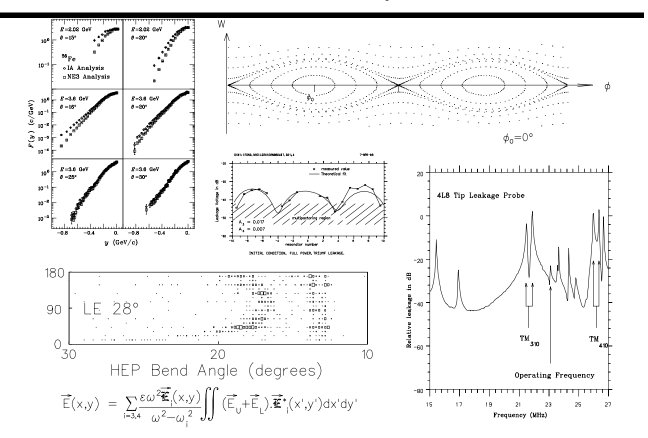
| TRIUMF | 4004 Wesbrook Mall, Vancouver, B.C., Canada V6T 2A3 | | | |
|---|---|--------------|--------------|------|
| Computing Document | J.L. Chuma | January 1998 | TRI-CD-93-01 | v1.3 |
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PHYSICA ® REFERENCE MANUAL

Mathematical Analysis and Data Visualization Software



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Contents

| 1 | INTRODUCTION | 1 |
|---|----------------------------|----|
| | 1.1 What is in this manual | 1 |
| 2 | COMMANDS | 4 |
| | 3DPLOT | 4 |
| | ALIAS | 5 |
| | ASSIGN | 5 |
| | BESTFIT | 6 |
| | Parameter types and sizes | 6 |
| | Weights | 6 |
| | Cycles | 7 |
| | BIN | 7 |
| | Weights | 7 |
| | Number of bins | 8 |
| | Lagrange | 8 |
| | Averages | 8 |
| | Increment only if empty | 9 |
| | Edge defined bins | 9 |
| | BIN2D | 9 |
| | Dimensions | 10 |
| | Extremes | 10 |
| | Weights | 10 |
| | Increment only if empty | 10 |
| | Defined by box corners | 10 |
| | BUFFER | 11 |
| | Parameters | 11 |
| | Reading the buffers | 11 |
| | Writing the buffers | 12 |

| Dynamic buffer | 1 | 2 |
|---------------------------------------|---|-----|
| Static buffer | 1 | 2 |
| Keypad buffer | 1 | 2 |
| CALL | 1 | 4 |
| User written subroutine description | 1 | 4 |
| Creating a shareable image | | 8 |
| CLEAR | | 9 |
| Alphanumerics | 1 | 9 |
| Toggle graphics | 1 | 9 |
| Clear the replot buffers only | | 9 |
| Do not clear the replot buffers | | 9 |
| COLOUR | | |
| Using a scalar for a colour number | | |
| CONTOUR | | |
| Contour level selection | | |
| Contour level colour | | |
| Contour labels | | |
| Saving contour levels and coordinates | | |
| Legend | | |
| Polar coordinates | | |
| Axes | | |
| Scattered points | | |
| Matrix data | | |
| COPY | | |
| Conditional copy | | |
| Unconditional copy | | |
| Appending with copy | | |
| DCL | | |
| UNIX equivalent | | |
| DEALIAS | | |
| DEFAULTS | | |
| Initialization file | | |
| | | |
| Reset windows | | |
| Default values | | |
| DENSITY | | |
| Density plot types | | |
| Axes | | |
| Matrix boundary | | |
| Zooming in | | |
| Derivatives | | |
| Profiles | | |
| Polar coordinates | | |
| Solid filled regions | 3 | ٠:۱ |

| | ndom points | 34 |
|-------|-----------------------------|----|
|] | fusion | 36 |
|] | hering patterns | 37 |
|] | xes | 40 |
| DEST | OY | 44 |
| Į | conditional | 44 |
| | nditional | |
| | | |
| (| and OFF | |
| | vice keywords | |
| | LaserJet | |
|] | Jet | |
| | ner bitmap devices | |
| | stScript devices | |
| | n plotters | |
| | ner vector plotters | |
| | S metafiles | |
| | splay files | |
| | Е | |
| | gitizing pad types | |
| | tional output variables | |
| | eparing for digitizing data | |
| | gitizing data | |
| | E | |
| | aphics window borders | |
| | oadcast messages | |
| | nfirmation requests | |
| | hoing from scripts | |
| | ving a variable's history | |
| | ırnaling input and output | |
| | ompting | |
|] | plotting | |
| | Window graphics replay | 57 |
| | out line recall shell | 58 |
| | cking commands in a file | 58 |
| DISPL | | 58 |
| | splay a message | 58 |
| | nt table | 59 |
| | ecial characters | 59 |
| | tch fill patterns | 59 |
| | te types | 59 |
| | tting symbols | 62 |
| | nus | 62 |

| EDGR | 69 |
|---|----|
| Open a drawing file | 69 |
| Edit a drawing file | 69 |
| Close a drawing file | 69 |
| Open a new frame | 70 |
| ELLIPSE | 70 |
| Output vectors | 70 |
| Replotting | 70 |
| Number of points | 71 |
| Explicitly defined | 71 |
| Fit an ellipse | 71 |
| ENABLE | 73 |
| Graphics window borders | 73 |
| Broadcast messages | 73 |
| Confirmation requests | 74 |
| Echoing from scripts | 74 |
| Saving a variable's history | 74 |
| Journaling input and output | 74 |
| Prompting | 75 |
| Replotting | 75 |
| X Window graphics replay | 75 |
| Input line recall shell | 75 |
| Stacking commands in a file | 76 |
| ERASEWINDOW | 76 |
| EXECUTE | 76 |
| Environment variables in file names | 77 |
| Filename extensions | 77 |
| script library | 77 |
| Comments | 77 |
| Echoing | 78 |
| Temporarily passing control to the keyboard | |
| Returning from a script | 78 |
| Aborting a script | |
| Passing parameters to a script | 78 |
| Prompting | 79 |
| Labels and GOTOs | 79 |
| DO loops | 79 |
| Conditional statements | 80 |
| EXTENSION | 81 |
| FIGURE | 82 |
| Line types | 82 |
| Fillable figures | 82 |
| X Windows | 83 |

| Units | 83 |
|---|----|
| Confirmation | 8 |
| Stack file | 8 |
| Circle | 8 |
| Arc | 84 |
| Wedge | 84 |
| Box | 8 |
| Polygon | 8 |
| Ellipse | 8 |
| Arrow | 8 |
| FILTER | 80 |
| Noise amplification caused by filtering | 8 |
| Median filter | 8 |
| Mean filter | 8' |
| Nonrecursive filters | |
| Recursive filters | 89 |
| FIT | 93 |
| Expression and parameters | 9 |
| Method | 9 |
| Normal distribution | 90 |
| Poisson distribution | 98 |
| Correlation and covariance | 99 |
| Confidence level of the fit | 10 |
| Number of iterations | 10 |
| Informational messages | 10 |
| Update after a fit | 10 |
| FMIN | 10 |
| Informational messages | 10 |
| FZERO | 10 |
| Muller's method | |
| Informational messages | |
| GENERATE | 10 |
| Increment and number of points given | |
| Maximum and number of points given | |
| Increment and maximum given | |
| Random numbers | 10 |
| GET | |
| The GPLOT keywords | |
| The PHYSICA keywords | |
| GLOBALS | |
| GRAPH | |
| Plotting symbols | |
| Axis scaling | 11 |

| Gr | h legend |
|---------------|---------------------------------------|
| Plo | ng data and axes |
| Plo | ng axes only |
| Plo | ng data only |
| Re | otting data on a common scale |
| His | grams |
| Po | coordinates |
| Er | bars |
| Fil | g |
| GRID . | |
| Po | coordinates |
| Dι | icate points |
| Int | polated grid |
| No | interpolated grid |
| Ma | ix from sparse data |
| HARDC | Υ |
| Pri | ing and saving |
| HELP . | |
| Pa | ng the output |
| Us | defined library |
| INPUT. | |
| Ve | ors |
| Ma | ix |
| INQUIR | |
| JOURN | |
| En | conment variables in file names |
| KEYWO | 0 |
| LABEL. | |
| Tu | ing off the labels |
| LEGENI | |
| Th | string portion of a legend entry |
| Th | ine segment portion of a legend entry |
| Th | rame box |
| Tra | sparency |
| Th | egend title |
| Sta | ıs |
| LINE | |
| Plo | ng units |
| X | ndows 14 |
| No | interactive drawing |
| LIST | |
| Pa | ng the output |
| | ng a matrix |

| LOAD | . 146 |
|--|-------|
| Restrictions | . 147 |
| Arguments | . 147 |
| Subroutines | |
| Subroutine example | |
| Functions | |
| Function example | |
| MAP | |
| HBOOK | |
| FIOWA | |
| YBOS | |
| MATRIX | |
| MONITOR | |
| Disabling/enabling graphics monitor output | |
| The generic terminal driver | |
| NEWS | |
| ORIENTATION | |
| The world coordinate system | |
| PEAK | |
| Choosing the data curve | |
| X Windows | |
| Code keys | |
| PICK | |
| X Windows | |
| Specifying the number of points | |
| Code keys | |
| Choosing the vertices of a polygon | |
| Matrices | |
| Determining regional counts for data sets | |
| PIEGRAPH | |
| | |
| Pie wedge filling | |
| PLOTTEXT | |
| Comments | |
| Command delimiters | |
| Continuation Lines | |
| Inserting a blank line | |
| Character height | |
| Line spacing | |
| Left margin | |
| Bolding | |
| Colour | . 172 |

|] | ont | 74 |
|-------|---|------------|
| (| Centre justification | 74 |
|] | eft justification | 75 |
|] | Eight justification $\dots \dots \dots$ | 76 |
|] | Iorizontal spacing | 77 |
| , | Sub-script mode | 78 |
| , | Super-script mode | 79 |
| , | Slanted mode | 79 |
|] | Iexadecimal mode | 30 |
| 1 | accents | 31 |
| POLY | GON | 32 |
| QUIT | | 33 |
| READ | | 33 |
|] | Environment variables in file names | 35 |
| (| Opening and closing files | 35 |
|] | Leading data into vectors | 36 |
|] | Leading data into scalars | 93 |
|] | Leading data into a matrix | 3 5 |
|] | Reading data into a string variable |)3 |
| REBI | 「 |)5 |
|] | Cebinning vectors |)5 |
|] | Cebinning matrices |) 6 |
| REFR | ESH |)7 |
| RENA | ME |) 8 |
| REPL | OT |) 8 |
| 1 | Vhat is saved |)8 |
| | strings |)9 |
|] | Enable/Disable |)9 |
| 1 | Vindows |)9 |
| | Clearing the graphics |)9 |
|] | Redraw all windows | 10 |
| RESIZ | E | 10 |
| REST | ORE | |
|] | Environment variables in file names | 11 |
|] | HYSICA sessions | 11 |
|] | TOWA data sets | 12 |
| 2 | TFIOWA data sets | 14 |
|] | IBOOK data sets | 14 |
| • | BOS data sets | 18 |
| | SR MUD data sets | 21 |
| | SR data sets | 23 |
|] | uSR data sets | 23 |
| | CHAOS data sets | 24 |

| RETU | RN |
|------|---|
| SAVE | |
| SCAL | AR |
| | Fit parameters |
| | Dummy variables |
| SCAL | ES |
| | Commensurate axis scaling |
| | Labeled tic marks |
| SET | |
| | How the SET command works |
| | GPLOT keywords |
| | The PHYSICA keywords |
| | I |
| | S |
| SORT | |
| | Associated vectors |
| | K |
| | Environment variables in file names $\dots \dots \dots$ |
| | Appending to a stack file |
| | Executing commands while stacking |
| | STICS |
| | nformational messages |
| | 8 |
| | Weights |
| | Definitions |
| | Moments |
| | inear correlation coefficient |
| | JS |
| | ACE |
| | Colour |
| | INAL |
| TEXT | |
| | Confirmation |
| | Stack files |
| | Text characteristics |
| | Justification and location $\dots\dots\dots\dots$ 2 |
| | Text Formats |
| | Replotting text |
| | Drawing the Date and Time |
| | Erasing text \dots 2 |
| | |
| | Bar definitions $\dots\dots\dots\dots\dots$ 3 |
| | String definitions $\ldots\ldots\ldots\ldots$ 2 |
| TIEN | O . |

| | TRANSFORM | 267 |
|---|-------------------------------------|-------------|
| | UNIQUE | 267 |
| | Indices | 268 |
| | USE | 269 |
| | Environment variables in file names | 270 |
| | VECTOR | 270 |
| | VOLUME | 270 |
| | Volume under a matrix | 271 |
| | WAIT | 271 |
| | WINDOW | 271 |
| | What are windows | 272 |
| | Boundaries | 272 |
| | Plotting units | 272 |
| | Defining a new window | 272 |
| | Pre-defined windows | 27 3 |
| | Windows and GPLOT | 27 3 |
| | Multiple window creation | 274 |
| | WORLD | 275 |
| | WRITE | 275 |
| | Environment variables in file names | 276 |
| | Appending to a file | 276 |
| | Formats | 276 |
| | Vectors | 276 |
| | Scalars | 277 |
| | Matrix | 278 |
| | String | 278 |
| | ZEROLINES | 278 |
| | | |
| 3 | OPERATORS | 281 |
| | Boolean operators | 281 |
| | Transpose | |
| | Reflect | |
| | Union | |
| | Intersection | |
| | Append | |
| | Outer product | |
| | Inner product | |
| 4 | FUNCTIONS | 900 |
| 4 | FUNCTIONS | 288 |
| | Element by element functions | 288 |
| | ATANO | 200 |

| ATAN2D | 90 |
|-------------------------------------|------------|
| RAN | 91 |
| ELTIME | 91 |
| DIM | 91 |
| MOD | 92 |
| SIGN | 92 |
| MIN | 93 |
| MAX | |
| Special mathematical functions | 94 |
| Airy's functions | |
| Beta functions | |
| Bessel functions | |
| Binomial coefficient | |
| Chebyshev polynomials | |
| Probability functions | |
| Cosine integral | |
| Sine integral | |
| Dawson's integral | |
| Digamma Psi function | |
| Dilogarithm | |
| Elliptic integrals | |
| Error function | |
| Exponential integrals | |
| Fermi-Dirac function | |
| Fisher's F -distribution function | |
| Fresnel integrals | |
| Gamma function | |
| Hermite polynomials | |
| Hypergeometric function | |
| Jacobi polynomials | |
| Kelvin functions | |
| | 03 803 |
| Legendre functions and polynomials | |
| | $04 \\ 04$ |
| Rademacher function | |
| | 05 05 |
| | 05 805 |
| | |
| | 06 06 |
| 1 0 | 06 |
| 0 1 | 09 |
| 8 | 09 |
| | 09 |
| 110/IB | |

| | UCASE | 310 |
|-----------|--|-------------------|
| | LCASE | 310 |
| | TCASE | 311 |
| | CHAR | 311 |
| | EXPAND | 311 |
| | VARNAME | 312 |
| | VARTYPE | 312 |
| | STRING | |
| | RCHAR | 313 |
| | TRANSLATE | |
| Nui | neric functions with string arguments | |
| | CLEN | |
| | ICHAR | |
| | EQS | |
| | NES | |
| | SUB | |
| | SUP | |
| | INDEX | |
| | EVAL | |
| Niii | meric analysis functions | |
| Ivai | AREA | |
| | DERIV | |
| | INTEGRAL | |
| | GAUSSJ | |
| | INVERSE | |
| | DET | |
| | IDENTITY | |
| | EIGEN | |
| | PFACTORS | |
| | FFT | |
| | IFFT | |
| | CONVOL | |
| | INTERP | |
| | SPLINTERP | |
| | SMOOTH | |
| | SPLSMOOTH | |
| | | |
| | SAVGOL | |
| . | JOIN | |
| rur | nctions that return a variable's characteristics | |
| | EXIST | |
| | | $\frac{336}{336}$ |
| | VLEN | |
| | FIRST | 337 |

| | | LAST | | | | | | | | | | | | | | | | | | | | | | | | | | | | 337 |
|---|------|---------------|------|------|-----|-----|---|---|-------|---|-----|---|---|-----|---|-----|---|---|-----|-----|---|---|-----|---|---|-----|---|---|---|-----|
| | | ICLOSE | | | | | | | | | | | | | | | | | | | | | | | | | | | | 337 |
| | | IEQUAL | | | | | | | | | | | | | | | | | | | | | | | | | | | | 337 |
| | | WHERE | | | | | | | | | | | | | | | | | | | | | | | | | | | | 337 |
| | Sha | pe changing | fund | ctio | ns | | | | | | | | | | | | | | | | | | | | | | | | | 338 |
| | | FOLD | | | | | | | | | | | | | | | | | | | | | | | | | | | | 338 |
| | | UNFOLD | | | | | | | | | | | | | | | | | | | | | | | | | | | | 338 |
| | | ROLL | | | | | | | | | | | | | | | | | | | | | | | | | | | | 339 |
| | | STEP | | | | | | | | | | | | | | | | | | | | | | | | | | | | 340 |
| | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | | oing function | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | | SUM | | | | | | | | | | | | | | | | | | | | | | | | | | | | 342 |
| | | PROD | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | | RSUM | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | | LOOP | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | | LOOI | • • | • • | • • | • • | • | • | • | • | • • | • | • | • • | • | • • | • | • | • • | • • | • | • | • • | • | • | • • | • | • | • | 343 |
| A | GP | LOT KEYWO | RD | S | | | | | | | | | | | | | | | | | | | | | | | | | • | 348 |
| | A. 1 | Summary | | | | | | | | | | | | | | | | | | | | | | | | | | | | 348 |
| | | General . | | | | | | | | | | | | | | | | | | | | | | | | | | | | 348 |
| | | Text | | | | | | | | | | | | | | | | | | | | | | | | | | | | 348 |
| | | x-axis | | | | | | | | | | | | | | | | | | | | | | | | | | | | 349 |
| | | y-axis | | | | | | | | | | | | | | | | | | | | | | | | | | | | 350 |
| | | Axis Box . | | | | | | | | | | | | | | | | | | | | | | | | | | | | 351 |
| | A.2 | General Ch | | | | | | | | | | | | | | | | | | | | | | | | | | | | 352 |
| | | PTYPE | | | | | | | | | | | | | | | | | | | | | | | | | | | | 352 |
| | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | 352 |
| | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | 352 |
| | | COLOUR . | | | | | | | | | | | | | | | | | | | | | | | | | | | | 352 |
| | | NUMBLD | | | | | | | | | | | | | | | | | | | | | | | | | | | | 352 |
| | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | 353 |
| | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | 353 |
| | | CHARA | | | | | | | | | | | | | | | | | | | | | | | | | | | | 353 |
| | | CHARSZ . | | | | | | | | | | | | | | | | | | | | | | | | | | | | 354 |
| | A.3 | Text | | | | | | | | | | | | | | | | | | | | | | | | | | | | 354 |
| | 11.0 | CURSOR . | | | | | | | | | | | | | | | | | | | | | | | | | | | | 354 |
| | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | 354 |
| | | TXTHIT | | | | | | | | | | | | | | | | | | | | | | | | | | | | 355 |
| | | XLOC | | | | | - | • | | - | | - | • | | • | | • | • | | | | - | | • | • | | | - | | 355 |
| | | *** 0.0 | | | | | | | | | | | | | | | | | | | | | | | | | | | | 355 |
| | Λ 1 | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | A.4 | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | | XAXIS | • • | • • | • • | • • | • | • | • | • | | • | • | • • | • | | • | • | • • | • | • | • | • • | • | • | | • | • | • | 355 |

