, on the next invocation of a routine to generate random numbers, the appropriate table will be reinitialized.

For more details on the shuffled and GFSR generators see the *Introduction* to Chapter 12 in the *PV-WAVE: IMSL Statistics Reference*.

Example

In this example, three separate simulation streams are used, each with a different form of the generator. Each stream is stopped and restarted. (Although this example is obviously an artificial one, there may be reasons for maintaining separate streams and stopping and restarting them because of the nature of the usage of the random numbers coming from the separate streams.)

```plaintext
nr = 5
iseed1 = 123457
iseed2 = 123457
```
iseed7 = 123457

; Begin first stream, iopt = 1 (by default)
RANDOMOPT, Set = iseed1
r = RANDOM(nr)
RANDOMOPT, Get = iseed1
PM, r, Title = 'First stream output'
First stream output
  0.966220
  0.260711
  0.766262
  0.569337
  0.844829
PRINT, 'output seed ', iseed1
output seed 1814256879

; Begin second stream, iopt = 2
RANDOMOPT, gen_opt = 2
RANDOMOPT, Set = iseed2
r = RANDOM(nr)
RANDOMOPT, Get = iseed2
RANDOM_TABLE, table, /Get
PM, r, Title = 'Second stream output'
Second stream output
  0.709518
  0.186145
  0.479442
  0.603839
  0.379015
PRINT, 'output seed ', iseed2
output seed 1965912801

; Begin third stream, iopt = 7
RANDOMOPT, gen_opt = 7
RANDOMOPT, Set = iseed7
r = RANDOM(nr)
RANDOMOPT, Get = iseed7
RANDOM_TABLE, itable, /Get, /GFSR
PM, r, Title = 'Third stream output'

Third stream output
  0.391352
  0.0262676
  0.762180
  0.0280987
  0.899731
PRINT, 'output seed ', iseed7
output seed   1932158269

; Reinitialize seed and resume first stream
RANDOMOPT, gen_opt = 1
RANDOMOPT, Set = iseed1
r = RANDOM(nr)
RANDOMOPT, Get = iseed1
pm, r, title = 'First stream output'

First stream output
  0.0442665
  0.987184
  0.601350
  0.896375
  0.380854
PRINT, 'output seed ', iseed1
output seed    817878095

; Reinitialize seed and table for shuffling and
; resume second stream
RANDOMOPT, gen_opt = 2
RANDOMOPT, Set = iseed2
RANDOM_TABLE, table, /Set
r = RANDOM(nr)
RANDOMOPT, Get = iseed2
PM, r, Title = 'Second stream output'

Second stream output
  0.255690
  0.478770
  0.225802
  0.345467
  0.581051

PRINT, 'output seed ', iseed2
output seed   2108806573

; Reinitialize seed and table for GFSR and
; resume third stream.
RANDOMOPT,  gen_opt = 7
RANDOMOPT,  Set = iseed7
RANDOM_TABLE, itable, /Set, /Gfsr
r = RANDOM(nr)
RANDOMOPT,  Get = iseed7
PM, r, Title = 'Third stream output'

Third stream output
  0.751854
  0.508370
  0.906986
  0.0910035
  0.691663

PRINT, 'output seed ', iseed7
output seed   1485334679

---

**RANDOM_NPP Function**

Generates pseudorandom numbers from a nonhomogeneous Poisson process.

**Usage**

\[ \text{result} = \text{RANDOM\_NPP}(\text{tbegin}, \text{tend}, ftheta, \text{theta_min}, \text{theta_max}, \text{neub}) \]
Input Parameters

\textit{tbegin} — Lower endpoint of the time interval of the process. \textit{tbegin} must be nonnegative. Usually, \textit{tbegin} = 0.

\textit{tend} — Upper endpoint of the time interval of the process. \textit{tend} must be greater than \textit{tbegin}.

\textit{ftheta} — Scalar string specifying a user-supplied function to provide the value of the rate of the process as a function of time. This function accepts one argument and must be defined over the interval from \textit{tbegin} to \textit{tend} and must be nonnegative in that interval.

\textit{theta\_min} — Minimum value of the rate function \textit{ftheta()} in the interval (\textit{tbegin}, \textit{tend}). If the actual minimum is unknown, set \textit{theta\_min} = 0.0.

\textit{theta\_max} — Maximum value of the rate function \textit{ftheta} in the interval (\textit{tbegin}, \textit{tend}). If the actual maximum is unknown, set \textit{theta\_max} to a known upper bound of the maximum. The efficiency of RANDOM\_NPP is less the greater \textit{theta\_max} exceeds the true maximum.

\textit{neub} — Upper bound on the number of events to be generated. In order to be reasonably sure that the full process through time \textit{tend} is generated, calculate \textit{neub} as \textit{neub} = \textit{x} + 10.0 \times \text{SQRT}(\textit{x}), where \textit{x} = \textit{theta\_max} \times (\textit{tend} - \textit{tbegin}).

Returned Value

A one dimensional array containing the times to events. If then length of the result is less that \textit{neub}, the time \textit{tend} is reached before \textit{neub} events are realized

Input Keywords

\textit{Double} — If present and nonzero, double precision is used.

Discussion

Routine RANDOM\_NPP simulates a one-dimensional nonhomogeneous Poisson process with rate function \textit{theta} in a fixed interval (\textit{tend} - \textit{tbegin}). Let \( \lambda(t) \) be the rate function and \textit{t0} = \textit{tbegin} and \textit{t1} = \textit{tend}. Routine RANDOM\_NPP uses a method of thinning a nonhomogeneous Poisson process
\{N^*(t), \ t \geq t_0\} \text{ with rate function } \lambda^*(t) \geq \lambda(t) \text{ in } (t_0, t_1], \text{ where the number of events, } N^*, \text{ in the interval } (t_0, t_1] \text{ has a Poisson distribution with parameter}

\[ \mu_0 = \int_{t_0}^{t_1} \lambda(t) \, dt \]

The function

\[ \Lambda(t) = \int_0^t \lambda^*(t) \, dt \]

is called the \textit{integrated rate function}. In RANDOM_NPP, \( \lambda^*(t) \) is taken to be a constant \( \lambda^*(= \theta_{\text{max}}) \) so that at time \( t_i \), the time of the next event \( t_{i+1} \) is obtained by generating and cumulating exponential random numbers

\[ E_{1,i}^*, E_{2,i}^*, \ldots, \]

with parameter \( \lambda^* \), until for the first time

\[ u_{j,i} \leq \left( t_i + E_{1,i}^* + \cdots + E_{j,i}^* \right) / \lambda^* \]

where the \( u_{j,i} \) are independent uniform random numbers between 0 and 1. This process is continued until the specified number of events, \( n_{\text{ub}} \), is realized or until the time, \( t_{\text{end}} \), is exceeded. This method is due to Lewis and Shedler (1979), who also review other methods. The most straightforward (and most efficient) method is by inverting the integrated rate function, but often this is not possible.

If \( \theta_{\text{max}} \) is actually greater than the maximum of \( \lambda(t) \) in \( (t_0, t_1] \), the routine will work, but less efficiently. Also, if \( \lambda(t) \) varies greatly within the interval, the efficiency is reduced. In that case, it may be desirable to divide the time interval into subintervals within which the rate function is less variable. This is possible because the process is without memory.

If no time horizon arises naturally, \( t_{\text{end}} \) must be set large enough to allow for the required number of events to be realized. Care must be taken, however, that \( f_{\theta} \) is defined over the entire interval.

After simulating a given number of events, the next event can be generated by setting \( \text{tbegin} \) to the time of the last event (the sum of the elements in the result)
and calling RANDOM_NPP again. Cox and Lewis (1966) discuss modeling applications of nonhomogeneous Poisson processes.

**Example**

In this example, RANDOM_NPP is used to generate the first five events in the time 0 to 20 (if that many events are realized) in a nonhomogeneous process with rate function

\[ \lambda(t) = 0.6342 \, e^{0.001427 \cdot t} \]

for \( 0 < t \leq 20 \).

Since this is a monotonically increasing function of \( t \), the minimum is at \( t = 0 \) and is 0.6342, and the maximum is at \( t = 20 \) and is

\[ 0.6342 \, e^{0.02854} = 0.652561. \]

```
.RUN
- FUNCTION ftheta_npp, t
-   return, .6342*exp(.001427*t)
- END
% Compiled module: FTHETA_NPP.

randomopt, set=123457
neub = 5
tmax = .652561
tmin = .6342
tbegin=0
tend=20

r = RANDOM_NPP(tbegin, tend, 'ftheta_npp', tmin, tmax, neub)
PM, r
0.0526598
0.407979
0.258399
0.0197666
0.167641
```
**RANDOM_ORDER Function**

Generates pseudorandom order statistics from a uniform (0, 1) distribution, or optionally from a standard normal distribution.

**Usage**

```plaintext
result = RANDOM_ORDER(ifirst, ilast, n)
```

**Input Parameters**

- **ifirst** — First order statistic to generate.
- **ilast** — Last order statistic to generate.
- **ilast** must be greater than or equal to **ifirst**. The full set of order statistics from **ifirst** to **ilast** is generated. If only one order statistic is desired, set **ilast = ifirst**.
- **n** — Size of the sample from which the order statistics arise.

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.
- **Uniform** — If present and nonzero, generate pseudorandom order statistics from a uniform (0, 1) distribution. (Default)
- **Normal** — If present and nonzero, generate pseudorandom order statistics from a standard normal distribution.

**Returned Value**

An array of length **ilast + 1 - ifirst** containing the random order statistics in ascending order.

The first element is the **ifirst** order statistic in a random sample of size **n** from the uniform (0, 1) distribution.

**Discussion**

Routine RANDOM_ORDER generates the **ifirst** through the **ilast** order statistics from a pseudorandom sample of size **n** from a uniform (0, 1) distribution. Depending on the values of **ifirst** and **ilast**, different methods of generation are used to achieve greater efficiency. If **ifirst = 1** and
ilast = n, that is, if the full set of order statistics are desired, the spacings between successive order statistics are generated as ratios of exponential variates. If the full set is not desired, a beta variate is generated for one of the order statistics, and the others are generated as extreme order statistics from conditional uniform distributions. Extreme order statistics from a uniform distribution can be obtained by raising a uniform deviate to an appropriate power.

Each call to RANDOM_ORDER yields an independent event. This means, for example, that if on one call the fourth order statistic is requested and on a second call the third order statistic is requested, the “fourth” may be smaller than the “third”. If both the third and fourth order statistics from a given sample are desired, they should be obtained from a single call to RANDOM_ORDER (by specifying ifirst less than or equal to 3 and ilast greater than or equal to 4).

If the keyword Normal is present and nonzero, then RANDOM_ORDER generates the ifirst through the ilast order statistics from a pseudorandom sample of size n, from a normal (0, 1) distribution.

Example

In this example, RANDOM_ORDER is used to generate the fifteenth through the nineteenth order statistics from a sample of size twenty.

```plaintext
r = random_order(15, 19, 20)
```

```plaintext
pm, r
0.706909
0.808627
0.874552
0.922146
0.957402
```

**RAND_TABLE_2WAY Function**

Generates a pseudorandom two-way table.

**Usage**

```
result = RAND_TABLE_2WAY (row_totals, col_totals)
```
Input Parameters

*row_totals* — One dimensional array containing the row totals.

*col_totals* — One dimensional array containing the column totals. (Input)

The elements of *row_totals* and *col_totals* must be nonnegative and must sum to the same quantity.

Returned Value

A N_ELEMENTS(*row_totals*) by N_ELEMENTS(*col_totals*) random matrix with the given row and column totals.

Discussion

Routine RAND_TABLE_2WAY generates pseudorandom entries for a two-way contingency table with fixed row and column totals. The method depends on the size of the table and the total number of entries in the table. If the total number of entries is less than twice the product of the number of rows and columns, the method described by Boyette (1979) and by Agresti, Wackerly, and Boyette (1979) is used. In this method, a work vector is filled with row indices so that the number of times each index appears equals the given row total. This vector is then randomly permuted and used to increment the entries in each row so that the given row total is attained.

For tables with larger numbers of entries, the method of Patefield (1981) is used. This method can be considerably faster in these cases. The method depends on the conditional probability distribution of individual elements, given the entries in the previous rows. The probabilities for the individual elements are computed starting from their conditional means.

Example

In this example, RAND_TABLE_2WAY is used to generate a two by three table with row totals 3 and 5, and column totals 2, 4, and 2.

\[
\begin{array}{ccc}
2 & 1 & 0 \\
0 & 3 & 2 \\
\end{array}
\]
**RAND_ORTH_MAT Function**

Generates a pseudorandom orthogonal matrix or a correlation matrix.

**Usage**

\[
result = \text{RAND\_ORTH\_MAT}(n)
\]

**Input Parameters**

\(n\) — The order of the matrix to be generated.

**Returned Value**

A two-dimensional array containing the \(n\) by \(n\) random correlation matrix.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.

*Eigenvalues* — A one-dimensional array of length \(n\) containing the eigenvalues of the correlation matrix to be generated. The elements of *Eigenvalues* must be positive, they must sum to \(n\), and they cannot all be equal.

*A\_Matrix* — A two-dimensional array containing \(n\) by \(n\) random orthogonal matrix. A random correlation matrix is generated using the orthogonal matrix input in *A\_Matrix*. The keyword *Eigenvalues* must also be supplied if *A\_Matrix* is used.

**Discussion**

Routine RAND\_ORTH\_MAT generates a pseudorandom orthogonal matrix from the invariant Haar measure. For each column, a random vector from a uniform distribution on a hypersphere is selected and then is projected onto the orthogonal complement of the columns already formed. The method is described by Heiberger (1978). (See also Tanner and Thisted 1982.)

If the keyword *Eigenvalues* is used, a correlation matrix is formed by applying a sequence of planar rotations to the matrix \(A^T D A\), where \(D = \text{diag}(\text{Eigenvalues}(0), \ldots, \text{Eigenvalues}(n-1))\), so as to yield ones along the diagonal. The planar rotations are applied in such an order that in the two by two matrix that determines the rotation, one diagonal element is less than 1.0 and one is greater
than 1.0. This method is discussed by Bendel and Mickey (1978) and by Lin and Bendel (1985).

The distribution of the correlation matrices produced by this method is not known. Bendel and Mickey (1978) and Johnson and Welch (1980) discuss the distribution.

For larger matrices, rounding can become severe; and the double precision results may differ significantly from single precision results.