| | XLOG | | • / | • | • | • | • | • | • | • | 357 |
|-------------|----------|------|-----|-------|-------|-------|---|-------|-------|-------|-------|-------|-------|-------|---|-------|---------|
| | NXGRID | | | | | | | | | | | | | | | | 357 |
| | XCROSS | | | | | | | | | | | | | | | | 359 |
| | XZERO . | | | | | | | | | | | | | | | | 359 |
| | XTICTP . | | | | | | | | | | | | | | | | 359 |
| | XTICA . | | | | | | | | | | | | | | | | 360 |
| | NLXINC | | | | | | | | | | | | | | | | 360 |
| | XTICL . | | | | | | | | | | | | | | | | 360 |
| | NSXINC | | | | | | | | | | | | | | | | 360 |
| | XTICS . | | | | | | | | | | | | | | | | 361 |
| | XMAX . | | | | | | | | | | | | | | | | 361 |
| | XVMAX . | | | | | | | | | | | | | | | | 361 |
| | XMIN | | | | | | | | | | | | | | | | 362 |
| | XVMIN . | | | • | | | | | | | | | | | | | 362 |
| | XMOD . | | | | | | | | | | | | | | | | 363 |
| | XOFF | | | | | | | | | | | | | | | | |
| | XLEADZ | | | | | | | | | | | | | | | | 364 |
| | XPAUTO | | | | | | | | | | | | | | | | 364 |
| | XPOW . | | | | | | | | | | | | | | | | 364 |
| | NXDIG . | | | | | | | | | | | | | | | | 365 |
| | NXDEC . | | | | | | | | | | | | | | | | 365 |
| | XNUMSZ | | | | | | | | | | | | | | | | 365 |
| | XNUMA | | | | | | | | | | | | | | | | 365 |
| | XITICA . | | | | | | | | | | | | | | | | 366 |
| | XITICL . | | | | | | | | | | | | | | | | 366 |
| A. 5 | y-axis . | | | | | | | | | | | | | | | | 366 |
| | YAXIS . | | | | | | | | | | | | | | | | 366 |
| | YLABSZ | | | | | | | | | | | | | | | | 368 |
| | YLOG | | | | | | | | | | | | | | | | 368 |
| | NYGRID | | | | | | | | | | | | | | | | 368 |
| | YCROSS | | | | | | | | | | | | | | | | 370 |
| | YZERO . | | | | | | | | | | | | | | | | 370 |
| | YTICTP . | | | | | | | | | | | | | | | | 370 |
| | YTICA . | | | | | | | | | | | | | | | | 370 |
| | NLYINC | | | | | | | | | | | | | | | | 371 |
| | YTICL . | | | | | | | | | | | | | | | | 371 |
| | NSYINC | | | | | | | | | | | | | | | | 371 |
| | YTICS . | | | | | | | | | | | | | | | | 372 |
| | YMAX . | | | | | | | | | | | | | | | | 372 |
| | YVMAX . | | | | | | | | | | | | | | | | 372 |
| | YMIN | | | | | | | | | | | | | | | | 373 |
| | YVMIN . | | | | | | | | | | | | | | | | 373 |
| | YMOD . | | | | | | | | | | | | | | | | 374 |

| E | INDEX | 401 |
|---|------------------------------|-------|
| D | USER ROUTINE SOURCE CODE | 391 |
| C | AlphaVMS COMMAND PROCEDURE | 389 |
| В | VAX/VMS COMMAND PROCEDURE | 384 |
| | LEFTIC | . 382 |
| | LEFNUM | |
| | TOPTIC | |
| | TOPNUM | |
| | RITTIC | . 381 |
| | RITNUM | |
| | BOTTIC | |
| | BOTNUM | |
| | YAXISA | |
| | YUAXIS | |
| | YLAXIS | |
| | XUAXIS | |
| | XLAXIS | |
| | BOX | |
| | YUWIND | |
| | YLWIND | |
| | XUWIND | |
| | XLWIND | . 377 |
| | A.6 Axis Box Characteristics | |
| | YITICL | |
| | YITICA | |
| | YNUMA | |
| | NYDEC | |
| | NYDIG | |
| | YPOW | |
| | YPAUTO | |
| | YLEADZ | |
| | YOFF | . 374 |

List of Tables

| 2.1 | The control key menu for the 3DPLOT command | 5 |
|------|--|----|
| 2.2 | Control keys recognized by the terminal interface | 13 |
| 2.3 | Function keys recognized by the terminal interface | 13 |
| 2.4 | Interpretation of the IATYPE array in user written subroutines | 15 |
| 2.5 | Interpretation of the ICODE array in user written subroutines | 16 |
| 2.6 | Colour names and associated colour numbers | 20 |
| 2.7 | Exceptions to the standard GPLOT defaults | 30 |
| 2.8 | Line type defaults in centimeters | 30 |
| 2.9 | Hatch pattern defaults in centimeters | 31 |
| 2.10 | Density plot types and their required qualifiers | 31 |
| 2.11 | Keywords used to destroy entire classes of variables | 46 |
| 2.12 | Plotting units for HPLASERJET devices | 48 |
| 2.13 | Plotting units for INKJET devices | 49 |
| 2.14 | Plotting units for PRINTRONIX, LA100, and THINKJET devices | 49 |
| 2.15 | PostScript paper sizes | 50 |
| 2.16 | Plotting units for POSTSCRIPT devices | 51 |
| 2.17 | Pen plotter paper sizes | 51 |
| | Plotting units for pen plotter devices | 52 |
| 2.19 | Plotting units for LNO3+ and IMAGEN devices | 52 |
| 2.20 | Plotting units for GKS graphics metafiles | 53 |
| 2.21 | Plotting units for display file graphics output | 53 |
| 2.22 | Mouse button definitions when digitizing data | 55 |
| | Keyboard key definitions when digitizing data | 55 |
| 2.24 | The PHYSICA keyword menu of default values | 63 |
| 2.25 | The full menu of GPLOT keywords, with values in centimeters | 64 |
| 2.26 | The short menu of GPLOT keywords, with values in centimeters | 65 |
| 2.27 | The menu of GPLOT <i>x</i> -axis characteristics, with values in centimeters | 66 |
| 2.28 | The menu of GPLOT y -axis characteristics, with values in centimeters | 67 |
| 2.29 | The menu of general GPLOT keywords, with values in centimeters | 68 |
| 2.30 | Geometric figures that can be drawn with the FIGURE command | 83 |
| 2.31 | Types of units recognized by the FIGURE command | 83 |

| | Various 1^{st} derivative nonrecursive filters |
|------|--|
| | Various 2 nd derivative nonrecursive filters |
| 2.34 | Various 3^{rd} derivative nonrecursive filters |
| 2.35 | Smoothing nonrecursive filters (quadratic) |
| 2.36 | Smoothing nonrecursive filters (quartic) |
| 2.37 | Smoothing nonrecursive filters (Spencer's formulae) 90 |
| 2.38 | Integrating recursive filters |
| 2.39 | The HISTYP keyword |
| 2.40 | Symmetric error bars |
| 2.41 | Asymmetric error bars |
| | HARDCOPY command print and save codes for bitmap devices |
| 2.43 | HARDCOPY command print and save codes for non-bitmap devices 129 |
| | Types of units recognized by the LEGEND command |
| 2.45 | The LINE command interactive menu |
| 2.46 | Monitor device types and corresponding keywords |
| 2.47 | Key codes for the PEAK command |
| 2.48 | Key codes for the PICK command |
| 2.49 | The pie wedge defining characteristics |
| 2.50 | Types of units recognized by the PIEGRAPH command 163 |
| 2.51 | PLOTTEXT command text formatting commands |
| 2.52 | Formatted text accent special characters |
| 2.53 | Variables that can be read and their required qualifiers |
| 2.54 | The ARROTYP code and corresponding arrow styles |
| 2.55 | Interpretations of the FILL keyword |
| | The hatch pattern defaults |
| 2.57 | Line type definitions |
| 2.58 | The line type defaults |
| 2.59 | The SHOWHISTORY keyword interpretation |
| | The font names |
| | The STATISTICS command extrema keywords |
| | The STATISTICS command central measure keywords |
| 2.63 | The STATISTICS command dispersion and skewness keywords 249 |
| 2.64 | Text justification interaction with CURSOR |
| 2.65 | Text menu and justification |
| 2.66 | TEXT command text formatting commands |
| 2.67 | The initial pre-defined windows |
| 3.68 | Boolean operators |
| | Other operators |
| 4.70 | Trigonometric functions |
| 471 | Basic element by element numeric functions 290 |

List of Figures

| 2.1 | Interpolating a fine mesh on the contours of a matrix | 26 |
|------|--|-------------|
| 2.2 | An example of a box type density plot with both x and y profiles | 42 |
| 2.3 | Examples of box type density plot with accentuated and delimited values | 43 |
| 2.4 | Box type density plots with scattered points and with a matrix | 44 |
| 2.5 | An example of a font table produced by the DISPLAY FONT command | 60 |
| 2.6 | The table of special characters | 61 |
| 2.7 | An example of the hatch fill patterns | 61 |
| 2.8 | An example of the default line types | 62 |
| 2.9 | The special plotting symbols | 62 |
| 2.10 | A FILTER example showing data smoothing | 92 |
| 2.11 | A FILTER example showing 1^{st} derivative $\dots \dots \dots$ | 93 |
| 2.12 | Finding a local minimum with the FMIN command | 103 |
| 2.13 | Finding roots with the FZERO command | 105 |
| 2.14 | Plotting error bars with the GRAPH command | 121 |
| 2.15 | Filling the area under a curve drawn with the GRAPH command | 122 |
| 2.16 | Histogram examples drawn with the GRAPH command | 123 |
| 2.17 | Automatic axis labels using the LABEL command | 137 |
| 2.18 | An example illustrating the graph LEGEND | 143 |
| | Pie chart wedge definition | |
| | An example of a pie chart | |
| 2.21 | Example accents on the letter "o" | 182 |
| 2.22 | An example demonstrating the POLYGON command | 183 |
| 2.23 | An example using the SCALES command | 229 |
| 2.24 | Arrow styles | 230 |
| | An example using the SLICES command | |
| 2.26 | Surface coordinate system | 255 |
| 2.27 | A SURFACE example | 256 |
| 2.28 | Text extent rectangle with two-character justification codes | 258 |
| | An example using the TEXT command | |
| | An example using the TILE command | |
| 2.31 | The initial pre-defined windows in PORTRAIT orientation | 27 3 |

| 2.32 | An example illustrating the ZEROLINES command | 30 |
|------|---|----|
| 4.33 | An example illustrating the DERIV function | 18 |
| 4.34 | An example illustrating the INTEGRAL function | 30 |
| 4.35 | An FFT example showing data smoothing | 27 |
| A.36 | Some <i>x</i> -axis characteristics | 56 |
| A.37 | Logarithmic x-axis examples | 58 |
| A.38 | Virtual axes examples | 33 |
| A.39 | Some y-axis characteristics | 37 |
| A.40 | Logarithmic y-axis examples | 36 |
| A.41 | The window and axis locations | 17 |

1 INTRODUCTION

PHYSICA provides a high level, interactive programming environment. The program constitutes a fully procedural programming language, with built-in user friendly graphics and sophisticated mathematical analysis capabilities. Combining an accessible user interface along with comprehensive mathematical and graphical features, makes PHYSICA a general purpose research tool for scientific, engineering and technical applications.

PHYSICA provides you with a wide range of mathematical and graphical operations. Over 200 mathematical functions are available, as well as over 30 operators, providing all of the standard operations of simple calculus, along with powerful curve fitting, filtering and smoothing techniques. The program employs a dynamic array management scheme allowing you a large number of arrays of unlimited size. Algebraic expressions are evaluated using a lexical scanner approach. These expressions can have up to 1500 "tokens," where a token is a literal constant, a variable name, a function name, or an operator. Array evaluations and assignments can be implemented in a simple, direct manner.

Line graphs, histograms and pie-charts, as well as contour, density and surface plots are available. Publication quality graphics can be easily obtained. You have complete control over the appearance of a drawing.

Initial development was for the VAX/VMS operating systems, but the program has been ported to AlphaVMS, ULTRIX, Digital Unix, Silicon Graphics IRIX, HP-UX, IBM AIX, SUNOS and Solaris, and most recently, PC Linux.

The user interacts with the program through the user interface, consisting of monitor dependent routines for display of messages and for reading user input, and device dependent routines for displaying drawings and obtaining hardcopies of user sessions. The user interface is a high-level command language that incorporates a simple to use and easy to learn syntax, based on context-free lexical scanners. The command language incorporates the basic elements of a structured programming language, including conditional branching, looping and subroutine calling constructs.

1.1 What is in this manual

The bulk of this manual is a reference guide to the program commands. These commands are discussed in alphabetical order.

The PHYSICA program provides the user with a large variety of analysis tools, including a fairly comprehensive set of operators and functions. The next chapter describes the operators that are available for use in expressions, followed by a chapter on the built-in functions

What is in this manual

that can also be used in any expression.

The first appendix contains definitions for the GPLOT graph and text plot characteristic keywords which are controlled by the SET and GET commands. Other appendices contain the command procedures for creating shareable images of user defined routines, for use with the VAX/VMS and AlphaVMS operating systems. This is followed by the default source code for these user defined routines.

Users are referred to the PHYSICA USER'S GUIDE for examples of program usage. Those who are familiar with the predecessor program, PLOTDATA, are referred to the PLOTDATA TO PHYSICA CONVERSION MANUAL for tips on converting PLOTDATA command macros to PHYSICA.

1.2 Conventions used in this manual

Examples of messages and prompts written by the program, as well as examples of user typed input are displayed in typewriter type style.

Commands and other reserved keywords are in UPPERCASE.

Curly brackets, { }, enclose parameters that are optional and/or have default values; and indicate that it is not necessary to enter these parameters. Vertical bars, |, separate choices for command parameters.

Curly brackets and vertical bars should *not* be entered with commands.

Parentheses, (), enclose formats. The back slash, \setminus , separates a command from a command qualifier or a parameter from its qualifier. The opening quote, ', and the closing quote, ', delimit literal strings.

Parentheses, the back slash and quotes *must* be included where indicated.

VMS usually refers to the OpenVMS operating system for either the VAX or the Alpha architectures.

UNIX refers to any UNIX like operating system, including Linux.

2 COMMANDS

3DPLOT

```
Syntax 3DPLOT x y z ipen { colr }

Defaults colr = current colour
```

The 3DPLOT command graphs the three vectors \mathbf{x} , \mathbf{y} , and \mathbf{z} in 3d space, displayed in 2 dimensions using a perspective projection. The vectors \mathbf{x} , \mathbf{y} , and \mathbf{z} should contain the (x,y,z) coordinates of the points. The ipen vector contains the codes for deciding what to draw at each coordinate. The colr vector contains the colour codes. The vectors must all be the same length.

This command is strictly interactive, it cannot be entered from batch mode. Note that if the graph seems to be a complete mess, try increasing the "eye to object distance" using the E key.

| ipen[j] | |
|---------|---|
| =2 | coordinate set j is connected by a line segment to j-1 |
| =3 | coordinate set j is not connected to j-1 |
| = 20 | a point is plotted at coordinate set j |
| < 0 | a GPLOT symbol ipen[j] is plotted at coordinate set j |

When drawing symbols, the size of the symbols can be controlled with the SET CHARSZ command before entering the 3DPLOT command, and changed inside the command by using the Z key. Use the DISPLAY PCHAR command to see the possible GPLOT special symbols.

If drawing line segments, colr[j] is the colour code for the last line segment j-1 to j. The first colour code is ignored. If drawing a point at the j_{th} coordinate, that is, ipen[j] = 20, or if drawing a symbol at the j_{th} coordinate, that is, ipen[j] < 0, then colr[j] is the colour of that point or symbol.

On X Window type monitors, the focus must be in the alphanumeric terminal window to use the menu.

The default axis number height is XNUMSZ, which can be changed with the SET XNUMSZ command before entering the 3DPLOT command, and changed inside the command by using the H key.

The current hardcopy device is disabled while the 3DPLOT command is active. The S key causes the current hardcopy device to be re-enabled, and the plot to be redrawn, thus allowing you to obtain a hardcopy, with the HARDCOPY command.

- < rotate left 1 angle increment</p>
- > rotate right 1 angle increment
- ^ rotate up 1 angle increment
- V rotate down 1 angle increment
- I zoom in
- A angle increment
- **H** axis number height
- **S** save for hardcopy
- Q quit

- L rotate left 360 degrees
- R rotate right 360 degrees
- U rotate up 360 degrees
- D rotate down 360 degrees
- 0 zoom out
- E eye to object distance
- Z symbol size
- M display this menu

Table 2.1: The control key menu for the 3DPLOT command

On X Window type monitors, the keyboard focus must be in the alphanumeric terminal window to use the menu.

ALIAS

| Syntax | ALIAS { newcommand command_string } |
|----------|-------------------------------------|
| Examples | ALIAS |
| | ALIAS RED COLOUR RED |
| | ALIAS APPEND WRITE\APPEND |
| | ALIAS CURVE GRAPH\NOAXES |
| | ALIAS AXES GRAPH\AXES |
| | ALIAS ACLEAR CLEAR\ALPHANUMERIC |

The ALIAS command creates new commands by equating a user defined keyword, newcommand, to a string, command_string. The command string must begin with a valid command, for example, it cannot be a file name that is to be used with other commands. Everything after newcommand is taken for the command_string, that is, no quotes should be used.

If the ALIAS command is entered with no parameters, then all of the current aliases will be displayed. The maximum number of aliases that can exist at one time is 100./indexalias!maximum number

Use the DEALIAS command to eliminate aliases.

ASSIGN

| Syntax | ASSIGN name logical |
|---------|----------------------------|
| Example | ASSIGN LASER_211 HP\$LASER |

The ASSIGN command is only relevant for VMS. It is equivalent to the DCL command: ASSIGN name logical

The logical assignment takes effect immediately, and remains in effect after PHYSICA is unloaded. Both parameters should be strings.

The ASSIGN command is useful for assigning new logical names to the output devices, that is, the logical name for a bitmap device or a plotter.