**Example**

In this example, RAND_ORTH_MAT is used to generate a 4 by 4 pseudorandom correlation matrix with eigenvalues in the ratio 1:2:3:4.

```plaintext
RANDOMOPT, set = 123457
a = RAND_ORTH_MAT(4)
ev = .4d0*[1.0d0, 2.0d0, 3.0d0, 4.0d0]
cor = RAND_ORTH_MAT(n, Eigenvalues = ev, A_Matrix= a)
PM, cor
  1.00000   -0.235786   -0.325795   -0.110139
-0.235786     1.00000    0.190564   -0.0172391
-0.325795     0.190564    1.00000   -0.435339
-0.110139   -0.0172391   -0.435339     1.00000
```

**RANDOM_SAMPLE Function**

Generates a simple pseudorandom sample from a finite population.

**Usage**

```
result = RANDOM_SAMPLE(nsamp, population)
```

**Input Parameters**

- **nsamp** — The sample size desired.
- **population** — A one or two dimensional array containing the population to be sampled. If either of the keywords *First_Call* or *Additional_Call* are specified, then *population* contains a different part of the population on each invocation, otherwise *population* contains the entire population.
**Returned Value**

\( nsamp \) by \( nvar \) array containing the sample, where \( nvar \) is the number of columns in the argument \( population \).

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.

- **First_Call** — If present and nonzero, then this is the first invocation with this data; additional calls to \( \text{RANDOM\_SAMPLE} \) may be made to add to the population. Additional calls should be made using the keyword **Additional_Call**. Keywords **Index** and **Npop** are required if **First_Call** is set. See Example 2.

- **Additional_Call** — If present and nonzero, then this is an additional invocation of \( \text{RANDOM\_SAMPLE} \), and updating for the subpopulation in \( \text{population} \) is performed. Keywords **Index**, **Npop** and **Sample** are required if **Additional_Call** is set. It is not necessary to know the number of items in the population in advance. **Npop** is used to cumulate the population size and should not be changed between calls to \( \text{RANDOM\_SAMPLE} \). See Example 2.

**Input/Output Keywords**

- **Index** — A one-dimensional array of length \( nsamp \) containing the indices of the sample in the population. Output if keyword **First_Call** is used. Input/Output if keyword **Additional_Call** is used.

- **Npop** — The number of items in the population. Output if keyword **First_Call** is used. Input/Output if keyword **Additional_Call** is used.

- **Sample** — An array of size \( nsamp \) by \( nvar \) containing the sample. Initially, the result of calling \( \text{RANDOM\_SAMPLE} \) with keyword **First_Call** is used for **Sample**.

**Discussion**

Routine \( \text{RANDOM\_SAMPLE} \) generates a pseudorandom sample from a given population, without replacement, using an algorithm due to McLeod and Bellhouse (1983).

The first \( nsamp \) items in the population are included in the sample. Then, for each successive item from the population, a random item in the sample is replaced by that item from the population with probability equal to the sample
size divided by the number of population items that have been encountered at that time.

### Example 1

In this example, RANDOM_SAMPLE is used to generate a sample of size 5 from a population stored in the matrix `population`.

```plaintext
RANDOMOPT, Set = 123457
pop = STATDATA(2)
samp = RANDOM_SAMPLE(5, pop)
PM, samp
```

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1764.00</td>
<td>36.4000</td>
<td></td>
</tr>
<tr>
<td>1828.00</td>
<td>62.5000</td>
<td></td>
</tr>
<tr>
<td>1923.00</td>
<td>5.80000</td>
<td></td>
</tr>
<tr>
<td>1773.00</td>
<td>34.8000</td>
<td></td>
</tr>
<tr>
<td>1769.00</td>
<td>106.100</td>
<td></td>
</tr>
</tbody>
</table>

### Example 2

Routine RANDOM_SAMPLE is now used to generate a sample of size 5 from the same population as in the example above except the data are input to RANDOM_SAMPLE one observation at a time. This is the way RANDOM_SAMPLE may be used to sample from a file on disk or tape. Notice that the number of records need not be known in advance.

```plaintext
RANDOMOPT, Set = 123457
pop = STATDATA(2)
samp = RANDOM_SAMPLE(5, pop(0, *), /First_Call, Index = ii, Npop=np)
FOR i=1,175 DO samp = RANDOM_SAMPLE(5, pop(i, *), /Additional_Call, $index = ii, npop = np, sample = samp)
PM, samp
```

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1764.00</td>
<td>36.4000</td>
<td></td>
</tr>
<tr>
<td>1828.00</td>
<td>62.5000</td>
<td></td>
</tr>
<tr>
<td>1923.00</td>
<td>5.80000</td>
<td></td>
</tr>
<tr>
<td>1773.00</td>
<td>34.8000</td>
<td></td>
</tr>
<tr>
<td>1769.00</td>
<td>106.100</td>
<td></td>
</tr>
</tbody>
</table>
**RAND_FROM_DATA Function**

Generates pseudorandom numbers from a multivariate distribution determined from a given sample.

**Usage**

\[
result = \text{RAND\_FROM\_DATA}(n\_random, x, nn)
\]

**Input Parameters**

\textit{n\_random} — Number of random multivariate vectors to generate.

\textit{x} — Two dimensional array of size \textit{nsamp} by \textit{ndim} containing the given sample.

\textit{nn} — Number of nearest neighbors of the randomly selected point in \textit{x} that are used to form the output point in the result.

**Returned Value**

\textit{n} by \textit{ndim} matrix containing the random multivariate vectors in its rows.

**Input Keywords**

\textit{Double} — If present and nonzero, double precision is used.

**Discussion**

Given a sample of size \textit{nsamp} of observations of a \textit{k-variate} random variable, RAND_FROM_DATA generates a pseudorandom sample with approximately the same moments as the given sample. The sample obtained is essentially the same as if sampling from a Gaussian kernel estimate of the sample density. (See Thompson 1989.) Routine RAND_FROM_DATA uses methods described by Taylor and Thompson (1986).

Assume that the (vector-valued) observations \(x_i\) are in the rows of \(x\). An observation, \(x_j\), is chosen randomly; its nearest \(m = \textit{nn}\) neighbors,

\[x_{j_1}, x_{j_2}, \ldots, x_{j_m}\]
are determined; and the mean

\[ \bar{x}_j \]

of those nearest neighbors is calculated. Next, a random sample

\( u_1, u_2, \ldots, u_m \) is generated from a uniform distribution with lower bound

\[
\frac{1}{m} - \sqrt{\frac{3(m-1)}{m^2}}
\]

and upper bound

\[
\frac{1}{m} + \sqrt{\frac{3(m-1)}{m^2}}
\]

\[
\sum_{i=1}^{m} u_i (x_{ji} - \bar{x}_j) + \bar{x}_j \]

delivered is

The process is then repeated until \( n \) such simulated variates are generated and stored in the rows of the result.

**Example**

In this example, RAND_FROM_DATA is used to generate 5 pseudorandom vectors of length 4 using the initial and final systolic pressure and the initial and final diastolic pressure from Data Set A in Afifi and Azen (1979) as the fixed sample from the population to be modeled. (Values of these four variables are in the seventh, tenth, twenty-first, and twenty-fourth columns of data set number nine in routine STATDATA, see Chapter 13: *Utilities* of this manual).

```
RANDOMOPT, Set = 123457
r = STATDATA(9)
x =FLTARR(113, 4)
x(*, 0) = r(*,6)
x(*, 1) = r(*,9)
x(*, 2) = r(*,20)
x(*, 3) = r(*,23)
r = RAND_FROM_DATA(5, x, 5)
```

CONT_TABLE Procedure

Sets up table to generate pseudorandom numbers from a general continuous distribution.