BESTFIT

| Syntax | BESTFIT pmin pmax penalty error parm pout |
|------------|--|
| | BESTFIT\CYCLES n pmin pmax penalty error parm pout |
| | BESTFIT\WEIGHTS w pmin pmax penalty error parm pout |
| | BESTFIT\CYCLES\WEIGHTS w n pmin pmax penalty error parm pout |
| Qualifiers | \CYCLES, \WEIGHTS |
| Defaults | \NOCYCLES, \NOWEIGHTS, pmin = 0 , pmax = 1 , penalty = 1 |

This command calculates parameters for a least-squares fit to an error vector using adjustable parameters.

Suppose you have an error vector, error, of length n. Suppose that there are m variable parameters and that the measured effect of a unit change for each of the parameters at each of the n locations is stored in matrix parm with n rows and m columns. Vector penalty, of length m, represents the penalty functions to changes of the m parameters. The larger penalty[i] the smaller the adjustment of the i^{th} parameter.

The optimal set of changes of the m parameters within their allowed range of pmin to pmax will be determined in the least-squares sense. The vector pout will contain the parameter changes giving this fit, and will be of length m.

Parameter types and sizes

The influence function parm must be a matrix. Suppose it has n rows and m columns. The parameter ranges, pmin and pmax, as well as the penalty function penalty, must be vectors with the same length, m, which is the number of parameters for the fit. The error vector error must be a vector of length n, which is the number of locations.

Weights

Syntax BESTFIT\WEIGHTS w pmin pmax penalty error parm pout BESTFIT\WEIGHTS\CYCLES w n pmin pmax penalty error parm pout

If a weight vector, w, is entered, you *must* indicate that it is there by using \WEIGHTS qualifier. The weight, w[i], corresponds to the importance of reducing the initial error to zero at the

 \mathbf{i}^{th} location. The weight array should be a vector of length n. The closer to zero the value of w[i], the looser will be the fit at the \mathbf{i}^{th} location. If the \CYCLES qualifier is also used, the weight array comes before the iteration number in the parameter list.

Cycles

```
Syntax BESTFIT\CYCLES n pmin pmax penalty error parm pout BESTFIT\CYCLES\WEIGHTS w n pmin pmax penalty error parm pout
```

The \CYCLES qualifier allows the user to specify the maximum number of iteration steps for the fit. When this maximum number is reached, the fit will stop. The fit will also stop if the fit is successful before this maximum iteration number is reached. If the \WEIGHTS qualifier is also used, the weight array comes before the iteration number in the parameter list.

BIN

```
Syntax BIN x xbin xcount
BIN\NBINS x xbin xcount n { xmin xmax }

Qualifiers \WEIGHTS, \EDGES, \NBINS, \DISCARD, \EMPTY, \AVERAGE, \LAGRANGE

Defaults \-WEIGHTS, \-EDGES, \-NBINS, \-DISCARD, \-EMPTY, \-AVERAGE,
\LAGRANGE xmin = min(x), xmax = max(x)
```

The BIN command sorts an input vector, \mathbf{x} , into a grid of bins and accumulates the counts per bin into an output vector, \mathbf{x} count. Each element of \mathbf{x} is considered only once, so elements are never counted as being in more than one bin. By default, the bins are defined by their centres, given in vector \mathbf{x} bin, which must be strictly monotonically increasing. If $n = \text{len}(\mathbf{x}$ bin), define the bin ranges, r_i

$$r_1 = \texttt{xbin}_1 - (\texttt{xbin}_2 - \texttt{xbin}_1)/2$$

$$r_i = \texttt{xbin}_i - (\texttt{xbin}_i - \texttt{xbin}_{i-1})/2 \quad \textbf{for} \quad i = 2, 3, \ldots, n$$

$$r_{n+1} = \texttt{xbin}_n + (\texttt{xbin}_n - \texttt{xbin}_{n-1})/2$$

For each $i=1,2,\ldots, \operatorname{len}(\mathtt{x})$, if $r_j \leq \mathtt{x}_i < r_{j+1}$ for some $j=1,2,\ldots,n$ then \mathtt{xcount}_j is incremented by 1, or by the weight, \mathtt{w}_i . By default, events below r_1 will be placed in the first bin, and events above r_{n+1} will be placed in the last bin. If the \DISCARD qualifier is used, events outside this range will be discarded.

Weights

```
Syntax BIN\WEIGHTS w x xbin xcount
BIN\AVERAGE\WEIGHTS w x xbin xcount
BIN\EMPTY\WEIGHTS w x xbin xcount
BIN\EDGES\WEIGHTS w x xbin xcount
BIN\EDGES\EMPTY\WEIGHTS w x xbin xcount
BIN\EDGES\AVERAGE\WEIGHTS w x xbin xcount
```

By default, a bin count is incremented by one (1) for every event that goes in a bin. If a weight vector is entered, you *must* indicate that it is there by using \WEIGHTS qualifier. The weight w must be a vector with the same length as x. The i^{th} event causes the bin count to be incremented by w_i .

Number of bins

```
Syntax BIN\NBINS x xbin xcount n { xmin xmax }

Defaults xmin = min(x), xmax = max(x)
```

By default, the bins are defined by their centres, given in vector xbin, which must be strictly monotonically increasing. If the \NBINS qualifier is used, the number of bins, n, is expected. A new vector, xbin, will be created which will have n elements. If the numbers xmin and xmax are not entered, they default to the minimum and maximum of vector x.

$$exttt{xbin}_i = exttt{xmin} + (i - rac{1}{2})(exttt{xmax} - exttt{xmin})/ exttt{n} ext{ for } i = 1, 2, \dots, exttt{n}$$

Lagrange

Syntax BIN\LAGRANGE x xbin xcount

If the \LAGRANGE qualifier is used, \WEIGHTS, \EDGES, \AVERAGES, and \EMPTY are not allowed.

If n = len(xbin), define the bin ranges, r_i

$$r_1 = exttt{xbin}_1 - (exttt{xbin}_2 - exttt{xbin}_1)/2$$

$$r_i = exttt{xbin}_i - (exttt{xbin}_i - exttt{xbin}_{i-1})/2 \quad ext{for} \quad i = 2, 3, \ldots, n$$

$$r_{n+1} = exttt{xbin}_n + (exttt{xbin}_n - exttt{xbin}_{n-1})/2$$

For each $i=1,2,\ldots$, len(x) find j so that $r_j \leq x_i < r_{j+1}$ for some $j=1,2,\ldots,n$. If j=n, then $x = (x_i - x b i n_j)/(r_{j+1} + r_j)/2$

Averages

```
Syntax BIN\AVERAGE x xbin xcount BIN\AVERAGE\EDGES x xbin xcount
```

The \AVERAGE qualifier means that the output xcount vector will contain the average value for each bin. An internal counter is kept for each bin, and the value for $xcount_i$ will be divided by the number of events in bin i before it is output. \AVERAGE can be used with \WEIGHTS and \EDGES, but not with \EMPTY.

Increment only if empty

```
Syntax BIN\EMPTY x xbin xcount BIN\EMPTY\EDGES x xbin xcount
```

The \EMPTY qualifier means that an event is counted in a bin only if that bin is empty. So only the first event encountered for each bin will be counted in that bin. \EMPTY can be used with \WEIGHTS and \EDGES, but not with \AVERAGE.

Edge defined bins

```
Syntax BIN\EDGES x xbin xcount
```

By default, bins are defined by their centres. If the \EDGES qualifier is used, bins are defined by their edges. The bin edges must be given in vector xbin. The length of xcount will be one less than the length of xbin. A weight vector, w, may be specified if you use the \WEIGHTS qualifier.

If $xbin_j \le x_i < xbin_{j+1}$ then $xcount_j$ is incremented either by one (1), or by the specified weight, w_i .

BIN2D

```
Syntax

BIN2D x y xbin ybin mc nx ny { xmin xmax ymin ymax }

BIN2D\MATRIX mdata mxin myin mout

Qualifiers \WEIGHTS, \EMPTY, \MATRIX, \XDISCARD, \YDISCARD, \DISCARD

Defaults \-WEIGHTS, \-MATRIX, \-EMPTY, \-DISCARD \

xmin = min(x), xmax = max(x), ymin = min(y), ymax = max(y)
```

The BIN2D command forms a matrix of bins of data by sorting the vectors x and y into grids of bins which are returned in vectors x bin and y bin. The accumulated matrix of total counts per bin is returned in matrix mc. If the numbers x min and x max are not entered, they default

to the minimum and maximum of x. Similarly, if the numbers ymin and ymax are not entered, they default to the minimum and maximum of y.

$$exttt{xbin}_i = exttt{xmin} + (i - rac{1}{2})(exttt{xmax} - exttt{xmin})/| exttt{nx}| ext{ for } i = 1, 2, \ldots, | exttt{nx}|$$

$$\mathtt{ybin}_j = \mathtt{ymin} + (j - \frac{1}{2})(\mathtt{ymax} - \mathtt{ymin})/|\mathtt{ny}| \quad \mathbf{for} \quad j = 1, 2, \dots, |\mathtt{ny}|$$

The (x_i, y_i) point will be accumulated in $mc_{i,j}$ where:

```
row: i = int((y_i - ymin)/(ymax - ymin)|ny|) + 1

column: j = int((x_i - xmin)/(xmax - xmin)|nx|) + 1
```

Dimensions

The lengths of x and y must be equal. If a weight vector, w, is supplied, it must also be the same length.

The vectors xbin and ybin and the matrix mc will be created. xbin will have |nx| elements, ybin will have |ny| elements, and matrix mc will have |nx| columns and |ny| rows.

Extremes

By default, events below xmin are placed in the first bin column, events above xmax are placed in the last bin column, events below ymin are placed in the first bin row, and events above ymax are placed in the last bin row. If the \DISCARD qualifier is used, events outside either of these ranges will be discarded. If the \XDISCARD qualifier is used, events below xmin are discarded, and events above xmax are discarded. If the \YDISCARD qualifier is used, events below ymin are discarded, and events above ymax are discarded.

Weights

```
Syntax BIN2D\WEIGHTS w x y xbin ybin mc nx ny { xmin xmax ymin ymax }
```

If a weight is entered, you *must* indicate that it is there by using \WEIGHTS qualifier. The weight w must be a vector. The i^{th} event causes the bin count to be incremented by w_i .

Increment only if empty

```
Syntax BIN2D\EMPTY x y xbin ybin mc nx ny { xmin xmax ymin ymax }
```

If the $\backslash \text{EMPTY}$ qualifier is used, an event is counted in a bin only if that bin is empty. Only the first event encountered for each bin will be counted in that bin. $\backslash \text{EMPTY}$ cannot be used with $\backslash \text{MATRIX}$.

Defined by box corners

```
Syntax BIN2D\MATRIX mdata mx my mc
```

The BIN2D\MATRIX command calculates the sum of the data points given by matrix mdata within a set of boxes. The x-coordinates of the boxes are given in matrix mx, the y-coordinates are given in matrix my. Matrices mx and my must be the same size. A data point is taken to be inside a box if it is interior or on an edge. Each data point is considered only once, so a data point is never taken to be in more than one of the boxes. The coordinates of the data points are the row and column indices, for example, mdata[3,4] is row 3 and column 4 so it is at (x,y) location (4,3). The x and y-coordinates in mx and my should be in this index space of coordinates. The qualifiers \EMPTY and \WEIGHTS cannot be used with \MATRIX.

See the PICK\MATRIX command for information on interactively choosing the above mentioned boxes.

BUFFER

```
Syntax BUFFER { n }
BUFFER\READ filename
BUFFER\WRITE filename

Qualifiers READ, WRITE, DYNAMIC, STATIC, KEYPAD

Defaults n = 20, \DYNAMIC
```

The BUFFER command controls the input line recall buffers:

- the dynamic buffer
- the static buffer
- the keypad buffer

Table 2.2 on page 13 shows the control keys recognized by this terminal interface. Table 2.3 on page 13 shows the function keys recognized by the terminal interface.

See the RESTORE\PHYSICA command, page 211, for information on restoring these buffers from previously saved sessions. There is an option to not restore these buffers, by using the \NOTTBUFFERS qualifier.

Parameters

If no parameters are entered and no qualifiers are used, the current length of the dynamic recall buffer is displayed. If n is entered, this will be the new length of the dynamic buffer as displayed when the PF1 key is typed. The value of n must be $0 < n \le 35$.

Reading the buffers

If the \READ qualifier is used, the dynamic buffer will be read from the specified file. You can also read the static buffer or the keypad buffer by also using the \STATIC qualifier or the \REYPAD qualifier. For example:

BUFFER\KEYPAD\READ FILE.DAT

will read the keypad buffer from FILE.DAT

Writing the buffers

If the \WRITE qualifier is used, the dynamic buffer will be written to the specified file. You can also write the static buffer or the keypad buffer by also using the \STATIC qualifier or the \KEYPAD qualifier. For example:

BUFFER\KEYPAD\WRITE FILE.DAT

will write the keypad buffer to FILE.DAT

Dynamic buffer

The dynamic buffer is a terminal interface which closely mimics the DCL command recall facility. The arrow, delete, backspace, and most control keys, work as in DCL. An input line is stored automatically in the dynamic buffer when a carriage return is typed. The line is stored at the top of the stack, with previously entered lines being pushed down the stack. The maximum length of the buffer stack is 35 lines. After 35 lines have been stored, the lines at the bottom of the stack begin dropping off and are lost.

Static buffer

The static buffer is similar to the dynamic buffer, but the lines in the static buffer are not updated automatically. To interactively enter a line into the static buffer, type the input line and then type the PF3 keypad key followed by control-L. You will be asked to enter a storage number. Enter a digit from 1 to 9 or a letter from A to Z, where A represents 10, B represents 11, and so on. To recall a line previously stored in the static buffer, type the PF3 keypad key, and then type the storage digit or letter of the desired line.

Keypad buffer

The keypad buffer allows the keypad keys: 0 - 9, period, comma, and minus, to be defined. Any of these keys can also be set up such that a carriage return is included. Thus, the

command is executed as soon as the key is typed. To interactively enter a keypad key definition, type the input line and then type the <enter> key on the keypad. You will be asked to type the keypad key to be loaded with the input line. If this is to include a carriage return, type the <enter> keypad key, otherwise type any other key to resume.

| key | action |
|---|--|
| control-^ | appended to a string recalls the last command containing it |
| control-A | toggles insert/overstrike mode |
| control-E | moves alphanumeric cursor to end of line |
| control-H | (BACKSPACE) moves alphanumeric cursor to the beginning of the input line |
| control-K | disables recall shell, a "!" in column 1 re-enables it |
| control-N | reads the dynamic recall buffer from a file |
| control-P | writes the dynamic recall buffer to a file |
| control-R | refreshes the current input line |
| control-X | (control-U) erases input line to the left of the alphanumeric cursor |
| Currently LINEFEED (control-J), ESC and TAB (control-I) are not enabled | |

Table 2.2: Control keys recognized by the terminal interface

| key | action |
|-------|--|
| PF1 | list and allows selection from dynamic recall buffer |
| PF2 | lists the HELP facility |
| PF3 | lists, loads (via control-L), and selects from static buffer |
| PF4 | invokes a simple desk calculator |
| ENTER | lists or loads the keypad buffer |
| F14 | toggles insert/overstrike mode |

Table 2.3: Function keys recognized by the terminal interface

CALL

```
Syntax CALL { SUBn } arg1 { arg2 ... arg15 }

Examples CALL SUB1 X Y

CALL X Y
```

The CALL command uses one of the following:

- a subroutine hardwired into the program
- a subroutine loaded dynamically with the LOAD command (VAX/VMS only), or
- a subroutine loaded at run time via a shareable image (VAX/VMS or AlphaVMS).

Functions can not be referenced with the CALL command, but the user written functions can be used wherever a function can be used in an expression.

If one of the keyword parameters SUB1, SUB2, ..., SUB8 is entered with the CALL command, either the built-in subroutine by default, or one of the eight subroutines which were loaded at run time via PHYSICA_USER_FUNCTIONS, a shareable image, will be used. The shareable image option is only available under VAX/VMS and AlphaVMS.

If none of these keyword parameters is used, then the object module that was loaded dynamically with the LOAD command will be used. The LOAD command is only available under VAX/VMS.

For UNIX users, the process to add user defined routines to physica follows:

- 1. edit the file phys_user.f to put in your versions of user1, ..., user8, sub1, ..., sub8
- 2. compile it, e.g., f77 -c phys_user.f
- 3. put it in the archive, e.g., ar -rsv physica.a phys_user.o
- 4. link the program with physica.link

The default set of eight (8) subroutines and eight (8) functions is listed in **Appendix D**. For VMS users the sources are provided in the file: PHYSICA\$DIR:PHYSICA_USER_FUNCTIONS.FOR For UNIX users the sources are provided in the file: phys_user.f Off-site UNIX users can find this file in the physica-link tar file.

User written subroutine description

If a subroutine is loaded at run time via the sharable image, its name *must* be one of SUB1, SUB2, ..., SUB8. If a subroutine is to be loaded dynamically via the LOAD command, its name

is irrelevant and can be anything the user desires. In either case, a user written subroutine *must* have the following form:

```
SUBROUTINE subname(IATYPE, ICODE, IUPDATE, IER, arg1, arg2,...)
INTEGER*4 IATYPE(15), ICODE(3,15), IUPDATE(15), IER
```

Other than the required arguments, IATYPE, ICODE, IUPDATE, and IER, there may be from 1 to 15 arguments in the subroutine argument list. The user is responsible for insuring that the correct number and type of arguments are used when actually employed with the CALL command. The parameters used in the CALL command, argI, which are passed as arguments to the subroutine, may be constants, scalars, vectors, matrices, literal quote strings, or *scalar* string variables. The number of arguments and the type of argument *must* agree with the actual subroutine.

All of the numeric aguments, except for the required integer arguments IATYPE, ICODE, IUPDATE, and IER, must be REAL*8. A string argument is passed as a LOGICAL*1 array.

Note: The integer arguments IATYPE, ICODE, IUPDATE, and IER should *not* be mentioned as parameters with the CALL command.

See the file: PHYSICA\$DIR:PHYSICA_USER_FUNCTIONS.FOR for some subroutine examples.

IATYPE

IATYPE is an INTEGER*4 array, length 15, that indicates the type of each of the subroutine arguments argI. See Table 2.4 on page 15.

| argument type | <pre>IATYPE(i)</pre> |
|---------------|----------------------|
| unfilled | -99 |
| string | -1 |
| scalar | 0 |
| vector | 1 |
| matrix | 2 |

Table 2.4: Interpretation of the IATYPE array in user written subroutines

ICODE

ICODE is an INTEGER*4 array, dimensioned 3 by 15, that indicates the dimension of each of the subroutine arguments argI. Never extend variables beyond their original size as passed to the subroutine. If a variable is shortened inside the subroutine, the subroutine must update the new dimensions in the ICODE array, so that PHYSICA can reduce the variable

dimensions appropriately. See Table 2.5 on page 16.

| argument type | ICODE(1,i) | ICODE(2,i) | ICODE(3,i) |
|---------------|------------|------------|------------|
| string | length | 0 | 0 |
| scalar | 0 | 0 | 0 |
| vector | length | 0 | 0 |
| matrix | nrows | ncolms | 0 |

Table 2.5: Interpretation of the ICODE array in user written subroutines

The ICODE array will be filled by PHYSICA with the current dimensions of the arguments, so the user written subroutine can check and, if necessary, update the dimensions of any of the subroutine arguments.

Never extend vectors, matrices, or string variables beyond their original sizes as passed to the user written subroutine. If a variable's size is shortened inside the subroutine, then the subroutine must update the ICODE array so that these variable dimensions can be reduced internally by PHYSICA upon return from the subroutine.

IUPDATE

IUPDATE is an INTEGER*4 array, length 15, that the user routine sets to indicate to PHYSICA whether one of the argI arguments has been modified inside that subroutine.

The default value for <code>IUPDATE(i)</code> is <code>0</code>. Set <code>IUPDATE(i)</code> to <code>1</code> to indicate that the <code>ith</code> argument, argI, has been modified. Never extend variables beyond their original size as passed to the subroutine. If a variable is shortened inside the subroutine, the subroutine must update the new dimensions in the <code>ICODE</code> array, so that <code>PHYSICA</code> can reduce the variable dimensions appropriately.

IER

IER is an INTEGER*4 variable that defaults to the value 0. Your routine can set IER to indicate to PHYSICA that an error has occured in the routine. Arithmetic errors, such as division by zero, over/underflow, will be asynchronously trapped. If other error tests are to be done inside the subroutine, the user flags the error by setting IER = -1 before the RETURN. If the CALL command was executed from within a script, this error flag causes PHYSICA to abort that script and control is passed back to the keyboard.

Numeric arguments

All the numeric arguments of your subroutine, except for the integer arguments IATYPE,

ICODE, IUPDATE, and IER, must be REAL*8. A string argument is passed as a LOGICAL*1 array. Dimension numeric array arguments with length 1, for example:

```
REAL*8 X(1), Y(1), Z(1)
```

String arguments

All the string arguments of your subroutine must be LOGICAL*1, and should be dimensioned 1, for example:

```
LOGICAL*1 LFILE(1)
```

You can convert this to a string, say, CHARACTER*80 CFILE, using the following method:

```
LENF = ICODE(1,i)
DO I = 1, LENF
   CFILE(I:I) = CHAR(LFILE(I))
END DO
```

where LFILE is the i^{th} argument.

Accessing matrix data

If a matrix is passed as an argument to a user written subroutine, the elements of the matrix can only be accessed using a calculated index. To access element m[i,j] of the matrix m, use m[i+(j-1)*nrows] for $i=1, \ldots, nrows$ and $j=1, \ldots, ncols$.

Example

If the command is CALL SUB3 A T X M, where A is a scalar, T is a string variable, X is a vector, and M is a matrix, the subroutine should begin as follows:

```
SUBROUTINE SUB3(IATYPE,ICODE,IUPDATE,IER,A,T,X,M)

INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER

REAL*8 A, X(1), M(1)

LOGICAL*1 T(1)

...

LENT =ICODE(1,2) ! the length of the string variable T

LENX =ICODE(1,3) ! the length of vector X

NROWS=ICODE(1,4) ! the number of rows of the matrix M

NCOLS=ICODE(2,4) ! the number of columns of the matrix M

...

RETURN

END
```

Creating a shareable image

The default set of eight (8) subroutines and eight (8) functions, listed in **Appendix D**, is provided in the file:

```
PHYSICA$DIR: PHYSICA_USER_FUNCTIONS.FOR
```

Copy this file to your own directory. For example:

```
$ copy PHYSICA$DIR:PHYSICA_USER_FUNCTIONS.FOR disk:[directory]TEST.FOR
```

Substitute your source code for the default routines. All eight subroutines and all eight functions *must* be present in your source code file.

A DCL command procedure must be executed *before* you invoke PHYSICA. This command procedure, the VAX/VMS version is listed in **Appendix B** while the AlphaVMS version is listed in **Appendix C**, can be found in: PHYSICA\$DIR:PHYSICA_USER_FUNCTIONS.COM

This procedure defines the logical name PHYSICA_USER_FUNCTIONS and creates the sharable image. There should be a system wide default definition of this logical name, so that if the shareable image is not in a user defined location, the default will be used from PHYSICA\$DIR.

Copy this procedure to your own directory. For example:

```
$ copy PHYSICA$DIR:PHYSICA_USER_FUNCTIONS.COM disk:[directory]TEST.COM
```

You will need to modify one line:

\$ define PHYSICA_USER_FUNCTIONS disk:[directory]TEST

In this line, replace disk: [directory] TEST with the actual location and name of your source code file. Execute the command procedure, which will produce, PHYSICA_USER_FUNCTIONS.EXE, a shareable image.

CLEAR

Syntax CLEAR

Qualifiers \ALPHANUMERIC, \TOGGLE, \REPLOTONLY, \NOREPLOT

Defaults clears graphics and replot option

By default, the CLEAR command clears the graphics. It also clears the replot buffers, that is, there will be nothing to replot until something is drawn again. Any hardcopies must be asked for *before* entering the CLEAR command.

Alphanumerics

Syntax CLEAR\ALPHANUMERIC

The CLEAR\ALPHANUMERIC command clears the alphanumeric, transparent, portion of the monitor screen only. This has no affect on graphics hardcopies or the REPLOT buffers.

Toggle graphics

Syntax CLEAR\TOGGLE

On a CIT467 terminal, the CLEAR\TOGGLE command will toggle the graphics screen on or off. The first time this command is entered, the graphics screen is turned off, and any graphics will disappear. The next time CLEAR\TOGGLE is entered, the graphics screen will be turned back on, and your graphics will reappear. This has no affect on graphics hardcopies or the REPLOT buffers.

Clear the replot buffers only

Syntax CLEAR\REPLOTONLY

The CLEAR\REPLOTONLY command only clears the replot buffers and does not affect the graphics. There is nothing to replot after CLEAR\REPLOTONLY until more data or text is drawn.

Do not clear the replot buffers

Syntax CLEAR\NOREPLOT

The CLEAR\NOREPLOT command only clears the graphics, it does not affect the replot buffers.

COLOUR

Syntax COLOUR colourname

COLOUR n

Defaults default colour is white

Examples COLOUR

COLOUR RED

 ${\tt COLOUR} \ {\tt R} \backslash {\tt SCALAR}$

The COLOUR command sets the graphics monitor colour and the associated graphics hardcopy colour number, for subsequent graphics. The colour number may, for example, be a plotter pen number.

If no parameter is entered, a list of the colour names and corresponding numbers is displayed. You may then enter a colour name or number. Table 2.6 shows the recognized colournames and their associated numbers.

Note: The colours black and white have always been a source of confusion. The colour black means the graphics screen background colour. The colour white means white if the background colour is black, and black if the background colour is white.

| name | number | name | number |
|-------|--------|---------|--------|
| black | 0 | | |
| white | 1 | yellow | 5 |
| red | 2 | cyan | 6 |
| green | 3 | magenta | 7 |
| blue | 4 | white | 8 |

Table 2.6: Colour names and associated colour numbers

Note: For pen plotters, the colour of the pens should be confirmed before submitting a plot file to a plotter.

Using a scalar for a colour number

If a scalar variable is to be used for the colour number, it must have the qualifier \SCALAR attached to it. For example:

COLR=5

COLOUR COLR\SCALAR

CONTOUR

```
Syntax CONTOUR { x y } v nctr { min { inc }}
CONTOUR\SPECIFIC { x y } v lvls

Qualifiers \SPECIFIC, \INTERPSIZE, \POLAR, \LEGEND, \COLOURS, \PARTIAL, \RESET, \CONTINUE, \BORDER, \AXES, \COORDINATES, \AREAS, \VOLUMES

Defaults x = [1;2;3;...] y = [1;2;3;...] \NOSPECIFIC, \NOINTERPSIZE, \NOPOLAR, \NOLEGEND \NOCOLOURS, \NOPARTIAL, \RESET, \BORDER, \AXES \NOCOORDINATES, \NOAREAS, \NOVOLUMES, \NOCONTINUE

Examples CONTOUR X Y Z 10 \CONTOUR M 10 MINVAL \CONTOUR\SPECIFIC\INTERPSIZE 10 X Y M LEVELS
```

The CONTOUR command draws contour lines for either data contained in a regular matrix or a scattered set of points contained in three vectors.

Contour level selection

```
Syntax CONTOUR { x y } v nctr { min { inc }}
CONTOUR\SPECIFIC { x y } v lvls
```

By default, the number of contours, nctr, must be provided. If $\mathtt{nctr} > 0$ and the increment is not specified, the actual number of contours drawn may not be the same as the number that was asked for, since "nice" contour levels will be selected and the range of values may not be neatly divisible by the requested number. If the minimum is provided, but not the increment, a "nice" value close to \mathtt{min} will be used instead of the actual data minimum. If the minimum and the increment are both specified, those exact values will be used for the contour levels.

Specific contour levels

Specific contour levels can be requested by using the \SPECIFIC qualifier. In this case, the vector lvls should contain the desired contour levels.

Exact contour levels

Exact contour levels can be requested in three ways.

- Use the \SPECIFIC qualifier and enter a vector containing the desired levels.
- Specify the minimum contour level, min, and the contour level increment, inc. This

```
produces a set of equally spaced contour levels, [min; min+inc; min+2*inc; ...; min+nctr*inc].
```

 Specify a negative number of contours, nctr < 0. This produces a set of equally spaced contour levels, as above, using the actual data minimum and the actual data maximum.

Zooming in

When the minimum and maximum contour levels are to be determined, by default, the entire range of the data is used. If the \PARTIAL qualifier is used, the minimum and maximum contour levels will be determined by the region contained within the axes. Thus, to zoom in on a particular region for more detail, pre-set the axis scales, using the SCALES command, before entering the CONTOUR command. Of course, the \PARTIAL qualifier does not apply when you request specific contour levels.

Contour level colour

Colour contours can be obtained using the \COLOUR qualifier. A colour vector, colr, is expected as the first parameter. The length of colr should be the same as the number of contours requested.

Contour labels

By default, contours are labeled with the actual contour level, using three significant digits. See the discussion on the \LEGEND qualifier for an alternate contour labeling facility.

Contour label size

The size of the contour labels is %LABSIZ. If labels are not desired on the contours, use the SET command to set %LABSIZ to zero before entering the CONTOUR command.

Contour label separation

The separation between contour labels is controlled with CNTSEP or %CNTSEP, which can be changed with the SET command. If %CNTSEP is set, the separation is a percentage of the height of the window, that is, YUWIND-YLWIND. If CNTSEP is set, the separation is expressed in

inches or centimeters, depending on the units. The default is %CNTSEP = 50.

Saving contour levels and coordinates

The contour levels are automatically stored in a vector named CCONT. If the \COORDINATES qualifier is used, the x and y coordinates of each contour level are stored in matrices named XCNT and YCNT. The number of points stored for each level is the first element of each column. For example, XCNT[1,nc] (=n1) is the number of points making up contour number nc, while XCNT[2:n1+1,nc] and YCNT[2:n1+1,nc] would contain the x and y coordinates of the x contour level. These vectors are then available to the user for plotting and/or manipulation. Each time the CONTOUR command is entered, these vectors are emptied and replaced, so if you wish to keep them, they should be renamed or copied into other vectors.

Legend

By default, the contours are labeled with the actual contour level, using three significant digits. If \NOAXES is used, then no legend is allowed.

If the \LEGEND qualifier is used, then the contours are labeled with an integer index and the list of indices corresponding to the actual contour levels, the legend, is plotted along the right side of the axes. If areas and/or volumes are requested, using \AREAS and/or \VOLUMES, these values will also appear in the legend.

If you zoom in on a contour plot, by setting the axis scales beforehand, and you use the \PARTIAL qualifier as well as the \AREAS qualifier, then the areas will be percentages of the area currently showing on the graph. By default, the areas are percentages of the total area.

To add more contours to a contour plot, re-issue the same contour command with the \CONTINUE qualifier and the legend will be continued from where it left off. You must have used the \NORESET qualifier on the previous CONTOUR command if you intend to use \CONTINUE on a succeeding CONTOUR command.

Legend size

The size of the legend characters is LEGSIZ. The value of LEGSIZ, or %LEGSIZ, can be changed with the SET command. The default value of %LEGSIZ is 1.6

Axis relocation

The legend requires the right end of the x-axis to be set to 75% of the window, that is, %XUAXIS is set to 75. The value of %XUAXIS can be changed with the SET command.

By default, %XUAXIS is reset to its former value after the CONTOUR command is finished. If the \normale not be reset.

Polar coordinates

By default, the vectors \mathbf{x} and \mathbf{y} are assumed to represent Cartesian coordinates. If the \POLAR qualifier is used, \mathbf{x} and \mathbf{y} are assumed to represent polar coordinates, with \mathbf{x} the radial component and \mathbf{y} the angular component, in degrees. The values are converted internally to rectangular coordinates, and the vectors are returned unchanged.

Axes

By default, axes are drawn for the contour plot. If the contour plot is to be overlayed on an existing set of axes, use the \NOAXES qualifier and no axes will be drawn. The axis scales will be left at their current values.

Scattered points

```
Syntax CONTOUR x y z nctr { min { inc }}
CONTOUR\SPECIFIC x y z lvls

Qualifiers \SPECIFIC, \POLAR, \LEGEND, \COLOURS, \PARTIAL \RESET, \AXES, \COORDINATES

Defaults \NOSPECIFIC, \NOPOLAR, \NOLEGEND, \NOCOLOURS, \NOPARTIAL, \RESET, \AXES, \NOCOORDINATES
```

If z is a vector, the vectors x and y are assumed to represent a scattered set of coordinates, where z[i] is the altitude corresponding to the coordinate location (x[i],y[i]). The vectors x and y must be entered if z is a vector.

Contours are computed by successive solution of quintic polynomial equations. The irregularly distributed data points are organized as triangles and the partial derivatives at each point are estimated from the function values of the neighboring points.

Areas and volumes cannot be calculated from scattered data.

Matrix data

Suppose that v is a matrix which has n columns and m rows. The vectors x and y are optional, and if entered, are used for scaling the axes. Each matrix element, v[i,j], is associated with the coordinates (x[j],y[i]). The length of x must be greater than or equal to x and the length of y must be greater than or equal to y.

If x and y are not entered, x defaults to the set [1;2;3;...;n], and y defaults to the set [1;2;3;...;m], so that matrix element m[i,j] is associated with the coordinates (j,i).

Minimum and maximum contour coordinates

The minimum and maximum x value for each contour are automatically stored in vectors named CXMIN and CXMAX; the minimum and maximum y value for each contour are automatically stored in vectors named CYMIN and CYMAX. These vectors are then available to the user for plotting and/or manipulation. Each time the CONTOUR command is entered, these vectors are emptied and replaced, so if you wish to keep them, they should be renamed or copied into other vectors.

Volume

If the \VOLUMES qualifier is used, the volume contained within each contour is calculated as a percentage of the total volume. The volume percentages are automatically stored in a vector named CVOLM. Each time the CONTOUR command is entered, this vector is emptied and replaced, so if you wish to keep it, it should be renamed or copied into another vector.

Area

If the \AREAS qualifier is used, the area contained within each contour is calculated as a percentage of the total area. The area percentages are automatically stored in a vector named CAREA. Each time the CONTOUR command is entered, this vector is emptied and replaced, so if you wish to keep it, it should be renamed or copied into another vector.

Area and volume calculation

The areas and volumes are calculated in the following way. Two dimensional, four point linear interpolation is used to calculate a fine mesh overlayed on the matrix, see Figure 2.1. Suppose the matrix has n columns and m rows and the size of the fine mesh is n_1 columns by m_1 rows. The total area is $n_1 \times m_1$ and the total volume is the sum of the interpolated values. Each point of the fine mesh is tested against the contour levels, and if a mesh point has a value greater than the contour level, a one is binned for the area vector and the mesh point value is binned for the volume vector. Finally, the area and volume vectors are normalized by conversion to percentages.

Figure 2.1: Interpolating a fine mesh on the contours of a matrix

Interpolation size

Suppose the matrix has n columns and m rows. The total size of the fine mesh, n_1 columns by m_1 rows, is defined by the following:

$$n_1 = (n-1) \times i_x + 1$$

 $m_1 = (m-1) \times i_y + 1$

where $i_x - 1$ is the number of interpolation points between matrix points in the x-direction, and $i_y - 1$ is the number of interpolation points between matrix points in the y-direction. The defaults are as below:

$$i_{x} \begin{vmatrix} 10 & \text{if} & n < 20 \\ 5 & \text{if} & 20 \leq n < 50 \\ 3 & \text{if} & 50 \leq n < 100 \\ 2 & \text{if} & 100 \leq n \end{vmatrix} \begin{vmatrix} 10 & \text{if} & m < 20 \\ 5 & \text{if} & 20 \leq m < 50 \\ 3 & \text{if} & 50 \leq m < 100 \\ 2 & \text{if} & 100 \leq m \end{vmatrix}$$

To over-ride these defaults, use the \INTERPSIZE qualifier, and enter ntrp as the first parameter. Both i_x and i_y will be set to ntrp.

Matrix boundary

By default, the boundary of the matrix is outlined within the axes. If this boundary is not desired, use the \NOBORDER qualifier.

```
COPY xin { xin1 ... } xout { xout1 ... } { IFF expression }
Syntax
Qualifiers \APPEND, \INDEX
Defaults
          \NOAPPEND, \NOINDEX
Examples COPY X Y Z XX YY ZZ
          COPY\APPEND X XX
          COPY X[1:10] Y[20:11:-1]
          COPY\INDEX X Y Z XX YY ZZ IFF (X>2)
```

The COPY command copies a subset of vector xin, into another vector, xout. By default, if xout already exists, the COPY command overlays the new data on the old.

Multiple input vectors can be entered, but there must be an output vector for each input vector. Vector xinI is copied into xoutI.

Conditional copy

```
COPY xin1 { xin2 ... } xout1 { xout2 ... } IFF expression
Syntax
```

If the keyword IFF is used, an expression is expected as the next, the last, parameter. In this case, all the input vectors must have the same length. Index ranges on the input vectors are not allowed. The expression does not have to involve any of the input vectors, but it must result in a one dimensional array which has the same length as the input vectors.

xinI[j] is copied into xoutI if and only if the j^{th} element of the expression is true. The expression is said to be true if its value is non-zero, and false if its value is zero.

By default, the elements of xinI are copied in order into xoutI, that is, if the expression

dictates that n elements of xinI are to be copied into xoutI then these will become the first n elements of xoutI.

If the \INDEX qualifier is used, then xinl[j] is copied to xoutl[j] if and only if the j^{th} element of the expression is true. If the j^{th} element of the expression is false, xoutl[j] is left as is, or is set to zero if j is greater than the original length of xoutl.

Unconditional copy

```
Syntax COPY xin1 { xin2 ... } xout1 { xout2 ... }
```

If no expression is entered, then index ranges may be used on the input and/or the output vectors. In some cases, the COPY command is equivalent to an assignment. For example:

```
COPY XX[10:1:-2] X[1:10:2] is equivalent to X[1:10:2]=XX[10:1:-2].
```

Appending with copy

If the \APPEND qualifier is used, the copied elements of xinI are appended onto the end of xoutI. If xoutI does not exist, COPY\APPEND is the same as COPY.