Usage

CONT_TABLE, f, iopt, ndata, table

Input Parameters

f — A scalar string specifying a user-supplied function to compute the cumulative distribution function. The argument to the function is the point at which the distribution function is to be evaluated.

iopt — Indicator of the extent to which table is initialized prior to calling CONT_TABLE.

<table>
<thead>
<tr>
<th>iopt</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>CONT_TABLE fills the last four columns of table. The user inputs the points at which the CDF is to be evaluated in the first column of table. These must be in ascending order.</td>
</tr>
<tr>
<td>1</td>
<td>CONT_TABLE fills the last three columns of table. The user supplied function f is not used and may be a dummy function; instead, the cumulative distribution function is specified in the first two columns of table. The abscissas (in the first column) must be in ascending order and the function must be strictly monotonically increasing.</td>
</tr>
</tbody>
</table>
$ndata$ — Number of points at which the CDF is evaluated for interpolation. $ndata$ must be greater than or equal to 4.

**Input/Output Parameters**

$table$ — $ndata$ by 5 table to be used for interpolation of the cumulative distribution function. The first column of $table$ contains abscissas of the cumulative distribution function in ascending order, the second column contains the values of the CDF (which must be strictly increasing), and the remaining columns contain values used in interpolation. The first row of $table$ corresponds to the left limit of the support of the distribution and the last row corresponds to the right limit of the support; that is, $table(0, 1) = 0.0$ and $table(ndata-1, 1) = 1.0$.

**Input Keywords**

$Double$ — If present and nonzero, double precision is used.

**Discussion**

Routine CONT_TABLE sets up a table that routine RAND_GEN_CONT (page 219) can use to generate pseudorandom deviates from a continuous distribution. The distribution is specified by its cumulative distribution function, which can be supplied either in tabular form in $table$ or by a function $f$. See the documentation for the routine RAND_GEN_CONT for a description of the method.

**Example**

For an example of using CONT_TABLE see the example for routine RAND_GEN_CONT (page 219).

---

**RAND_GEN_CONT Function**

Generates pseudorandom numbers from a general continuous distribution.

**Usage**

\[
result = \text{RAND\_GEN\_CONT}(n, table)
\]
**Input Parameters**

- **n** — Number of random numbers to generate.

- **table** — A two-dimensional array setup using CONT_TABLE to be used for interpolation of the cumulative distribution function. The first column of `table` contains abscissas of the cumulative distribution function in ascending order, the second column contains the values of the CDF (which must be strictly increasing beginning with 0.0 and ending at 1.0) and the remaining columns contain values used in interpolation.

**Returned Value**

An array of length `n` containing the random deviates.

**Input Keywords**

- **Double** — If present and nonzero, double precision is used.

**Discussion**

Routine RAND_GEN_CONT generates pseudorandom numbers from a continuous distribution using the inverse CDF technique, by interpolation of points of the distribution function given in `table`, which is set up by routine CONT_TABLE (page 218). A strictly monotone increasing distribution function is assumed. The interpolation is by an algorithm attributable to Akima (1970), using piecewise cubics. The use of this technique for generation of random numbers is due to Guerra, Tapia, and Thompson (1976), who give a description of the algorithm and accuracy comparisons between this method and linear interpolation. The relative errors using the Akima interpolation are generally considered very good.

**Example**

In this example, RAND_GEN_CONT (page 219) is used to set up a table for generation of beta pseudorandom deviates. The CDF for this distribution is computed by the routine BETACDF (Chapter 11). The table contains 100 points at which the CDF is evaluated and that are used for interpolation. Notice that two warnings are issued during the computations for this example.

```fortran
FUNCTION cdf, x
    return, BETACDF(x, 3., 2.)
```

RAND_GEN_CONT Function 219
```
iopt = 0
ndata = 100;
table = FLTARR(100, 5)
x = 0.0;
table(*,0) = FINDGEN(100)/100.
CONT_TABLE, 'cdf', iopt, ndata, table
RANDOMOPT, Set = 123457

r = RAND_GEN_CONT(5, table)
% BETACDF: Note: STAT_ZERO_AT_X
  Since "X" = 0.000000e+00 is less than or equal to zero,
  the distribution function is zero at "x".
% CONT_TABLE: Warning: STAT_SECOND_COL_TABLE3
CDF in the second column of table did not begin at 0.0
and end at 1.0, but they have been adjusted. Prior
to adjustment, table(0, 1) = 0.000000e+00 and
table(ndata-1, 1) = 9.994079e-01.

PM, r
0.92079391
0.46412855
0.76678398
0.65357975
0.81706959
```

**DISCR_TABLE Function**

Sets up table to generate pseudorandom numbers from a general discrete distribution.

**Usage**

```
result = DISCR_TABLE(prf, del, ndx, imin, nmass)
```
**Input Parameters**

*prf* — A scalar string specifying a user-supplied function to compute the probability associated with each mass point of the distribution. The argument to the function is the point at which the probability function is to be evaluated. The argument to the function can range from *imin* to the value at which the cumulative probability is greater than or equal to $1.0 - del$.

*del* — Maximum absolute error allowed in computing the cumulative probability. Probabilities smaller than *del* are ignored; hence, *del* should be a small positive number. If *del* is too small, however, $cumpr(nmass-1)$ must be exactly 1.0 since that value is compared to $1.0 - del$.

*nndx* — The number of elements of *cumpr* available to be used as indexes. *nndx* must be greater than or equal to 1. In general, the larger *nndx* is, to within sixty or seventy percent of *nmass*, the more efficient the generation of random numbers using RAND_GEN_DISCR will be.

**Input/Out Parameters**

*imin* — Scalar containing the smallest value the random deviate can assume. By default, *prf* is evaluated at *imin*. If this value is less than *del*, *imin* is incremented by 1 and again *prf* is evaluated at *imin*. This process is continued until $prf(imin) \geq del$. *imin* is output as this value and *result*(0) is output as $prf(imin)$.

*nmass* — Scalar containing the number of mass points in the distribution. Input, if keyword *CUM_probs* is used; otherwise, output. By default, *nmass* is the smallest integer such that $prf(imin + nmass - 1) > 1.0 - del$. *nmass* does include the points $imin + j$ for which $prf(imin + j) < del$, for $j = 0, 1, \ldots,$ $imin_{out} - imin_{in}$, where $imin_{in}$ denotes the input value of *imin* and $imin_{out}$ denotes its output value.

**Returned Value**

Array, *cumpr*, of length $nmass + nndx$ containing in the first *nmass* positions, the cumulative probabilities and in some of the remaining positions, indexes to speed access to the probabilities.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.
**Cum_Probs** — One dimensional array of length \( nmass \) containing the cumulative probabilities to be used in computing the index portion of the result. If the keyword *Cum_Probs* is used, *prf* is not used and may be a dummy function.

**Discussion**

Routine DISCR_TABLE sets up a table that routine RAND_GEN_CONT (page 219) uses to generate pseudorandom deviates from a discrete distribution. The distribution can be specified either by its probability function *prf* or by a vector of values of the cumulative probability function. Note that *prf* is not the cumulative probability distribution function. If the cumulative probabilities are already available in *Cum_Probs*, the only reason to call DISCR_TABLE is to form an index vector in the upper portion of the result so as to speed up the generation of random deviates by the routine RAND_GEN_CONT.