DCL

Syntax DCL

The DCL command is only relevant for VMS. The DCL command enters DCL mode by spawning a subprocess. To return to the program, type the DCL command RETURN. When in DCL mode, any VMS command may be entered, for example, edit a file, run a program.

The first time the DCL command is entered, a subprocess is spawned, which can take some time. If the word RETURN is typed, the subprocess is not destroyed and a subsequent DCL command will attach to this subprocess. Attaching to a subprocess is very fast. If the word LOGOFF is typed, the subprocess is destroyed, so that a subsequent DCL command will have to spawn a new subprocess.

UNIX equivalent

For UNIX users, just type control-z to suspend the program, then type bg to put the program into the background. To return to PHYSICA, type fg.

DEALIAS

Syntax DEALIAS ALL

DEALIAS aliascommand

The DEALIAS command allows the user to eliminate aliases that were created with the ALIAS command. If the keyword ALL is entered, all aliases will be eliminated. To display all aliases, enter the ALIAS command with no parameters.

DEFAULTS

Syntax DEFAULTS

Qualifiers \INITIALIZE, \WINDOWS

Defaults initialization file not executed, windows reset

Examples DEFAULTS

DEFAULTS\INIT

The DEFAULTS command resets the original PHYSICA defaults.

Initialization file

If the \INITIALIZE qualifier is used, the initialization script file is executed after the standard defaults have been set. That command script is also executed automatically at the time PHYSICA is run.

It is possible to have individualized sets of PHYSICA defaults by means of initialization script files. Create a script file and assign its name *before* running the program.

VMS: The file assigned to the logical name PHYSICA\$INIT is executed.

\$ DEFINE PHYSICA\$INIT your_initfile

You could include this assignment in your DCL login command file.

UNIX: The file assigned to the environment variable PHYSICA_INIT is executed.

% setenv PHYSICA_INIT your_initfile

If PHYSICA_INIT is undefined, the file .physicarc in the current directory

is executed. If this file doesn't exist, the file \$HOME/.physicarc is executed.

No further action is taken if this file doesn't exist.

Reset windows

By default, the windows are reset to their original definitions, see Table 2.67 on page 274. If the \-\windows qualifier is used, the windows will be left with their current definitions.

Default values

The PHYSICA keywords and their default values shown in Table 2.24 on page 63. These defaults are the standard GPLOT defaults with the exceptions as listed in Table 2.7.

| NLXINC= 2 | NLYINC = 2 | no plotting symbol |
|--------------|--------------|--------------------|
| NSXINC = 1 | NSYINC = 1 | autoscaling on |
| %XLAXIS = 15 | %YLAXIS = 15 | |
| %XUAXIS=95 | %YUAXIS = 90 | FONT = TSAN |
| %XNUMSZ= 3 | %YNUMSZ = 3 | |

Table 2.7: Exceptions to the standard GPLOT defaults

The line types are reset to their original specifications, as shown in Table 2.8 in centimeters. See the SET LINES command on page 228 for information on changing the line type definitions. See the DISPLAY command for information on how to view examples of the line types.

| line type | p1 | p2 | рЗ |
|-----------|------|------|------|
| 1 | 0.00 | 0.00 | 0.00 |
| 2 | 0.07 | 0.00 | 0.00 |
| 3 | 0.50 | 0.30 | 0.00 |
| 4 | 0.50 | 0.30 | 0.10 |
| 5 | 0.30 | 0.30 | 0.00 |
| 6 | 0.30 | 0.30 | 0.10 |
| 7 | 0.20 | 0.20 | 0.00 |
| 8 | 0.20 | 0.20 | 0.05 |
| 9 | 0.05 | 0.20 | 0.00 |
| 10 | 0.05 | 0.30 | 0.00 |

Table 2.8: Line type defaults in centimeters

The hatch patterns are reset to their original specifications, as shown in Table 2.9 in centimeters. See the SET HATCH command on page 228 for information on changing the hatch pattern definitions. See the DISPLAY command for information on how to view examples of the hatch patterns.

DENSITY

```
Syntax DENSITY { x y } v

Qualifiers \POLAR, \PARTIAL, \DERIV, \PROFILE, \XPROFILE, \YPROFILE, \BORDER, \AXES, \LOG

Defaults \-POLAR, \-PARTIAL, \-DERIV, \-PROFILE, \BORDER, \AXES, \-LOG

Examples DENSITY M

DENSITY X Y Z
```

| pattern | spac | | |
|---------|------|------|-------|
| number | 1 | 2 | angle |
| 1 | 0.01 | | 0 |
| 2 | 0.01 | | 90 |
| 3 | 0.05 | | 0 |
| 4 | 0.05 | | 90 |
| 5 | 0.10 | | 0 |
| 6 | 0.10 | | 90 |
| 7 | 0.20 | | 45 |
| 8 | 0.20 | | -45 |
| 9 | 0.20 | 0.10 | 45 |
| 10 | 0.20 | 0.10 | -45 |

Table 2.9: Hatch pattern defaults in centimeters

The DENSITY command produces a density plot for either data contained in a matrix or a scattered set of points contained in three vectors.

If the \LOG qualifier is used, the base 10 logarithm of the data is used for the plot. If the \LEGEND qualifier is also used, the scales on the legend will be in the form 10^n . The data is divided into levels whose boundaries are always integral powers of 10. The number of levels will vary depending on the original data.

Density plot types

There are five types of density plot available. The default, requiring no special qualifier, is solid filled regions in colour. Other types are chosen by using the appropriate qualifier. Refer to Table 2.10.

| density plot types | required qualifier |
|-------------------------------------|------------------------------|
| solid filled regions in colour | none (default) |
| random points | $\backslash \mathtt{RANDOM}$ |
| dithering with points (grey scales) | \POINTS |
| diffusion with points (grey scales) | \DIFFUSION |
| scaled rectangles | \BOXES |

Table 2.10: Density plot types and their required qualifiers

Axes

By default, axes are drawn for the density plot. If the density plot is to be overlayed on an

existing set of axes, use the \NOAXES qualifier and no axes will be drawn. The axis scales will be left at their current values.

Matrix boundary

The \boxnown BORDER qualifier is valid only if matrix data is entered. By default, the rectangular boundary of the matrix is outlined within the axes. If you do not want this boundary to be drawn, use the \noberdelta BORDER qualifier.

Zooming in

By default, the entire range of possible density levels will be used to determine the minimum and maximum density levels. If the \PARTIAL qualifier is used, the minimum and maximum density levels will be determined by the region contained within the axes. To zoom in on a particular region for more detail, pre-set the axis scales, using the SCALES command, before entering the DENSITY command.

Derivatives

If the \DERIV qualifier is used, the derivative of the data is used for plotting instead of the raw data itself.

Profiles

The qualifiers \PROFILE, \XPROFILE and \YPROFILE are valid only if matrix data is entered.

If the \XPROFILE qualifier is used, the columns of the matrix are summed, the sums are normalized to be between 0 and 1, and a histogram of the normalized sums is drawn horizontally across the top of the graph.

If the \mathbb{YPROFILE} qualifier is used, the rows of the matrix are summed, the sums are normalized to be between 0 and 1, and a histogram of the normalized sums is drawn vertically along the right side of the graph.

If the \PROFILE qualifier is used, both the horizontal and vertical profiles are drawn.

When a profile is drawn, the axis borders must also be set to allow space for the profiles, that is, %XUAXIS is set to 65% if a legend is present or 85% if there is no legend, and %YUAXIS is set to 80.

By default, %XUAXIS and %YUAXIS are reset to their former values. If the \NORESET qualifier is used, the axis locations are not reset.

Polar coordinates

By default, the vectors \mathbf{x} and \mathbf{y} are assumed to represent Cartesian coordinates. If the \POLAR qualifier is used, \mathbf{x} and \mathbf{y} are assumed to represent polar coordinates, with \mathbf{x} the radial component and \mathbf{y} the angular component, in degrees. The values are converted internally to rectangular coordinates, and the vectors are returned unchanged.

Solid filled regions

```
Syntax DENSITY { x y } v { p1 p2 }
Qualifiers \POLAR, \LEGEND, \PARTIAL, \DERIV, \PROFILE, \XPROFILE, \YPROFILE,
\BORDER, \HISTOGRAM, \RESET, \AXES

Defaults if v is a matrix: x = [1;2;3;...], y = [1;2;3;...],
\p1 = 0, p2 = 1, \NOPOLAR, \BORDER, \AXES, \NOLEGEND, \NOPROFILE,
\RESET
```

Solid filled regions in colour is the default density type. No qualifiers are needed to produce this type of drawing. The range of values of the matrix is divided into eight (8) equal levels, and a different colour is associated with each level. By default, a value is interpolated at each pixel location within the matrix region so as to give smoothly joined regions.

PostScript output

For PostScript output, set the POSTRES keyword to the appropriate resolution for your hard-copy device, using the SET command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark inkjet printer has a resolution of 360 dpi. A "good" picture can be obtained with POSTRES = 180 and LINTHK = 2.

Input variables

If v is a vector, the parameters x and y are expected and must be vectors. x and y are assumed to represent a scattered set of points, where v[i] is the altitude corresponding to the location (x[i],y[i]). A matrix is interpolated on these scattered points by means of a Thiessen triangulation of the plane. The minimum length of the three vectors x, y, and v will be used.

If v is a matrix, the parameters x and y default to [1;2;3;...], but if entered they must be vectors. Each matrix element, m[i,j], is associated with the coordinates (x[j],y[i]). The length of x must be greater than or equal to the number of columns of v and the length of y must be greater than or equal to the number of rows. The vectors x and y are used for scaling the axes.

Not interpolated solid fill

If the \HISTOGRAM qualifier is used, each data location is represented by a rectangle of colour, centred on the data location. The regions are not smoothly joined.

Changing the range of values

The optional parameters p1 and p2 can be used to broaden or shrink the range of data values. If v_{max} is the maximum value of the data and v_{min} is the minimum value of the data, the full colour range will be from a minimum of $min = \mathtt{p1} \times (v_{max} - v_{min}) + v_{min}$ to a maximum of $max = \mathtt{p2} \times (v_{max} - v_{min}) + v_{min}$. If v is a data value and if $v < \mathtt{p1} \times (v_{max} - v_{min}) + v_{min}$, that data value is treated as v_{min} . If $v > \mathtt{p1} \times (v_{max} - v_{min}) + v_{min}$, that data value is treated as v_{max} . The default values are: $\mathtt{p1} = 0$ and $\mathtt{p2} = 1$.

Legend

If the \LEGEND qualifier is used, a legend is drawn along the right side of the axes. The legend requires the right end of the x-axis to be set to 75% of the window, that is, %XUAXIS is set to 75.

When a y-profile is drawn, using the \PROFILE qualifier or the \YPROFILE qualifier, the right edge of the axis box must allow space for the profile as well as a possible legend. If a y-legend profile and a legend are present, then %XUAXIS is set to 65. If a y-legend profile is present but not a legend, then %XUAXIS is set to 85.

The value of %XUAXIS can be changed with the SET command. By default, %XUAXIS is reset to its former value after the DENSITY command. If the \normale normal qualifier is used, the axis location is not reset.

The numeric legend entries are written using the LEGFRMT format and with height given by LEGSIZ, both of which are changed with the SET command. The default values are: LEGFRMT = 1PE10.3 and %LEGSIZ = 1.6.

Random points

To obtain the random points type of density plot, use the \RANDOM qualifier.

A value is interpolated at every pixel location within the data region and the value is then normalized to lie between 0 and 1. This normalized value is compared to a randomly generated number. That pixel location is lit up if the square of the normalized value is greater than the random number.

PostScript output

For PostScript output, set the POSTRES keyword to the appropriate resolution for your hard-copy device, using the SET command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark inkjet printer has a resolution of 360 dpi. A "good" picture can be obtained with POSTRES = 180 and LINTHK = 2.

Input variables

If v is a vector, the parameters x and y are expected and must be vectors. x and y are assumed to represent a scattered set of points, where v[i] is the altitude corresponding to the location (x[i],y[i]). A matrix is interpolated on these scattered points by means of a Thiessen triangulation of the plane. The minimum length of the three vectors x, y, and v will be used.

If v is a matrix, the parameters x and y default to [1;2;3;...], but if entered they must be vectors. Each matrix element, m[i,j], is associated with the coordinates (x[j],y[i]). The length of x must be greater than or equal to the number of columns of v and the length of y must be greater than or equal to the number of rows. The vectors x and y are used for scaling the axes.

Random points in colour

If the \COLOUR qualifier is used with \RANDOM , the range of data values is divided into eight equal levels and a different colour is associated with each level. The pixel is lit in the colour corresponding to the relative size of the interpolated value.

Changing the range of values

The optional parameters p1 and p2 can be used to broaden or shrink the range of data values. If v_{max} is the maximum value of the data and v_{min} is the minimum value of the data, the full colour range will be from a minimum of $min = \mathtt{p1} \times (v_{max} - v_{min}) + v_{min}$ to a maximum of $max = \mathtt{p2} \times (v_{max} - v_{min}) + v_{min}$. If v is a data value and if $v < \mathtt{p1} \times (v_{max} - v_{min}) + v_{min}$, that data value is treated as v_{min} . If $v > \mathtt{p1} \times (v_{max} - v_{min}) + v_{min}$, that data value is treated as v_{max} . The default values are: $\mathtt{p1} = 0$ and $\mathtt{p2} = 1$.

Example

```
R=MOD([0:143],4)+1
SORT\UP R
T=MOD([0:143],36)*10
DENSITY\RANDOM R*COSD(T) R*SIND(T) EXP(-R/2)*COSD(180*(R-1))
```

Diffusion

```
Syntax DENSITY\DIFFUSION { x y } v { p1 p2 }
Qualifiers \POLAR, \PARTIAL, \DERIV, \PROFILE, \XPROFILE, \YPROFILE, \BORDER, \RESET, \AXES

Defaults if v is a matrix: x = [1;2;3;...], y = [1;2;3;...], \NOPOLAR, \BORDER, \AXES, \NOPROFILE, \RESET
p1 = 0, p2 = 1
```

To obtain the diffusion type of density plot, use the \DIFFUSION qualifier.

Diffusion is a form of digital halftoning. A threshold is fixed at $\frac{1}{2}$. Data values are interpolated at each pixel location, and then normalized to be between 0 (white) and 1 (black). The resulting binary output value is compared with the original grey level value. The difference is called the error for that location. The signal consisting of past error values is passed through an error filter to produce a correction factor to be added to future input values. Thus, errors are diffused over a weighted neighborhood.

PostScript output

For PostScript output, set the PostRes keyword to the appropriate resolution for your hard-copy device, using the Set command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark InkJet printer has a resolution of 360 dpi. A "good" picture can be obtained with Postres = 180 and Linthk = 2.

Input variables

If v is a vector, the parameters x and y are expected and must be vectors. x and y are assumed to represent a scattered set of points, where v[i] is the altitude corresponding to the location (x[i],y[i]). A matrix is interpolated on these scattered points by means of a Thiessen triangulation of the plane. The minimum length of the three vectors x, y, and v will be used.

If v is a matrix, the parameters x and y default to [1;2;3;...], but if entered they must be

vectors. Each matrix element, m[i,j], is associated with the coordinates (x[j],y[i]). The length of x must be greater than or equal to the number of columns of v and the length of y must be greater than or equal to the number of rows. The vectors x and y are used for scaling the axes.

Changing the range of values

The optional parameters p1 and p2 can be used to broaden or shrink the range of data values. If v_{max} is the maximum value of the data and v_{min} is the minimum value of the data, the full colour range will be from a minimum of $min = \text{p1} \times (v_{max} - v_{min}) + v_{min}$ to a maximum of $max = \text{p2} \times (v_{max} - v_{min}) + v_{min}$. If v is a data value and if $v < \text{p1} \times (v_{max} - v_{min}) + v_{min}$, that data value is treated as v_{max} . If $v > \text{p1} \times (v_{max} - v_{min}) + v_{min}$, that data value is treated as v_{max} . The default values are: p1 = 0 and p2 = 1.

Example

```
R=MOD([0:143],4)+1
SORT\UP R
T=MOD([0:143],36)*10
DENSITY\DIFFUSION R*COSD(T) R*SIND(T) EXP(-R/2)*COSD(180*(R-1))
```

Dithering patterns

```
DENSITY\POINTS { x y } v { p1 p2 }

DENSITY\POINTS\DITHER d { x y } v { p1 p2 }

DENSITY\POINTS\LEVELS lvl { x y } v { p1 p2 }

DENSITY\POINTS\LEVELS\DITHER d lvl { x y } v { p1 p2 }

Qualifiers \POLAR, \LEGEND, \PARTIAL, \DERIV, \PROFILE, \XPROFILE, \YPROFILE, \BORDER, \RESET, \AXES, \DITHER, \CONTOURS, \LEVELS, \AREAS, \VOLUMES, \LINES, \EQUALLY_SPACED

Defaults if v is a matrix: x = [1;2;3;...], y = [1;2;3;...], d = [1;1;2;1;2;2;3;2;3;3;4;3;4;4;5;5;6;6;0;0], \NOPOLAR, \BORDER, \NOLEGEND, \NOPROFILE, \RESET p1 = 0, p2 = 1, ten equally spaced contour levels
```

To obtain the dithering pattern type of density plot, use the \POINTS qualifier.

By default, the range of data values is divided into ten (10) equally spaced levels and a different dithering pattern is associated with each level. A value is interpolated at every pixel location within the bounds of the data region to determine the level for that point. The dithering pattern for that level then determines whether that pixel is to be lit up. Thus, the

boundaries of the data are divided up into different dithering pattern regions.

PostScript output

For PostScript output, set the PostRes keyword to the appropriate resolution for your hard-copy device, using the Set command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark inkjet printer has a resolution of 360 dpi. A "good" picture can be obtained with Postres = 180 and Linthk = 2.

Input variables

If v is a vector, the parameters x and y are expected and must be vectors. x and y are assumed to represent a scattered set of points, where v[i] is the altitude corresponding to the location (x[i],y[i]). A matrix is interpolated on these scattered points by means of a Thiessen triangulation of the plane. The minimum length of the three vectors x, y, and v will be used.

If v is a matrix, the parameters x and y default to [1;2;3;...], but, if entered, they must be vectors. Each matrix element, m[i,j], is associated with the coordinates (x[j],y[i]). The length of x must be greater than or equal to the number of columns of v and the length of y must be greater than or equal to the number of rows. The vectors x and y are used for scaling the axes.

Changing the range of values

The optional parameters p1 and p2 can be used to broaden or shrink the range of data values. If v_{max} is the maximum value of the data and v_{min} is the minimum value of the data, the full colour range will be from a minimum of $min = \text{p1} \times (v_{max} - v_{min}) + v_{min}$ to a maximum of $max = \text{p2} \times (v_{max} - v_{min}) + v_{min}$. If v is a data value and if $v < \text{p1} \times (v_{max} - v_{min}) + v_{min}$, that data value is treated as v_{min} . If $v > \text{p1} \times (v_{max} - v_{min}) + v_{min}$, that data value is treated as v_{max} . The default values are: p1 = 0 and p2 = 1.

Dithering pattern definition

The default dithering pattern vector is:

```
[ 1;1; 2;1; 2;2; 3;2; 3;3; 4;3; 4;4; 5;5; 6;6; 0;0 ]
```

A user defined dithering pattern can be entered by using the \DITHER qualifier and entering the dithering pattern vector, d, as the first parameter.

The dithering pattern is determined by pairs of numbers from d, so the number of dithering

patterns defined by d is $\frac{1}{2}$ the length of d.

For pattern number i, every $d[2 \times i - 1]^{th}$ pixel is lit up horizontally, and every $d[2 \times i]^{th}$ pixel is lit up vertically. For example, if d[1] = 1 and d[2] = 1, then for level 1 every pixel is lit up, while if d[3] = 2 and d[4] = 3, then for level 2 every second pixel is lit up horizontally and every third pixel is lit up vertically.

Legend

If the \LEGEND qualifier is used, a legend is drawn along the right side of the axes. The legend requires the right end of the x-axis to be set to 75% of the window, that is, %XUAXIS is set to 75.

When a y-profile is drawn, using the \PROFILE qualifier or the \YPROFILE qualifier, the right edge of the axis box must allow space for the profile as well as a possible legend. If a y-legend profile and a legend are present, then %XUAXIS is set to 65. If a y-legend profile is present but not a legend, then %XUAXIS is set to 85.

The value of %XUAXIS can be changed with the SET command. By default, %XUAXIS is reset to its former value after the DENSITY command. If the \normale normaliser is used, the axis location is not reset.

The numeric legend entries are written using the LEGFRMT format and with height given by LEGSIZ, both of which are changed with the SET command. The default values are: LEGFRMT = 1PE10.3 and %LEGSIZ = 1.6.

Contours

By default, a contour line is drawn around the boundary of each dithering pattern region. If the \NOLINES qualifier is used, then these contour lines will not be drawn.

If the qualifier \CONTOURS is used, an automatically created vector named DENS\$CONT will be made which will contain the boundary values of each region. If there are N regions, the length of DENS\$CONT will be N+1.

User specified contour levels

A specific set of contour levels can be entered by using \LEVELS and entering a vector of contour level values, 1v1, as the first parameter, unless the \DITHER qualifier is also used, in which case the contour level vector should be the second parameter. If both are used, and the length of the dithering vector is N, the length of the level vector must be $\frac{N}{2}-1$. Suppose that v_{min} and v_{max} are the minimum and maximum of the data v. The level vector must be

strictly monotonically increasing, with $lvl[1] > v_{min}$ and $lvl[\#] < v_{max}$.

The \EQUALLY_SPACED qualifier only applies to the case of a dithering type density plot with legend, where the user supplies the contour levels, for example:

```
DENSITY\POINTS\LEVELS\LEGEND\EQUALLY_SPACED lvl x y m
```

If the \EQUALLY_SPACED qualifier is used, the legend boxes will all be the same size, irregardless of the values specified in the levels vector, lvl.

Areas and volumes

If the \AREAS qualifier is used, an automatically created vector named DENS\$AREA will be made which will contain the percentage areas contained within each region. If the \VOLUMES qualifier is used, an automatically created vector named DENS\$VOLM will be made which will contain the percentage volumes contained within each region. If there are N regions, the length of DENS\$AREA and DENS\$VOLM will both be N. Also, the sum of the elements of each of these vectors will always be 100, that is, sum(DENS\$AREA[j],j,1:N) = 100.

Example

```
R=MOD([0:143],4)+1
SORT\UP R
T=MOD([0:143],36)*10
D=[ 1;1; 2;2; 4;4; 7;7; 11;11; 0;0 ]
DENSITY\LEGEND\POINTS\DITHER D R*COSD(T) R*SIND(T) EXP(-R/2)*COSD(180*(R-1))
```

Boxes

```
Syntax DENSITY\BOXES { x y } v { p1 p2 { q1 q2 { r }}}
Qualifiers \POLAR, \PARTIAL, \DERIV, \PROFILE, \XPROFILE, \YPROFILE, \BORDER, \RESET, \AXES

Defaults if v is a matrix: x = [1;2;3;...], y = [1;2;3;...], p1 = 0, p2 = 1, q1 = 0, q2 = 1, r = 1, \NOPOLAR, \BORDER, \AXES, \NOPROFILE, \RESET
```

To obtain the scaled rectangles type of density plot, use the \BOXES qualifier.

Input variables

If v is a vector, the parameters x and y are expected and must be vectors. x and y are as-

sumed to represent a scattered set of points. A box is drawn, centred at location (x[i],y[i]) with relative size determined by v[i]. No internal matrix is interpolated with the scaled rectangle type of density plot. The minimum length of the three vectors x, y, and v will be used.

If v is a matrix, the parameters x and y default to [1;2;3;...], but if entered they must be vectors. A box is drawn, centred at location (x[j],y[i]) with relative size determined by v[i,j]. The length of x must be greater than or equal to the number of columns of v and the length of y must be greater than or equal to the number of rows.

Delimiting the range of values

The optional parameters p1 and p2 can be used to select a window of values from within the box size range, min to max, as defined above. Suppose that v is the data value at (x,y). A box is drawn at (x,y) if and only if $p1 < \frac{v - min}{max - min} < p2$. The default values are: p1 = 0 and p2 = 1.

Accentuating a range of values

The optional parameters q1 and q2 can be used to accentuate a range of values. If v_{max} is the maximum value of the data and v_{min} is the minimum value of the data, the full box size range will be from a minimum of $min = \text{q1} \times (v_{max} - v_{min}) + v_{min}$ to a maximum of $max = \text{q2} \times (v_{max} - v_{min}) + v_{min}$. The default values are: q1 = 0 and q2 = 1.

Box size scale factor

The optional parameter r is a scale factor which controls the size of the boxes. For each box, the width and height is multiplied by r. The default value is: r = 1.

Filled boxes

The boxes can be filled. Use the SET FILL command to change the fill type and pattern. See Table 2.55 on page 233 for a description of the interpretations of the FILL keyword.

Examples

The following script produces Figure 2.2.

```
X=[ 1; 0; 1; 0; .2; .3; .5; .8]
Y=[ 5; 5; 0; 0; 1;1.5; 2.5; 4]
Z=[ 10; 10; 10; 10; -100; 10; -100; 500]
GRID\XYOUT X Y Z M XOUT YOUT
DENSITY\BOXES\PROFILES XOUT YOUT M
```

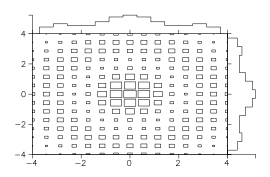


Figure 2.2: An example of a box type density plot with both x and y profiles

The following script produces Figure 2.3.

```
.2; .3;
   1;
        0; 1; 0;
                                .5; .8]
Y=[5;5;0;0;
                      1;1.5; 2.5;
                                     4]
Z=[ 10; 10; 10; 10; -100; 10; -100; 500]
GRID\XYOUT X Y Z M XOUT YOUT
WINDOW 5
LABEL\XAXIS 'DENSITY\BOXES XOUT YOUT M O 1 O 1'
DENSITY\BOXES XOUT YOUT M O 1 O 1
WINDOW 7
LABEL\XAXIS 'DENSITY\BOXES XOUT YOUT M .5 1 0 1'
DENSITY\BOXES XOUT YOUT M .5 1 0 1
WINDOW 6
LABEL\XAXIS 'DENSITY\BOXES XOUT YOUT M O 1 .5 1'
DENSITY\BOXES XOUT YOUT M 0 1 .5 1
WINDOW 8
LABEL\XAXIS 'DENSITY\BOXES XOUT YOUT M .5 1 .5 1'
DENSITY\BOXES XOUT YOUT M .5 1 .5 1
```

The following script produces Figure 2.4.

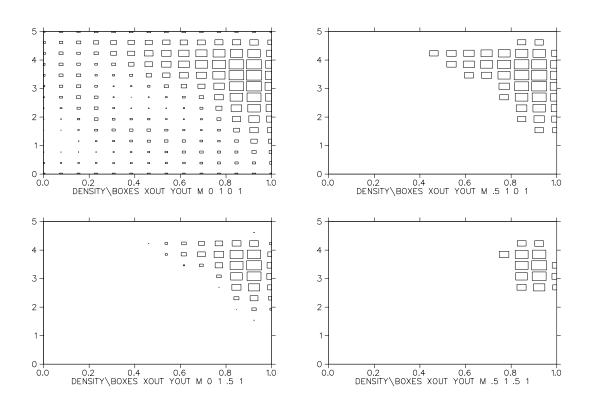
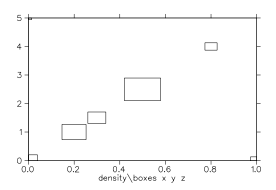


Figure 2.3: Examples of box type density plot with accentuated and delimited values

```
X=[ 1; 0; 1; 0;
                      .2; .3;
                                .5;
                                     .8]
Y=[ 5; 5;
            0; 0;
                      1;1.5;
                                     4]
                               2.5;
Z=[ 10; 20; 30; 40;
                      50; 40;
                               70; 30]
WINDOW 5
LABEL\XAXIS 'density\boxes x y z'
DENSITY\BOXES X Y Z
WINDOW 7
GRID\XYOUT X Y Z M XOUT YOUT
LABEL\XAXIS 'density\boxes xout yout m'
DENSITY\BOXES XOUT YOUT M
```



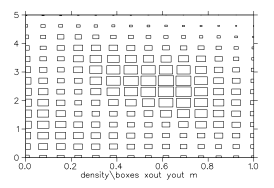


Figure 2.4: Box type density plots with scattered points and with a matrix

DESTROY

```
Syntax DESTROY v1 { v2 ... } { IFF expression }

Qualifier \EXPAND

Default \-EXPAND

Examples DESTROY X Y Z A B C M1 M2 M3 T1 T2 T3
DESTROY *V M1 M2 *T
DESTROY X Y Z IFF (X>=2)&(X<=4)
DESTROY X[1:10] Y[2:20:2] T[3][A:B] Z
```

The DESTROY command eliminates subsets of vectors or string variables, or it destroys scalars or matrices. Up to twenty-nine (29) variable names may be entered.

Unconditional

```
Syntax DESTROY v1 { v2 ... }

Qualifier \EXPAND

Default \-EXPAND
```

When scalars or matrices are entered, they are completely destroyed. Index ranges are not allowed on scalars or matrices. These variables are simply eliminated. By default, variable names are not constructed or expanded.

If vectors are entered with no index ranges, they are entirely eliminated. Subsets of vectors or of string variables may be eliminated by including an index range.

Expand names

By using the \EXPAND qualifier, you can enter names that must be constructed, or string variables that can be expanded.

For example, suppose that scalar I has the value 2, and the string variable TXT has the value $^{\prime}X2^{\prime}$.

```
DESTROY\EXPAND 'X'//RCHAR(I) will destroy X2
DESTROY TXT will destroy TXT
DESTROY\EXPAND TXT will destroy X2
```

Index ranges

Index ranges are *not* allowed on scalars or matrices. Index ranges are *not* allowed if an expression is entered.

Subsets of vectors or string variables may be eliminated by including an index range. For example:

```
DESTROY X[5:25:5] Y Z T[1:10]
```

will eliminate elements 5, 10, 15, 20, and 25 from vector X and eliminate characters 1 through 10 of string variable T. The variables X and T will be compressed. Variables Y and Z will be entirely destroyed.

Classes of variables

There are various keywords to simplify the elimination of groups of variables. Refer to Table 2.11. These keywords cannot be used with an expression, but they can be used along with specific variable names.

Conditional

```
Syntax DESTROY v1 { v2 ... } IFF expression
```

| keyword | result |
|---------|--|
| * | all variables will be destroyed |
| *V | all vectors will be destroyed |
| *S | all scalars will be destroyed |
| *M | all matrices will be destroyed |
| *T | all string variables will be destroyed |

Table 2.11: Keywords used to destroy entire classes of variables

If the keyword IFF is used, an expression is expected as the next, the last, parameter. Elements of *vectors only* can be eliminated conditional on the value of an expression. Matrices, scalars and string variables are not allowed, and index ranges on the input vectors are also *not* allowed with a conditional expression. The expression does not have to involve any of the input vectors, but it must result in a one dimensional array which has the same length as the input vectors.

vI[j] is eliminated if and only if the j^{th} element of the expression is true. The expression is said to be true if its value is non-zero, and false if its value is zero.

Examples

```
DESTROY X[5:25:5] Y Z *M T[1:10] TXT[3][2:5]
```

will destroy elements 5, 10, 15, 20, and 25 from vector X; eliminate characters 1 through 10 of string variable T; and eliminate characters 2 through 5 of the third string of array string variable TXT. The variables X, T, and TXT[3] will be compressed. Variables Y and Z will be entirely destroyed. All matrices will be eliminated.

It is possible to delete from a vector all elements that have a value between M and N, assuming that M and N are scalars.

```
DESTROY X IFF (X>M)&(X<N)
```

The expression is evaluated for each element of X and if the expression is true, the corresponding element is deleted from the vector X. Other vectors can also be entered. For example:

```
DESTROY X Y Z IFF (X>M)&(X<N)
```

If the j^{th} element of the expression is true, then elements X[j], Y[j] and Z[j] will be deleted. In this way, sets of data can be kept together.

DEVICE

Syntax DEVICE keyword

Defaults initial graphics hardcopy device: HP LaserJet 150 dpi

Examples DEVICE HPLASER 300

DEVICE HPPLOTTER

DEVICE\COLOUR POSTSCRIPT DEVICE\GREY POSTSCRIPT A

The DEVICE command is used to select a graphics hardcopy output device. If a device type is chosen, as opposed to entering OFF or ON, then the graphics will be cleared. The initial default graphics hardcopy device is an HP LaserJet bitmap at 150 dpi resolution.

The graphics hardcopy device determines the world coordinate plotting units. The device should be chosen before opening an EDGR file.

ON and OFF

If the user is not interested in graphics hardcopy, it is recommended that hardcopy generation be turned OFF, since graphics on the monitor is noticeably faster when a hardcopy is not made.

If DEVICE OFF is entered, output to the hardcopy bitmap or file will be disabled. The graphics will not be cleared, but, while the output is disabled, no hardcopy output will be available. To re-enable output to a previously chosen device type, use the DEVICE ON command. The device type that was previously chosen will again be available. The graphics will not be cleared.

Device keywords

If the DEVICE command is entered with no parameters, the current graphics hardcopy device is displayed. A table of devices and their corresponding code numbers is also displayed, and you can then enter your choice, either by name or by code number.

| bitmap | | vector | | display | |
|--------------------------|-----|-------------------------------------|-------|---------|----|
| devices | | plotters | | files | |
| | | | | | |
| HPLaserJet {100 150 300} | 1-3 | <pre>HPPlotter {A B C D E}</pre> | 12-16 | VT640 | 36 |
| Inkjet $\{1 2 3 4 5\}$ | 4-8 | Houston {A B C D E} | 17-21 | VT241 | 37 |
| HPThinkJet | 9 | LNO3+ | 22 | Cit467 | 38 |
| La100 | 10 | Imagen | 23 | TK4010 | 39 |
| Printronix | 11 | GKS | 24 | TK4107 | 40 |
| | | ${\tt PostScript \{A B C D E A4\}}$ | 25-30 | PT100G | 41 |
| | | Roland {A B C D E} | 31-35 | Seiko | 42 |

HPLaserJet

```
egin{array}{ll} {\it Syntax} & {\it DEVICE HPLASERJET \{ dpi \} } \\ {\it Default} & {\it dpi} = 150 \\ \end{array}
```

The optional parameter, dpi, is only applicable with HPLaserJet. It controls the resolution of the hardcopy, specifying the dots per inch. The valid resolutions are: dpi = 100, 150, 300. If dpi is not entered, 150 is assumed.

Refer to Table 2.12 to see the plotting units for the HPLaserJet bitmap graphics hardcopy devices.

| orientation | units | horizontal | vertical |
|-------------|-------------|------------|----------|
| LANDSCAPE | centimeters | 27.94 | 21.59 |
| | inches | 11.00 | 8.50 |
| PORTRAIT | centimeters | 21.59 | 27.94 |
| | inches | 8.50 | 11.00 |

Table 2.12: Plotting units for HPLASERJET devices

InkJet

The HP PaintJet and the LJ250 are colour bitmap devices that allow for a single plot to cover from one (1) to a maximum of five (5) continuous pages. To choose the number of pages, use the optional parameter np. If graphics is being monitored on a colour monitor, the colours used on the INKJET will be the same as on the monitor, except that black and white are interchanged.

Refer to Table 2.13 to see the plotting units for the InkJet bitmap graphics hardcopy devices.

| | pages = 1 pages > | | pages = 1 | | > 1 |
|-------------|-------------------|------------|-----------|------------|----------|
| orientation | units | horizontal | vertical | horizontal | vertical |
| LANDSCAPE | centimeters | 26.67 | 19.05 | 27.94 | 20.32*np |
| | inches | 10.50 | 7.50 | 11.00 | 8.00*np |
| PORTRAIT | centimeters | 19.05 | 26.67 | 20.32*np | 27.94 |
| | inches | 7.50 | 10.50 | 8.00*np | 11.00 |

Table 2.13: Plotting units for INKJET devices

Other bitmap devices

Syntax DEVICE PRINTRONIX

DEVICE LA100 DEVICE THINKJET

The PRINTRONIX, LA100, and THINKJET are bitmap devices.

Refer to Table 2.14 to see the plotting units for the PRINTRONIX, LA100, and THINKJET bitmap graphics hardcopy devices.

| orientation | units | horizontal | vertical |
|-------------|-------------|------------|----------|
| LANDSCAPE | centimeters | 25.00 | 19.00 |
| | inches | 9.84 | 7.48 |
| PORTRAIT | centimeters | 19.00 | 25.00 |
| | inches | 7.48 | 9.84 |

Table 2.14: Plotting units for PRINTRONIX, LA100, and THINKJET devices

PostScript devices

Syntax DEVICE POSTSCRIPT { A | B | C | D | E | A4 } Qualifiers \FLIP, \COLOUR, \GREY

Defaults \-FLIP, \-COLOUR, \-GREY, paper size A

The optional parameter refers to paper sizes. Refer to Table 2.15. To see the plotting units for the PostScript devices, refer to Table 2.16.

Resolution

| paper | | |
|-------|----------------------|----------------------|
| size | centimeters | inches |
| A | 21.59×27.64 | 8.50×11.00 |
| В | 27.94×43.18 | 11.00×17.00 |
| C | 43.18×55.88 | 17.00×22.00 |
| D | 55.88×86.36 | 22.00×34.00 |
| E | 86.36×111.76 | 34.00×44.00 |
| A4 | 21.00×29.70 | 8.27×11.69 |

Table 2.15: PostScript paper sizes

The resolution of your PostScript hardcopy output can be changed with the SET POSTRES command. The default value for POSTRES is 180 dpi (dots per inch). This applies to dot filled text characters and to dot types of DENSITY plots.