**Example 1**

In this example, DISCR_TABLE is used to set up a table to generate pseudorandom variates from the discrete distribution:

\[
\begin{align*}
Pr(X = 1) &= .05 \\
Pr(X = 2) &= .45 \\
Pr(X = 3) &= .31 \\
Pr(X = 4) &= .04 \\
Pr(X = 5) &= .15
\end{align*}
\]

In this simple example, we input the cumulative probabilities directly using keyword *Cum_Probs* and request 3 indexes to be computed \((nndx = 4)\). Since the number of mass points is so small, the indexes would not have much effect on the speed of the generation of the random variates.

```plaintext
function PRF, x
  return, 0
end
cum_probs = [.05, .5, .81, .85, 1]
cumpr = DISCR_TABLE('PRF', 0.00001, 4, 1, 5, cum_probs = cum_probs)
PM, cumpr
  0.0500000
```
Example 2

This example, DISCR_TABLE is used to set up a table to generate binomial variates with parameters 20 and 0.5. The routine BINOMIALPDF (Chapter 11, Probability Distribution and Inverses) is used to compute the probabilities.

```
FUNCTION PRF, ix
    RETURN, BINOMIALPDF(ix, 20, .5)
END
```

cumpr = DISCR_TABLE('PRF', 0.00001, 12, 0, 21)

PM, cumpr

```
1.90735e-05
0.000200272
0.00128746
0.00590802
0.0206938
0.0576583
0.131587
0.251722
0.411901
0.588099
0.748278
0.868413
0.942342
0.979306
0.994092
```
**RAND_GEN_DISCR Function**

Generates pseudorandom numbers from a general discrete distribution using an alias method or optionally a table lookup method.

**Usage**

\[
\text{result} = \text{RAND_GEN_DISCR}(n, \text{imin}, \text{nmass}, \text{probs})
\]

**Input Parameters**

- **n** — Number of random numbers to generate.
- **imin** — Smallest value the random deviate can assume. This is the value corresponding to the probability in \(\text{probs}(0)\).
- **nmass** — Number of mass points in the discrete distribution.
- **probs** — Array of length \(\text{nmass}\) containing probabilities associated with the individual mass points. The elements of \(\text{probs}\) must be nonnegative and must sum to 1.0.
If the keyword *Table* is used, then *probs* is a vector of length at least *nmass* + 1 containing in the first *nmass* positions the cumulative probabilities and, possibly, indexes to speed access to the probabilities. Routine DISCR_TABLE (page 221) can be used to initialize *probs* properly. If no elements of *probs* are used as indexes, *probs* (*nmass*) is 0.0 on input. The value in *probs*(0) is the probability of *imin*. The value in *probs* (*nmass*-1) must be exactly 1.0 (since this is the CDF at the upper range of the distribution.)

**Returned Value**

An integer array of length *n* containing the random discrete deviates.

**Input Keywords**

*Double* — If present and nonzero, double precision is used.

*Table* — If present and nonzero, generate pseudorandom numbers from a general discrete distribution using a table lookup method. If this keyword is used, then *probs* is a vector of length at least *nmass* + 1 containing in the first *nmass* positions the cumulative probabilities and, possibly, indexes to speed access to the probabilities. Routine DISCR_TABLE (page 221) can be used to initialize *probs* properly.

**Discussion**

Routine RAND_GEN_DISCR generates pseudorandom numbers from a discrete distribution with probability function given in the vector *probs*; that is

\[
Pr(X = i) = p_j
\]

for \(i = i_{\text{min}}, i_{\text{min}} + 1, \ldots, i_{\text{min}} + n_m - 1\) where \(j = i - i_{\text{min}} + 1, p_j = \text{probs}(j)\),

\(i_{\text{min}} = \text{imin}\), and \(n_m = \text{nmass}\).

The algorithm is the *alias* method, due to Walker (1974), with modifications suggested by Kronmal and Peterson (1979).

If the keyword *Table* is used, RAND_GEN_DISCR generates pseudorandom deviates from a discrete distribution, using the table *probs*, which contains the cumulative probabilities of the distribution and, possibly, indexes to speed the search of the table. The DISCR_TABLE (page 221) can be used to set up the table *probs*. RAND_GEN_DISCR uses the inverse CDF method to generate the variates.
Example 1

In this example, RAND_GEN_DISCR is used to generate five pseudorandom variates from the discrete distribution:

\[ Pr(X = 1) = 0.05 \]
\[ Pr(X = 2) = 0.45 \]
\[ Pr(X = 3) = 0.31 \]
\[ Pr(X = 4) = 0.04 \]
\[ Pr(X = 5) = 0.15 \]

probs = [.05, .45, .31, .04, .15]
n = 5
imin = 1
nmass = 5
RANDOMOPT, Set_seed = 123457
r = RAND_GEN_DISCR(n, imin, nmass, probs)
PM, r

3
2
2
3
5

Example 2

In this example, DISCR_TABLE (page 221) is used to set up a table and then RAND_GEN_DISCR is used to generate five pseudorandom variates from the binomial distribution with parameters 20 and 0.5.

FUNCTION PRF, ix
   RETURN, BINOMIALPDF(ix, 20, .5)
END
imin = 0
nmass = 21
RANDOMOPT, Set_seed = 123457
cumpr = DISCR_TABLE(‘prf’, 0.00001, 12, imin, nmass)
\[ r = \text{RAND\_GEN\_DISCR}(n, \text{imin}, \text{nmass, cumpr}, /\text{table}) \]

PM, \( r \)

\begin{verbatim}
  14
  9
  12
  10
  12
\end{verbatim}

### RANDOM_ARMA Function

Generates a time series from a specific ARMA model.

**Usage**

\[
\begin{align*}
\text{result} &= \text{RANDOM\_ARMA}(n, \text{nparams}) \\
\text{result} &= \text{RANDOM\_ARMA}(n, \text{nparams, ar}) \\
\text{result} &= \text{RANDOM\_ARMA}(n, \text{nparams, ma}) \\
\text{result} &= \text{RANDOM\_ARMA}(n, \text{nparams, ar, ma})
\end{align*}
\]

**Input Parameters**

- \( n \) — Number of observations to be generated. Parameter \( n \) must be greater than or equal to one.
- \( \text{nparams} \) — One-dimensional array containing the parameters \( p \) and \( q \) consecutively. \( \text{nparams}(0) = p \), where \( p \) is the number of autoregressive parameters. Parameter \( p \) must be greater than or equal to zero. \( \text{nparams}(1) = q \), where \( q \) is the number of moving average parameters. Parameter \( q \) must be greater than or equal to zero.
- \( \text{ar} \) — One-dimensional array of length \( p \) containing the autoregressive parameters.
- \( \text{ma} \) — One-dimensional array of length \( q \) containing the moving average parameters.

**Returned Value**

- \( \text{result} \) — One-dimensional array of length \( n \) containing the generated time series.
Input Keywords

**Double** — If present and nonzero, double precision is used.

**Const** — Overall constant. See the Discussion section.

Default: \( Const = 0 \)

**Var_Noise** — If present (and **Input_Noise** is not used), the noise \( a_t \) will be generated from a normal distribution with mean 0 and variance \( Var_Noise \). Keywords **Var_Noise** and **Input_Noise** can not be used together.