Upside down drawings

If the \FLIP qualifier is used, the drawing will come out upside down on the paper. This is not mirror image. This feature is included to facilitate the insertion of PostScript plots into TeX or \mathbb{M}EX documents.

Colour

The \COLOUR qualifier only applies to PostScript output. The \COLOUR qualifier means colour changes will be inserted into the PostScript output. The default is \-COLOUR.

Grey scale

The \GREY qualifier only applies to PostScript output. The \GREY qualifier means colour changes will be inserted into the PostScript output as grey scales. The default is \-GREY.

Pen plotters

```
Syntax DEVICE HPPLOTTER { A | B | C | D | E }
DEVICE HOUSTON { A | B | C | D | E }
DEVICE ROLAND { A | B | C | D | E }

Defaults paper size A
```

The optional parameter refers to paper sizes. Refer to Table 2.17.

Refer to Table 2.18 to see the plotting units for the pen plotter type graphics hardcopy

| paper | | LANDSCAPE | | PORTRAIT | |
|-------|-------------|------------|----------|------------|----------|
| size | units | horizontal | vertical | horizontal | vertical |
| Α | centimeters | 25.00 | 19.00 | 19.00 | 25.00 |
| | inches | 9.84 | 7.48 | 7.48 | 9.84 |
| В | centimeters | 40.64 | 25.40 | 25.40 | 40.64 |
| | inches | 16.00 | 10.00 | 10.00 | 16.00 |
| С | centimeters | 53.34 | 40.64 | 40.64 | 53.34 |
| | inches | 21.00 | 16.00 | 16.00 | 21.00 |
| D | centimeters | 83.82 | 53.34 | 53.34 | 83.82 |
| | inches | 33.00 | 21.00 | 21.00 | 33.00 |
| E | centimeters | 109.22 | 83.82 | 83.82 | 109.22 |
| | inches | 43.00 | 33.00 | 33.00 | 43.00 |
| A4 | centimeters | 27.16 | 18.46 | 18.46 | 27.16 |
| | inches | 10.69 | 7.27 | 7.27 | 10.69 |

Table 2.16: Plotting units for POSTSCRIPT devices

| paper | | |
|-------|----------------------|-------------------|
| size | centimeters | inches |
| A | 21.59×27.64 | 8.5×11.0 |
| В | 27.94×43.18 | 11.0×17.0 |
| С | 43.18×55.88 | 17.0×22.0 |
| D | 55.88×86.36 | 22.0×34.0 |
| E | 86.36×111.76 | 34.0×44.0 |

Table 2.17: Pen plotter paper sizes

devices.

| paper | | LANDSCAPE | | PORTRAIT | |
|-------|-------------|------------|----------|------------|----------|
| size | units | horizontal | vertical | horizontal | vertical |
| A | centimeters | 25.00 | 19.00 | 19.00 | 25.00 |
| | inches | 9.84 | 7.48 | 7.48 | 9.84 |
| В | centimeters | 40.64 | 25.40 | 25.40 | 40.64 |
| | inches | 16.00 | 10.00 | 10.00 | 16.00 |
| С | centimeters | 53.34 | 40.64 | 40.64 | 53.34 |
| | inches | 21.00 | 16.00 | 16.00 | 21.00 |
| D | centimeters | 83.82 | 53.34 | 53.34 | 83.82 |
| | inches | 33.00 | 21.00 | 21.00 | 33.00 |
| E | centimeters | 109.22 | 83.82 | 83.82 | 109.22 |
| | inches | 43.00 | 33.00 | 33.00 | 43.00 |

Table 2.18: Plotting units for pen plotter devices

Other vector plotters

Syntax DEVICE LN03+

DEVICE IMAGEN

The LNO3+ and IMAGEN are vector plotters.

Refer to Table 2.19 to see the plotting units for the LNO3+ and IMAGEN vector plotter graphics hardcopy devices.

| orientation | units | horizontal | vertical |
|-------------|-------------|------------|----------|
| LANDSCAPE | centimeters | 25.40 | 19.05 |
| | inches | 10.00 | 7.50 |
| PORTRAIT | centimeters | 19.05 | 25.40 |
| | inches | 7.50 | 10.00 |

Table 2.19: Plotting units for LNO3+ and IMAGEN devices

GKS metafiles

Syntax DEVICE GKS

GKS refers to GKS metafiles, which are available only at sites where PHYSICA is linked with a local GKS library. In this case, there will likely be an interpreter program which allows the metafile to be replayed onto various printers and terminals.

Refer to Table 2.20 to see the plotting units for the GKS graphics metafiles.

| orientation | units | horizontal | vertical |
|-------------|-------------|------------|----------|
| LANDSCAPE | centimeters | 25.40 | 19.05 |
| | inches | 10.00 | 7.50 |
| PORTRAIT | centimeters | 19.05 | 25.40 |
| | inches | 7.50 | 10.00 |

Table 2.20: Plotting units for GKS graphics metafiles

Display files

| Syntax | DEVICE | VT640 |
|--------|--------|--------|
| | DEVICE | VT241 |
| | DEVICE | CIT467 |
| | DEVICE | TK4010 |
| | DEVICE | TK4107 |
| | DEVICE | PT100G |
| | DEVICE | GR1105 |

If a monitor type is chosen as a hardcopy device, a "display file" will be created. The graphics will be redisplayed when this file is typed, in DCL mode, on the appropriate terminal type. Any graphics that was drawn after choosing the display file will be replayed on the monitor screen, including any clearing of the graphics. The name of the file that is created will be displayed. The file is named after the plotter type.

Refer to Table 2.21 to see the plotting units for the display file type of graphics output.

| orientation | units | horizontal | vertical |
|-------------|-------------|------------|----------|
| LANDSCAPE | centimeters | 27.94 | 21.59 |
| | inches | 11.00 | 8.50 |
| PORTRAIT | centimeters | 21.59 | 27.94 |
| | inches | 8.50 | 11.00 |

Table 2.21: Plotting units for display file graphics output

| DIG | ITI | ZE |
|-----|-----|----|
|-----|-----|----|

```
Syntax DIGITIZE { xout yout { codes }}
```

The DIGITIZE command digitizes points off of a graph that is attached to a digitizing pad. To make use of the digitizer, simply secure the graph to the pad, enter the DIGITIZE command, and follow the directions.

Digitizing pad types

The only digitizing pad type that is currently supported is the Digi-Pad, Type 5A, made by GTCO Corporation, attached to a terminal.

Optional output variables

If the two optional variable names, xout and yout, are entered, then two vectors will be created. Any recorded points will be saved in these two vectors, with the horizontal axis values stored in xout and the vertical axis values stored in yout. If the optional variable name, codes, is entered, then a vector will be created with a code number saved in this vector for each recorded point. See the following table for the meanings of these code numbers.

| | code |
|-----------------|------|
| recorded point | 1 |
| marked point | 2 |
| connected point | 3 |

Preparing for digitizing data

When the DIGITIZE command is entered, the graphics screen will be cleared and instructions will be displayed on the monitor screen. You will first be asked to give names for the four digitizer's crosshair buttons. Enter four labels, one for each of the mouse buttons, with a maximum of 15 characters per label, separated by blanks or commas. You will then be asked to type each button to enable the program to coordinate the labels with the buttons.

Now you will be asked to place the digitizer's crosshair on some point of the graph where you know the graph coordinates and to press a button. Usually, this point is the lower left corner of the graph. Then you will be asked to enter the x and y graph coordinates of this point. Repeat this process for two more points, ensuring that the third point is not collinear with the first two points. Usually, the points one enters are the lower right corner and the upper right corner of the graph. These three points define the transformation for the graph and the angle of the axes.

Digitizing data

Now you are ready to digitize data off of the graph. It is a good idea to position the crosshair on the lower left and upper right corners and check the values there.

The action that is taken at any time is determined by the mouse button or the keyboard key that is typed. See Table 2.22 for the mouse buttons and their corresponding definitions,

and see Table 2.23 for the keyboard keys and their corresponding definitions.

| button | definition |
|--------|---|
| 1 | digitize a point and display the x and y values, |
| | do not save these values |
| 2 | digitize and display values as above; record these values in the two (optional) vectors xout and yout |
| 3 | digitize, display, and record values as above; place a small marker at the chosen point |
| 4 | digitize, display, record, and mark values as above; also connect this point to the last recorded point by drawing a line segment |

Table 2.22: Mouse button definitions when digitizing data

| key | definition |
|-----|--|
| F | write the values that have been recorded so far to a file, enter the |
| | file name when asked |
| ? | display the menu of key control codes |
| / | clear the alphanumeric monitor screen |
| Q | quit; the screen will be cleared |

Table 2.23: Keyboard key definitions when digitizing data

DISABLE

| Svntax | DISABLE | kawword |
|--------|---------|---------|
| Svinax | DISADLE | Keyword |

The DISABLE command allows you to disable certain features denoted by keyword. Use the ENABLE command to re-enable those features.

Graphics window borders

Syntax DISABLE BORDER

Default enabled

Disabling BORDER means that the rectangular borders that delimit the hardcopy page boundary and the window edges will not be drawn. These rectangles do not appear on any hardcopies, but may be considered to interfere with the graphics monitor display.

Broadcast messages

Syntax DISABLE BROADCAST

Default the terminal state when PHYSICA is invoked

The DISABLE BROADCAST command is only relevant for VMS. The DISABLE BROADCAST command is equivalent to the DCL command:

\$ SET TERMINAL/NOBROADCAST

This prevents broadcast messages from being accepted by the terminal monitor. If the user is not interested in receiving broadcast messages, for example, "New mail", then it is recommended that broadcast mode be turned off.

Confirmation requests

Syntax DISABLE CONFIRM

Default enabled

If CONFIRM is disabled, no confirmation will be requested from the TEXT, FIGURE, LEGEND FRAME, and WINDOW commands. The CONFIRM setting can be over-ridden on a specific command by using the \CONFIRM (or the \NOCONFIRM) qualifier.

Echoing from scripts

Syntax DISABLE ECHO

Default disabled

Disabling ECHO means that commands that are entered via a command script file will not be displayed on the monitor screen as they are executed.

Local effect

If the ENABLE ECHO command is encountered within a script, echoing is done only while within that script. For example, suppose you have echoing disabled at the keyboard entry level and you execute a script which has ENABLE ECHO within it. Subsequent lines that are read from that script will be echoed, but when that script is finished executing, echoing will be disabled again.

Saving a variable's history

Syntax DISABLE HISTORY

Default enabled

Disabling HISTORY means that when a variable is altered, it's history will not be updated. A variable's history is displayed with the SHOW command. This feature was included because variables altered within large DO loops can have their histories updated so many times that virtual memory limits will be exceeded. Even if HISTORY is disabled, new variables will still have initial history lines.

Journaling input and output

Syntax DISABLE JOURNAL

DISABLE JOURNAL\MACRO

Default enabled, journal file: PHYSICA. JOURNAL, script journaling disabled

DISABLE JOURNAL means that the journal file is to be closed. Subsequent journaling of program output and user input will be disabled. DISABLE JOURNAL\MACRO means that journaling of script commands and output is disabled, but interactively entered input and resultant output will still be enabled. Enter ENABLE JOURNAL, to reopen the last journal file that was open and append subsequent journal entries to this file. Enter the JOURNAL command to open a new journal file.

Prompting

Syntax DISABLE PROMPTING

Default enabled

Disabling PROMPTING means that commands will not prompt you for input when you leave something out or enter some incorrect parameter.

Replotting

Syntax DISABLE REPLOT

Default enabled

Disabling REPLOT means that commands that subsequent graphs and text will not be stored for replotting. See the REPLOT for more information. The REPLOT setting can be over-ridden on a specific command by using the \REPLOT (or the \NOREPLOT) qualifier.

X Window graphics replay

Syntax DISABLE REPLAY

Default enabled

Disabling REPLAY means that the X Window System graphics replay storage is disabled, hence

subsequent graphics will not be available for replay. For example, if the graphics window size is changed, the graphics displayed in that window will not be re-displayed. The virtual memory space for storing graphics vectors is allocated dynamically. Enter DISABLE REPLAY to save virtual memory space for other uses, such as large data arrays. The REPLAY keyword only applies if an X Window type monitor is being used. See the MONITOR command for more information.

Input line recall shell

Syntax DISABLE SHELL

Default enabled

Disabling SHELL means that the input line recall shell is turned off. You will not be able to recall keyboard input lines when the shell has been disabled. When the SHELL is re-enabled, the buffer of input lines will be available again. It is useful to disable the shell when reading data across a network, since the terminal I/O may become corrupted if the shell is enabled. See the BUFFER command, page 11, for more information on the input line recall shell.

Stacking commands in a file

Syntax DISABLE STACK

Default disabled

Disabling STACK means that subsequently entered commands will not be written to the specified stack file. See the STACK command for more information.

DISPLAY

```
Syntax DISPLAY 'message'
DISPLAY FONT { fontname }
DISPLAY SPECIAL
DISPLAY HATCH
DISPLAY LINES
DISPLAY PCHAR
DISPLAY MENU keyword
```

The DISPLAY command either displays a message on the monitor screen, or interprets the command parameter as a keyword and draws a corresponding table or displays a corresponding list.

Display a message

Syntax DISPLAY 'message'

If the command parameter is not a recognized keyword, it is assumed to be a message to be displayed on the monitor screen. The message can be a literal string, enclosed in quotes, or a string variable. The message will be displayed even if ECHO is disabled.

Note: What is written to the journal file when the DISPLAY string command is encountered in a macro:

If JOURNAL is enabled, the DISPLAY string command will write the string to the journal file the same way it is written to the monitor screen.

If JOURNAL\MACRO is enabled, the DISPLAY command itself will also be written to the journal file.

Font table

```
Syntax DISPLAY FONT { fontname }
```

If the keyword FONT is entered, and if a fontname is entered, the font table for the specified fontname is drawn. If no fontname is entered, the table for the current font is drawn. To view a list of the font names, enter the GET FONT command. See Figure 2.5 for an example font table.

Special characters

```
Syntax DISPLAY SPECIAL
```

If the keyword SPECIAL is entered, the table of special character names that can be included as format commands in strings is drawn. This table is shown in Figure 2.6.

Hatch fill patterns

```
Syntax DISPLAY HATCH
```

If the keyword HATCH is entered, the currently defined hatch fill patterns with their corresponding numbers are drawn. For example, see Figure 2.7.

The hatch fill patterns can be used for filling text characters, histogram bars, pie charts, etc. See the SET HATCH command for information on redefining the hatch patterns.

Line types

```
Syntax DISPLAY LINES
```

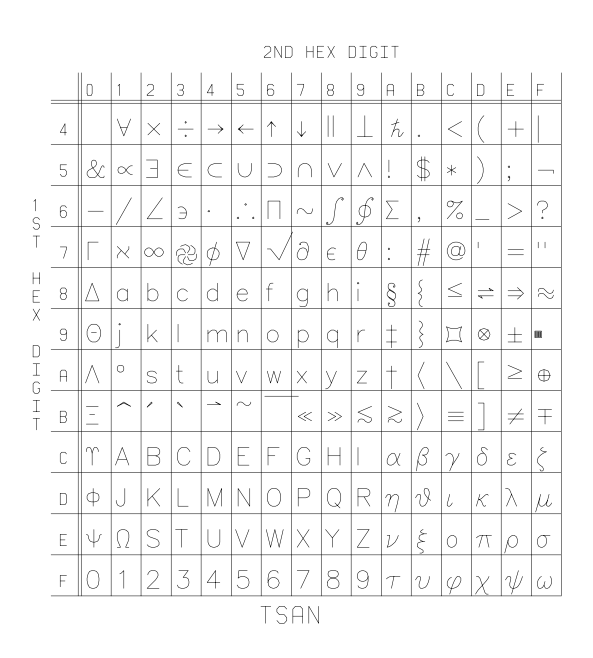


Figure 2.5: An example of a font table produced by the <code>DISPLAY FONT</code> command

| Name | Uppercase | Lowercase | Name U | ppercase | Lowercase | Name Up | percas | Lowercase |
|-----------|------------------------------------|----------------------------|------------|----------------|--------------|------------|-------------------|-------------------|
| Alpha | A | α | Overline | _ | _ | Degree | o | 0 |
| Beta | В | β | Leftarrow | ← | ← | Dagger | † | † |
| Gamma | Γ̈́ | γ | Uparrow | ↑ | ↑ | Ddagger | + | |
| Delta | Δ | δ | Downarrow | į. | j | S | 8 | 8 |
| Epsilon | Ē | € | Rightarrow | | , → | Langle | l | ί |
| Zeta | \bar{z} | ξ | Parallel | II. | | Rangle |), |), |
| Eta | H | $\overset{\circ}{\eta}$ | Perp | Ï | | Riharpoons | , = | , = |
| Theta | Θ | $\overset{\cdot }{	heta }$ | Mid | Ī | 1 | Vector | _ | |
| Tota | Ĭ | i | Squarebull | et 🖢 | | Nea | _ | _ |
| Карра | K | κ | Box | | | Therefore | | |
| Lambda | Λ | λ | Sum | $\bar{\Sigma}$ | Σ | Angle | Ž | Ž |
| Mu | M | μ | 00 | 4 | _ | Vee | V | _ V |
| Nu | N | ν | Prod | Π | П | Wedge | ٨ | ^ |
| Xi | Ξ | Ę | 11.00 | | | Cdot | | • |
| Omicron | ō | ò | Int | | ſ | Infty | ∞ | 00 |
| Pi | Π | au | | J | J | In | € | € |
| Rho | P | ρ | | 1 | | Ni | Э | Э |
| Sigma | Σ | σ | Surd | | \checkmark | Propto | ~ | ∝ |
| Tau | $\overline{\overline{\mathbf{T}}}$ | au | | V | | Exists | 3 | 3 |
| Upsilon | Ť | $\stackrel{\cdot}{v}$ | Oint | ∮ | ∮ | Forall | ¥ | A |
| Phi | Φ | $\overset{\circ}{\phi}$ | Plus | + | + | Neg | ≠ | ≠ |
| Chi | X | χ | Minus | _ | _ | Equiv | = | = |
| Psi | Ψ | $\hat{m{\psi}}$ | Pm | ± | ± | Approx | ≈ | ≈ |
| Omega | Ω | $\overset{r}{\omega}$ | Mo | Ŧ | Ŧ | Sim | ~ | ~ |
| Vartheta | v | บิ | Times | × | × | Lt | < | < |
| Varphi | φ | φ | Div | ÷ | ÷ | Gt | > | > |
| Varepsilo | | ε | Oplus | () | | LI | « | « |
| Aleph | × × | × | Otimes | 8 | 8 | Ga | >> | >> |
| Tlogo | ₩ | ₩ | Сар | \cap | \cap | Lsimeq | ≲ | ≲ |
| Nabla | ∇ | ∇ | Subset | <u> </u> | <u> </u> | Gsimeq | ≳ | ≳ |
| Partial | ð | ð | Сир | Ū | Ū | Lea | \leq | ≤ |
| Hbar | ħ | ħ | Supset | 5 | 5 | Gea | ≥ | ≥ |

Figure 2.6: The table of special characters

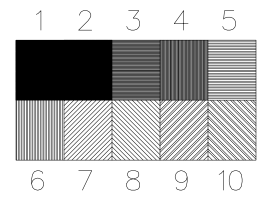


Figure 2.7: An example of the hatch fill patterns

If the keyword LINES is entered, the currently defined line types with their corresponding numbers are drawn. For an example of the default line types, see Figure 2.8.