Default: \( Var_Noise = 1.0 \)

**Input_Noise** — One-dimensional array of length \( n + \max(Ar_Lags(i)) \) containing the random noises. Keywords **Input_Noise** and **Var_Noise** can not be used together. Keywords **Input_Noise** and **Output_Noise** can not be used together.

**Ar_Lags** — One-dimensional array of length \( p \) containing the order of the non-zero autoregressive parameters.

Default: \( Ar_Lags = [1, 2, \ldots, p] \)

**Ma_Lags** — One-dimensional array of length \( q \) containing the order of the non-zero moving average parameters.

Default: \( Ma_Lags = [1, 2, \ldots, q] \)

**W_Init** — One-dimensional array of length \( \max(Ar_Lags(i)) \) containing the initial values of the time series.

Default: \( W_Init(*) = Const/(1 - ar(0) - ar(1) - \ldots - ar(p - 1)) \)

**Accept_Reject** — If present and nonzero, the random noises will be generated from a normal distribution using an acceptance/rejection method. If keyword **Accept_Reject** is not used, the random noises will be generated using an inverse normal CDF method. This argument will be ignored if keyword **Input_Noise** is used.

Output Keywords

**Output_Noise** — Named variable into which a one-dimensional array of length \( n + \max(Ma_Lags(i)) \) containing the random noises is stored.

Discussion

Function RANDOM_ARMA simulates an ARMA\((p, q)\) process, \( \{W_t\} \), for
\[ t = 1, 2, \ldots, n. \text{ The model is} \]
\[ \phi(B)W_t = \theta_0 + \theta(B)A_t \quad t \in \mathbb{Z} \]
\[ \phi(B) = 1 - \phi_1 B - \phi_2 B^2 - \ldots - \phi_p B^p \]
\[ \theta(B) = 1 - \theta_1 B - \theta_2 B^2 - \ldots - \theta_q B^q \]

Let \( \mu \) be the mean of the time series \( \{W_t\} \). The overall constant \( \theta_0 \) (\text{Const}) is
\[ \theta_0 = \begin{cases} \mu & p = 0 \\ \mu(1 - \sum_{i=1}^{p} \phi_i) & p > 0 \end{cases} \]

Time series whose innovations have a nonnormal distribution may be simulated by providing the appropriate innovations in \text{Input\_Noise} and start values in \text{W\_Init}.

The time series is generated according to the following model:
\[
X(i) = \text{Const} + \text{ar}(0) \times X(i - \text{Ar\_Lags}(0)) + \ldots + \\
\text{ar}(p - 1) \times X(i - \text{Ar\_Lags}(p - 1)) + \\
\text{A}(1) - \text{ma}(0) \times A(i - \text{Ma\_Lags}(0)) - \ldots - \\
\text{ma}(q - 1) \times A(i - \text{Ma\_Lags}(q - 1))
\]
where the constant is related to the mean of the series, \( \bar{W} \)
\[ \bar{W} \]
as follows:
\[ \text{Const} = \bar{W} - (1 - \text{ar}(0) - \ldots - \text{ar}(q - 1)) \]

\[ X(t) = W(t), \quad t = 0, 1, \ldots, n - 1 \]
and
\[ W(t) = \text{W\_Init}(t + p), \quad t = -p, -p + 1, \ldots, -2, -1 \]
and A is either Input_Noise (if Input_Noise is used) or Output_Noise (otherwise).

**Example 1**

In this example, RANDOM_ARMA is used to generate a time series of length five, using an ARMA model with three autoregressive parameters and two moving average parameters. The start values are 0.1000, 0.0500, and 0.0375.

```plaintext
RANDOMOPT, set = 123457
n = 5
nparams = [3, 2]
ar = [0.5, 0.25, 0.125]
ma = [-0.5, -0.25]
r = RANDOM_ARMA(n, nparams, ar, ma)
PM, r, Format = "(5F10.3)", $
Title = "ARMA random deviates"
   ARMA random deviates
   0.637  0.317 -0.366 -2.122 -1.407
```

**Example 2**

In this example, a time series of length 5 is generated using an ARMA model with 4 autoregressive parameters and 2 moving average parameters. The start values are 0.1, 0.05 and 0.0375.

```plaintext
RANDOMOPT, set = 123457
n = 5
nparams = [3, 2]
ar = [0.5, 0.25, 0.125]
ma = [-0.5, -0.25]
winit = [0.1, 0.05, 0.0375]
theta0 = 1
avar = 0.1
r = RANDOM_ARMA(n, nparams, ar, ma, Accept_Reject, $
   W_Init = wi, Const = theta0, $
   Var_Noise = avar)
PM, r, Format = "(5F10.3)", $
```

---

230 Chapter 2: New Commands PV-WAVE 7.5 New Features Guide
Title = " ARMA random deviates:

ARMA random deviates:
1.467   1.788   2.459   3.330   3.941

Warning Errors
STAT_RNARM_NEG_VAR — VAR(a) = “Var_Noise” = #, VAR(a) must be greater than 0. The absolute value of # is used for VAR(a).

**FAURE_INIT Function**

Initializes the structure used for computing a shuffled Faure sequence.

**Usage**

\[
result = \text{FAURE\_INIT}(ndim)
\]

**Input Parameters**

- **ndim** — The dimension of the hyper-rectangle.

**Returned Value**

A structure that contains information about the sequence.

**Input Keywords**

**Base** — The base of the Faure sequence.
- Default: The smallest prime greater than or equal to \( ndim \).

**Skip** — The number of points to be skipped at the beginning of the Faure sequence. Default:

\[
\left\lfloor \frac{m}{2} \right\rfloor
\]

where

\[
m = \left\lfloor \log B / \log base \right\rfloor
\]
and $B$ is the largest representable integer.

**Discussion**

Discrepancy measures the deviation from uniformity of a point set.

The discrepancy of the point set

$$x_1, \ldots, x_n \in [0,1]^d, \ d \geq 1,$$

is

$$D_n^{(d)} = \sup \left\{ \frac{\lambda(E;n)}{n} - \lambda(E) \right\},$$

where the supremum is over all subsets of $[0,1]^d$ of the form

$$E = \left[ 0, t_1 \right) \times \ldots \times \left[ 0, t_d \right), \ 0 \leq t_j \leq 1, \ 1 \leq j \leq d,$$

$\lambda$ is the Lebesgue measure, and

$$\left( E; n \right)$$

is the number of the $x_j$ contained in $E$.

The sequence $x_1, x_2, \ldots$ of points $[0,1]^d$ is a low-discrepancy sequence if there exists a constant $c(d)$, depending only on $d$, such that

$$D_n^{(d)} \leq c(d) \left( \frac{\log n}{n} \right)^d$$

for all $n > 1$.

Generalized Faure sequences can be defined for any prime base $b \geq d$. The lowest bound for the discrepancy is obtained for the smallest prime $b \geq d$, so the keyword `Base` defaults to the smallest prime greater than or equal to the dimension.

The generalized Faure sequence $x_1, x_2, \ldots$, is computed as follows:
Write the positive integer $n$ in its $b$-ary expansion,

$$n = \sum_{i=0}^{\infty} a_i(n)b^i$$

where $a_i(n)$ are integers,

$$0 \leq a_i(n) < b$$

The $j$-th coordinate of $x_n$ is

$$x_n^{(j)} = \sum_{k=0}^{\infty} \sum_{d=0}^{\infty} c_{kd}^{(j)} a_d(n) b^{-k-1}, \quad 1 \leq j \leq d$$

The generator matrix for the series,

$$c_{kd}^{(j)}$$

is defined to be

$$c_{kd}^{(j)} = j^d - k^c_{kd}$$

and

$$c_{kd}$$

is an element of the Pascal matrix,

$$c_{kd} = \begin{cases} 
\frac{d!}{c!(d-c)!} & k \leq d \\
0 & k > d 
\end{cases}$$

It is faster to compute a shuffled Faure sequence than to compute the Faure sequence itself. It can be shown that this shuffling preserves the low-discrepancy property.

The shuffling used is the $b$-ary Gray code. The function $G(n)$ maps the positive integer $n$ into the integer given by its $b$-ary expansion.
The sequence computed by this function is \( x(G(n)) \), where \( x \) is the generalized Faure sequence.

**Example**

In this example, five points in the Faure sequence are computed. The points are in the three-dimensional unit cube.

Note that \( \text{FAURE\_INIT} \) is used to create a structure that holds the state of the sequence. Each call to \( \text{FAURE\_NEXT\_PT} \) returns the next point in the sequence and updates the state structure.

```c
state = \text{FAURE\_INIT}(3)
p = \text{FAURE\_NEXT\_PT}(5, \text{state})
```

<table>
<thead>
<tr>
<th>PM</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.333689</td>
<td>0.492659</td>
</tr>
<tr>
<td>0.667022</td>
<td>0.825992</td>
</tr>
<tr>
<td>0.778133</td>
<td>0.270436</td>
</tr>
<tr>
<td>0.111467</td>
<td>0.603770</td>
</tr>
<tr>
<td>0.444800</td>
<td>0.937103</td>
</tr>
</tbody>
</table>

---

**FAURE\_NEXT\_PT Function**

Computes a shuffled Faure sequence.

**Usage**

\( \text{result} = \text{FAURE\_NEXT\_PT}(\text{npts}, \text{state}) \)

**Input Parameters**

- \( \text{npts} \) — The number of points to generate in the hyper-rectangle.
- \( \text{state} \) — State structure created by a call to \( \text{FAURE\_INIT} \).

**Returned Value**

An array of size \( \text{npts} \) by \( \text{state\_dim} \) containing the \( \text{npts} \) next points in the shuffled Faure sequence.
**Input Keywords**

*Double* — If present and nonzero, double precision is used.

**Output Keywords**

*Skip* — The current point in the sequence. The sequence can be restarted by initializing a new sequence using this value for *Skip*, and using the same dimension for *ndim*.

**Discussion**

Discrepancy measures the deviation from uniformity of a point set.

The discrepancy of the point set

\[ x_1, ..., x_n \in [0,1]^d, d \geq 1, \]

is

\[ D_n^{(d)} = \sup_{E} \left| \frac{A(E;n)}{n} - \lambda(E) \right|, \]

where the supremum is over all subsets of \([0,1]^d\) of the form

\[ E = \left[ 0, t_1 \right] \times \cdots \times \left[ 0, t_j \right], 0 \leq t_j \leq 1, 1 \leq j \leq d, \]

\( \lambda \) is the Lebesque measure, and

\( (E;n) \)

is the number of the \( x_j \) contained in \( E \).

The sequence \( x_1, x_2, \ldots \) of points \([0,1]^d\) is a low-discrepancy sequence if there exists a constant \( c(d) \), depending only on \( d \), such that

\[ D_n^{(d)} \leq c(d) \left( \frac{\log n}{n} \right)^d \]

for all \( n > 1 \).
Generalized Faure sequences can be defined for any prime base \( b \geq d \). The lowest bound for the discrepancy is obtained for the smallest prime \( b \geq d \), so the keyword Base defaults to the smallest prime greater than or equal to the dimension.

The generalized Faure sequence \( x_1, x_2, \ldots \), is computed as follows:

Write the positive integer \( n \) in its \( b \)-ary expansion

\[
n = \sum_{i=0}^{\infty} a_i(n) b^i
\]

where \( a_i(n) \) are integers,

\[
0 \leq a_i(n) < b
\]

The \( j \)-th coordinate of \( x_n \) is

\[
x_n^{(j)} = \sum_{k=0}^{\infty} \sum_{d=0}^{\infty} c_k^{(j)} a_d(n) b^{-k-1}, \quad 1 \leq j \leq d
\]

The generator matrix for the series,

\[
c_k^{(j)}
\]

is defined to be

\[
c_k^{(j)} = j d - k c_k^{(j)}
\]

and

\[
c_k^{(j)}
\]

is an element of the Pascal matrix,

\[
c_{k,d} = \begin{cases} 
  d! & k \leq d \\
  \frac{d!(d-c)!}{c!} & k > d \\
  0 & k > d
\end{cases}
\]
It is faster to compute a shuffled Faure sequence than to compute the Faure sequence itself. It can be shown that this shuffling preserves the low-discrepancy property.

The shuffling used is the \textit{b-ary} Gray code. The function $G(n)$ maps the positive integer $n$ into the integer given by its \textit{b-ary} expansion.

The sequence computed by this function is $x(G(n))$, where $x$ is the generalized Faure sequence.

\textbf{Example}

In this example, five points in the Faure sequence are computed. The points are in the three-dimensional unit cube.

Note that \texttt{FAURE\_INIT} is used to create a structure that holds the state of the sequence. Each call to \texttt{FAURE\_NEXT\_PT} returns the next point in the sequence and updates the state structure.

\begin{verbatim}
state = FAURE\_INIT(3)
p = FAURE\_NEXT\_PT(5, state)
PM, p
0.333689  0.492659  0.0640654
0.667022  0.825992  0.397399
0.778133  0.270436  0.175177
0.111467  0.603770  0.508510
0.444800  0.937103  0.841843
\end{verbatim}
Updates to Existing Functionality

This chapter discusses the following topics:

- **Updated PV-WAVE Functions and Procedures** — Descriptions of new keywords and parameters that have been added to PV-WAVE functions and procedures. (page 239)

- **Updated PV-WAVE:IMSL Statistics Functions and Procedures** — Description of new functionality that has been added to PV-WAVE:IMSL Statistics functions and procedures. (page 241)

- **Updated PV-WAVE:IMSL Mathematics Functions and Procedures** — Description of new functionality that has been added to PV-WAVE:IMSL Mathematics functions and procedures. (page 242)

---

**Updated PV-WAVE Functions and Procedures**

This section describes new keywords and parameters that have been added to PV-WAVE functions and procedures.

---

**JOURNAL Procedure**

Keywords

* Nobuffer — If present and nonzero, output lines should be written immediately to the journal file without the normal file buffering.**
NoBlock Keyword for PV-WAVE VDA Tool Procedures

A new keyword, NoBlock, has been added for all standard VDA tool procedures (see the PV-WAVE Reference).

Keywords

NoBlock — If specified, the event loop (WwLoop) that is started by the VDA Tool will use the given value. By default a value of 1 (non-blocking loop) will be used. However, it may be necessary in certain circumstances to force a non-blocking loop by specifying NoBlock = 2. (See WwLoop in the PV-WAVE Application Developer’s Guide for more information.)

YLabelCenter Keyword

Used With Routines: AXIS, BAR, BAR2D, BAR3D, CONTOUR, CONTOUR2, OPLot, PLOT, PLOT FIELD, SURFACE

Corresponding System Variable: None.

Controls whether the top and bottom major tick labels on a Y axis will be positioned within the boundaries of the axis box or centered across from the corresponding major tick.

If this keyword is set, the top and bottom Y axis major tick labels will be centered vertically with corresponding major ticks. If this keyword is not set, the default behavior is to position the top and bottom Y axis major tick labels within the boundaries of the axis box.

!Version System Variable

The !Version system variable now has a field called Revision. !Version.Revision will either be a null string for major PV-WAVE releases, e.g. 7.00, 7.01, 7.10, or a letter denoting the patch revision level, e.g. 'a', 'b', etc. Printing the complete version number for PV-WAVE is now:

PRINT, !Version.release + !Version.revision
Updated PV-WAVE:IMSL Statistics Functions

This section describes new features that has been added to PV-WAVE:IMSL Statistics functions. For details, see the PV-WAVE: IMSL Statistics Reference.

RANDOM Function

Added several new distributions, including multinomial and stable. Also added support for random permutations and samples.

RANDOMOPT Function

Added support for generating substream seeds, and support for a Generalized Feedback Shift Register (GFSR) generator.
Updated PV-WAVE:IMSL Mathematics Functions

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Added several new distributions, including multinomial and stable. Also added support for random permutations and samples.

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Index

A

AFFINE function 5
array
  shift along one dimension 43
  subset 44
arrays
  compute logical AND 24
  expanding into higher dimensions 14
  extrema 17
  homogeneous region, counting 7
  homogeneous region, isolating 6
  OR together 25
  resample to new dimensions 40
  sorting contents of 46
  transformation 5

B

binomial distributions 210, 212, 213, 216, 221, 225
BLOB function 6
BLOBCOUNT function 7
bond functions
  accrued interest maturity 131
  accrued interest period 133
  bond equivalent yield 135
  convexity 137
  coupon date 163, 165
  coupon days 139
  coupon days - next 145
  coupon numbers 141
  depreciation accounting period 147, 148
  discount price 150
  discount rate 152
  discount yield 154
  duration 157
  interest rate security 159
  Macaulay duration 161
  maturity received 172
  price maturity 170
  price periodic 167
  settlement db 143
tbill price 174
tbill yield 175
  year fraction 176
  yield maturity 178
  yield periodic 180
boundary 8
BOUNDARY function 8

C

Cartesian product 9
compare 41
converted 66
correlation matrix 212, 216
CPROD function 9
CURVATURES function 10

D

DERIVN function 11
DICOM 12
differentiation 11
dimensions of array, expanding 14
distribution functions
  binomial probability 200
divisor 19

E

EUCLIDEAN function 13
expand an array 14
EXPAND function 14
EXPON function 16
exponentiation 16
EXTREMA function 17

F

factor 18
FACTOR function 18
Faure 185, 188, 233, 237
Faure sequence 184, 187, 232, 235
faure_next_point 187, 235
financial functions
cumulative interest 97
cumulative principal 99
depreciation
double-declining balance 102
straight-line 104
sum-of-years 105
variable declining balance 106
dollar decimal 108
dollar fraction 109
effective rate 110
future value 111
future value schedule 113
interest payment 114
interest rate annuity 115
internal rate of return 117
internal rate of return schedule 118
modified internal rate of return 120
net present value 121
nominal rate 123
number of periods 124
payment 125
present value 127
present value schedule 128
principal payment 130
floor 20
G
GCD function 19
general discrete distribution 95, 200, 209,
210, 213, 218, 219, 221, 225, 226
Gray code 186, 190, 234, 238
GREAT_INT function 20
greatest integer 20
gird 21
GRIDN function 21
H
HISTN function 22
histogram 22
hyper-rectangle 95
I
images
24-bit, rendering 38
INDEX_AND function 24
INDEX_OR function 25
integration 95
INTERPOLATE function 26
interpolation 27
scattered data 26
INTRP function 27
J
JACOBIAN function 28
K
Kalman filtering 190
L
LCM function 29
least integer 45
Lebesque measure 185, 188, 233, 236
list
returning 30
LISTARR function 30
low-discrepancy 186, 189, 234, 238
M
mathematical function
boundary 8
common divisor 19
compare 41
exponentiation 16
greatest integer 20
histogram 22
least common multiple 29
least integer 45
minimize 31
moments 33
multiply 37
neighbors 34
prime factorization 18
primes 36
resample array 40
sign of passed values 42
subset an array 44
vector, replicate 39
mathematics. See PV-WAVE:IMSL
Mathematics
maximum likelihood estimates 197
minimize 31
MINIMIZE function 31
MOLEC function 32
molecular model 32
MOMENT function 33
moments 33
multiple 29
multiply 37
multivariate distribution 216

N
neighbors 34
NEIGHBORS function 34
NORMALS function 35

P
PRIME function 36
prime numbers 36
printing
  Windows 90
product 37
PRODUCT function 37
pseudorandom numbers 205, 216, 218, 219, 221, 225, 226
pseudorandom order statistics 209
pseudorandom orthogonal matrix 212
pseudorandom sample 213
PV-WAVE
  Database Connection
    new routines 91
PV-WAVE:IMSL Mathematics
  basic statistics 184
  new functions 95
  quadrature 95
  special functions 97
    updated functions 242
PV-WAVE:IMSL Statistics
  new routines 190
  probability distribution functions and inverses 200
  regression 201
  time series and forecasting 190
    updated functions 241
Q
quasi-Monte Carlo 95
R
RENDER24 function 38
REPLV function 39
RESAMP function 40
S
SAME function 41
SGN function 42
SHIF function 43
sign of passed values 42
SLICE function 44
SMALL_INT function 45
sorting
  array contents 46
SORTN function 46
state vector 191
statespace model 190
statistics. See PV-WAVE:IMSL Statistics
T
tick marks
  centering Y label 240
transformation
  affine 5
24-bit image data
  rendering 38
V
vector
  replicate 39
vtkADDATTRIBUTE Procedure 47
vtkAXES Procedure 48
vtkCAMERA Procedure 50
vtkCLOSE Procedure 51
vtkCOLORBAR Procedure 52
vtkCOMMAND Procedure 53
vtkERASE Procedure 54
vtkGRID Procedure 55
vtkHEDGEHOG Procedure 56
vtkINIT Procedure 58
vtkLIGHT Procedure 59
vtkPLOTS Procedure 60
vtkPOLYDATA Procedure 62
vtkPOLYSHADE Procedure 63
vtkPPMREAD Function 65
vtkPPMWRITE Procedure 66
vtkRECTILINEARGRID Procedure 67
vtkRENDERWINDOW Procedure 68
vtkSCATTER Procedure 70
vtkSLICEVOL Procedure 72
vtkSTRUCTUREDGRID Procedure 74
vtkSTRUCTUREDPOINTS Procedure 75
vtkSURFACE Procedure 76
vtkSURFGEN Procedure 79
vtkTEXT Procedure 81
vtkTVRD Function 82
vtkUNSTRUCTUREDGRID Procedure 83
vtkWDELETE Procedure 84
vtkWINDOW Procedure 85
vtkWRITEVRML Procedure 87
vtkWSET Procedure 88

W

WgOrbit procedure 89
WIN32_PICK_PRINTER function 90
Windows
   printer dialog 90