Figure 2.8: An example of the default line types

The line types can be used for drawing data curves on a graph. See the SET LINES command for information on redefining the line types.

Plotting symbols

Syntax DISPLAY PCHAR

If the keyword PCHAR is entered, the plotting symbols with their corresponding numbers will be drawn. These are the special plotting symbols that can be chosen with the SET PCHAR command. See Figure 2.9 for an example.

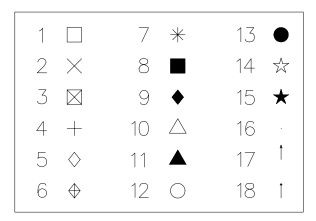


Figure 2.9: The special plotting symbols

Menus

```
Syntax DISPLAY MENU PHYSICA
DISPLAY MENU FULL
DISPLAY MENU SHORT
DISPLAY MENU GENERAL
DISPLAY MENU XAXIS
```

DISPLAY MENU YAXIS

If the keyword MENU is entered, a table of values for special PHYSICA keywords or tables of values for GPLOT keywords is displayed. These menus are displayed in alphanumeric mode on the monitor screen. Refer to **Appendix A** for descriptions of how each of the GPLOT keywords can affect a graph. The plot characteristics: MASK, ALIAS, PMODE, PTYPE, and ERRBAR should *not* be changed in PHYSICA, as these are altered internally by various commands.

PHYSICA

The PHYSICA specific keywords are described in the SET command section, page 228, and the GET command section, page 107.

Table 2.24 shows the default values for the PHYSICA specific keywords.

```
= 2.10
VERSION
VERSIONDATE = January 16, 1998
CNTSEP =
             10.795
                        50.000% | LABSIZ =
                                                   0.324
                                                              1.500%
LEGSIZ =
              0.345
                         1.600%
                                    LEGFRMT = 1PE10.3
ERRFILL =
              0.000
                                    ARROWID =
                                                   0.150
                                    ARROTYP =
ARROLEN =
              0.200
                                                   0.000
TENSION =
              1.000
                                  | FILL
                                                       0
SPEED
                 20
                                    SEED
                                                      12345
POSTRES =
                180
                                    WIDTH
                                                      80
              0.000
XPREV
                                    YPREV
                                                   0.000
NCURVES =
                  0
                                    WRAP
                                    MAXHISTORY =
SHOWHISTORY =
                  5
                                                      5
Current font = TSAN
Current plotting symbol= 0
```

Table 2.24: The PHYSICA keyword menu of default values

FULL

The full GPLOT keyword menu is displayed when the DISPLAY MENU FULL command is entered. This menu requires the monitor to be set to a width of 132, which is done automatically when the command is entered. If you want the monitor to be set back to a width of 80 after issuing the MENU FULL command, use the SET WIDTH command, page 228. Table 2.25 on page 64 displays the PHYSICA default values for the GPLOT keywords.

| MASK | = | -4.000 | | BOX | = | 1.000 | | CHARSZ | = | 0.190 | 1.000% | CHARA | = | 0.000 | 0.000% |
|--------|---|---------|----------|--------|---|---------|----------|--------|---|--------|---------|--------|---|--------|---------|
| PMODE | = | 1.000 | | ALIAS | = | 1.000 | | HISTYP | = | 0.000 | | LINTYP | = | 1.000 | |
| PTYPE | = | 0.000 | | COLOUR | = | 1.000 | | CURSOR | = | 1.000 | | ERRBAR | = | 0.000 | |
| XLWIND | = | 0.000 | 0.000% | XUWIND | = | 25.400 | 100.000% | XLAXIS | = | 3.810 | 15.000% | XUAXIS | = | 24.130 | 95.000% |
| NXDIG | = | 5.000 | | NXDEC | = | -1.000 | | XPOW | = | 0.000 | | XPAUTO | = | 1.000 | |
| XNUMSZ | = | 0.572 | 3.000% | XLABSZ | = | 0.572 | 3.000% | XTICL | = | 0.381 | 2.000% | XTICS | = | 0.190 | 1.000% |
| XTICA | = | 270.000 | -90.000% | XCROSS | = | 0.000 | | XMIN | = | 0.000 | | XMAX | = | 10.000 | |
| NLXINC | = | 2.000 | | NSXINC | = | 1.000 | | XAUTO | = | 2.000 | | XITICL | = | 0.572 | 3.000% |
| XITICA | = | 270.000 | -90.000% | XNUMA | = | 0.000 | 0.000% | XTICTP | = | 1.000 | | BOTTIC | = | 1.000 | |
| BOTNUM | = | 0.000 | | TOPTIC | = | -1.000 | | TOPNUM | = | 0.000 | | NXGRID | = | 0.000 | |
| XAXIS | = | 1.000 | | XAXISA | = | 0.000 | | XLOG | = | 0.000 | | XZERO | = | 0.000 | |
| YLWIND | = | 0.000 | 0.000% | YUWIND | = | 19.050 | 100.000% | YLAXIS | = | 2.857 | 15.000% | YUAXIS | = | 17.145 | 90.000% |
| NYDIG | = | 5.000 | | NYDEC | = | -1.000 | | YPOW | = | 0.000 | | YPAUTO | = | 1.000 | |
| YNUMSZ | = | 0.572 | 3.000% | YLABSZ | = | 0.572 | 3.000% | YTICL | = | 0.381 | 2.000% | YTICS | = | 0.190 | 1.000% |
| YTICA | = | 90.000 | 90.000% | YCROSS | = | 0.000 | | YMIN | = | 0.000 | | YMAX | = | 10.000 | |
| NLYINC | = | 2.000 | | NSYINC | = | 1.000 | | YAUTO | = | 2.000 | | YITICL | = | 0.572 | 3.000% |
| YITICA | = | 90.000 | 90.000% | YNUMA | = | -90.000 | -90.000% | YTICTP | = | 1.000 | | LEFTIC | = | 1.000 | |
| LEFNUM | = | 0.000 | | RITTIC | = | -1.000 | | RITNUM | = | 0.000 | | NYGRID | = | 0.000 | |
| YAXIS | = | 1.000 | | YAXISA | = | 90.000 | | YLOG | = | 0.000 | | YZERO | = | 0.000 | |
| TXTHIT | = | 0.572 | 3.000% | TXTANG | = | 0.000 | | XLOC | = | 12.700 | 50.000% | YLOC | = | 9.525 | 50.000% |
| | | | | | | | | | | | | | | | |

Table 2.25: The full menu of GPLOT keywords, with values in centimeters

SHORT

If the DISPLAY MENU SHORT command is entered, a short summary table is displayed. See Table 2.26 on page 65. This does not require the terminal to be set to a width of 132.

XAXIS

If the DISPLAY MENU XAXIS command is entered, a table of the x-axis characteristics is displayed. See Table 2.27 on page 66. This does not require the terminal to be set to a width of 132.

YAXIS

If the DISPLAY MENU YAXIS command is entered, a table of the *y*-axis characteristics is displayed. See Table 2.28 on page 67. This does not require the terminal to be set to a width of 132.

GENERAL

If the DISPLAY MENU GENERAL command is entered, a table of the general GPLOT keywords is displayed. See Table 2.29 on page 68. This does not require the terminal to be set to a width of 132.

| MASK | = | -4.000 | | CHARSZ | = | 0.190 | | 1.000% |
|----------|---|--------|----------|--------|---|--------|---|----------|
| PMODE | = | 1.000 | I | HISTYP | = | 0.000 | | |
| LINTYP | = | 1.000 | LINTHK = | 1.000 | - | COLOUR | = | 1.000 |
| XLWIND | = | 0.000 | 0.000% | XUWIND | = | 25.000 | | 100.000% |
| XLAXIS | = | 3.750 | 15.000% | XUAXIS | = | 23.750 | | 95.000% |
| NXDIG | = | 5.000 | I | NXDEC | = | -1.000 | | |
| XPOW | = | 0.000 | I | XPAUTO | = | 1.000 | | |
| XMIN | = | 0.000 | 0.000 | XMAX | = | 10.000 | | 10.000 |
| NLXINC | = | 2.000 | I | NSXINC | = | 1.000 | | |
| OTUAX | = | 2.000 | I | XLOG | = | 0.000 | | |
| YLWIND | = | 0.000 | 0.000% | YUWIND | = | 19.000 | | 100.000% |
| YLAXIS | = | 2.850 | 15.000% | YUAXIS | = | 17.100 | | 90.000% |
| NYDIG | = | 5.000 | I | NYDEC | = | -1.000 | | |
| YPOW | = | 0.000 | I | YPAUTO | = | 1.000 | | |
| YMIN | = | 0.000 | 0.000 | YMAX | = | 10.000 | | 10.000 |
| NLYINC | = | 2.000 | I | NSYINC | = | 1.000 | | |
| OTUAY | = | 2.000 | I | YLOG | = | 0.000 | | |
| XLOC | = | 12.500 | 50.000% | YLOC | = | 9.500 | | 50.000% |
| | | | + | | | | | |
| KLABEL = | | | | | | | | |
| LABEL = | | | | | | | | |

Table 2.26: The short menu of GPLOT keywords, with values in centimeters

| XLWIND | = | 0.000 | 0.000% | | XUWIND | = | 25.000 | 100.000% |
|--------|---|---------|----------|-----|--------|---|---------|----------|
| XLAXIS | = | 3.750 | 15.000% | | XUAXIS | = | 23.750 | 95.000% |
| XAXIS | = | 1.000 | | | XAXISA | = | 0.000 | |
| | | | | -+- | | | | |
| NXDIG | = | 5.000 | | | NXDEC | = | -1.000 | |
| XPOW | = | 0.000 | | 1 | XPAUTO | = | 1.000 | |
| NLXINC | = | 2.000 | | - | NSXINC | = | 1.000 | |
| | | | | -+- | | | | |
| XNUMSZ | = | 0.570 | 3.000% | | XNUMA | = | 0.000 | 0.000% |
| XTICL | = | 0.380 | 2.000% | | XTICS | = | 0.190 | 1.000% |
| XTICA | = | 270.000 | -90.000% | | XITICA | = | 270.000 | -90.000% |
| XITICL | = | 0.570 | 3.000% | | XTICTP | = | 1.000 | |
| BOTTIC | = | 1.000 | | - | BOTNUM | = | 0.000 | |
| TOPTIC | = | -1.000 | | | TOPNUM | = | 0.000 | |
| XMIN | = | 0.000 | | - | XMAX | = | 10.000 | |
| XVMIN | = | 0.000 | | - | XVMAX | = | 10.000 | |
| | | | | -+- | | | | |
| NXGRID | = | 0.000 | | - | XCROSS | = | 0.000 | |
| XLOG | = | 0.000 | | - | XZERO | = | 0.000 | |
| XMOD | = | 0.000 | | - | XLEADZ | = | 0.000 | |
| XOFF | = | 0.000 | | | XLABSZ | = | 0.570 | 3.000% |

Table 2.27: The menu of GPLOT *x*-axis characteristics, with values in centimeters

| YLWIND | = | 0.000 | 0.000% | - | YUWIND | = | 19.000 | 100.000% |
|--------|---|--------|---------|-----|--------|---|---------|----------|
| YLAXIS | = | 2.850 | 15.000% | | YUAXIS | = | 17.100 | 90.000% |
| YAXIS | = | 1.000 | | | YAXISA | = | 90.000 | |
| | | | | -+- | | | | |
| NYDIG | = | 5.000 | | | NYDEC | = | -1.000 | |
| YPOW | = | 0.000 | | | YPAUTO | = | 1.000 | |
| NLYINC | = | 2.000 | | 1 | NSYINC | = | 1.000 | |
| | | | | -+- | | | | |
| YNUMSZ | = | 0.570 | 3.000% | | YNUMA | = | -90.000 | -90.000% |
| YTICL | = | 0.380 | 2.000% | | YTICS | = | 0.190 | 1.000% |
| YTICA | = | 90.000 | 90.000% | | YITICA | = | 90.000 | 90.000% |
| YITICL | = | 0.570 | 3.000% | | YTICTP | = | 1.000 | |
| LEFTIC | = | 1.000 | | | LEFNUM | = | 0.000 | |
| RITTIC | = | -1.000 | | | RITNUM | = | 0.000 | |
| YMIN | = | 0.000 | | | YMAX | = | 10.000 | |
| YVMIN | = | 0.000 | | 1 | YVMAX | = | 10.000 | |
| | | | | -+- | | | | |
| NYGRID | = | 0.000 | | | YCROSS | = | 0.000 | |
| YLOG | = | 0.000 | | | YZERO | = | 0.000 | |
| YMOD | = | 0.000 | | | YLEADZ | = | 0.000 | |
| YOFF | = | 0.000 | | | YLABSZ | = | 0.570 | 3.000% |

Table 2.28: The menu of GPLOT y-axis characteristics, with values in centimeters

| CHARSZ | = | 0.190 | 1.000% | | CHARA | = | 0.000 | 0.000% |
|--------|---|--------|---------|-----|--------|---|-------|---------|
| BOX | = | 1.000 | | 1 | HISTYP | = | 0.000 | |
| LINTYP | = | 1.000 | | 1 | LINTHK | = | 1.000 | |
| COLOUR | = | 1.000 | | 1 | CLIP | = | 1.000 | |
| NUMBLD | = | 0.000 | | 1 | | | | |
| | | | | -+- | | | | |
| CURSOR | = | 1.000 | | | | | | |
| TXTHIT | = | 0.570 | 3.000% | 1 | TXTANG | = | 0.570 | 3.000% |
| XLOC | = | 12.500 | 50.000% | 1 | YLOC | = | 9.500 | 50.000% |
| | | | | -+- | | | | |
| XLABEL | = | | | | | | | |
| YLABEL | = | | | | | | | |
| | | | | -+- | | | | |
| XLABSZ | = | 0.570 | 3.000% | | YLABSZ | = | 0.570 | 3.000% |

Table 2.29: The menu of general GPLOT keywords, with values in centimeters

EDGR

The EDGR command interfaces to the graphical editor. The keyword determines what action is taken by EDGR.

Please refer to the EDGR USER'S GUIDE for details on using the graphical editor.

Open a drawing file

```
Syntax EDGR OPEN { filename }
```

If the keyword OPEN is entered, an EDGR drawing file will be opened. If the file name is not entered, it will be interactively requested. Do *not* give a file extension. For example, just enter FILE, do not enter FILE.extension. All graphics subsequently done will be entered into the drawing file, that is, filename.DWG and filename.DWT, until EDGR CLOSE, or EDGR OPEN again, is entered.

Commands that alter the plotting units should be entered *before* opening an EDGR file:

```
ORIENTATION, SET UNITS, and DEVICE.
```

Note: EDGR has its own hardcopy facility, so it is suggested that the user disable the graphics hardcopy output before opening an EDGR drawing. Use the command: DEVICE OFF.

Edit a drawing file

```
Syntax EDGR EDIT
```

If the keyword EDIT is entered, the graphical editor is invoked which allows you to edit your drawing.

Close a drawing file

```
Syntax EDGR CLOSE
```

If the keyword CLOSE is entered, and if a drawing file has been previously opened with

the EDGR OPEN command, then this command will close that file, no more graphics will be inserted into that file. If no file is currently open, then this command does nothing.

Open a new frame

```
Syntax EDGR FRAME
```

If the keyword FRAME is entered, and if a drawing file has been previously opened with the EDGR OPEN command, this command will open another frame within that file. If no file is currently open, then this command does nothing.

ELLIPSE

```
Syntax ELLIPSE a b cx cy angle
ELLIPSE\FIT xin yin

Qualifiers \FIT, \NPTS, \XYOUT, \REPLOT

Defaults \-FIT, \-NPTS, \-XYOUT, \REPLOT
```

The ELLIPSE command can uniformly populate the perimeter of an ellipse in two ways:

- 1. Given the major axis radius, the minor axis radius, centre coordinates and angle of the major axis
- 2. First fit an ellipse so a certain fraction of the data points are within the ellipse, then determine the major axis radius, the minor axis radius, centre coordinates and angle of the major axis

Output vectors

```
Syntax ELLIPSE\XYOUT a b cx cy angle xout yout ELLIPSE\FIT\XYOUT xin yin xout yout
```

By default, the ellipse perimeter will be plotted automatically. It is assumed that a graph has been drawn already. The ellipse will be overlayed on this graph, with no plotting symbol.

If the \XYOUT qualifier is used, then two output vector names, xout and yout, are expected. No automatic plotting is done, and the horizontal and vertical coordinates of the ellipse perimeter will be placed in these two vectors.

Replotting

The REPLOT command will replot any curves that have been drawn as well as the automati-

cally drawn ellipse, all on a common scale large enough to accommodate all curves.

If the \NOREPLOT qualifier is used, the automatically drawn ellipse will not be stored in the replot buffers, and thus will not be available for replotting.

Number of points

```
Syntax ELLIPSE\NPTS a b cx cy angle n ELLIPSE\FIT\NPTS xin yin n
```

By default, the ellipse perimeter is populated by 260 points. If the \NPTS qualifier is used, the number of points with which to populate the perimeter is expected. This number should be divisible by four (4).

Explicitly defined

```
Syntax ELLIPSE a b cx cy angle

Qualifiers \nPTS, \XYOUT, \REPLOT

Defaults \-NPTS, \-XYOUT, \REPLOT

Examples ELLIPSE\NOREPLOT MAJOR MINOR XCENT YCENT ANG
ELLIPSE\XYOUT\NPTS MAJOR MINOR XC YC ANG N XOUT YOUT
```

By default, the input parameters are assumed to be scalars representing the major axis radius, a, the minor axis radius, b, the coordinates of the centre, cx and cy, and the angle of the ellipse, angle in degrees, measured counter-clockwise from the horizontal.

Parameter order

The order in which the qualifiers appear is irrelevant. The order in which the command parameters appear is fixed: a b xc yc angle { npts } { xout yout }

Fit an ellipse

Syntax ELLIPSE\FIT xin yin

ELLIPSE\FIT\FRACTION xin yin frac

ELLIPSE\FIT\PARAMETERS xin yin a b cx cy angle

Qualifiers \NPTS, \XYOUT, \REPLOT, \FRACTION, \PARAMETERS, \MESSAGES

Defaults \-NPTS, \-XYOUT, \REPLOT, \-FRACTION, \-PARAMETERS, \MESSAGES

Examples ELLIPSE\FIT\FRAC\NPTS XIN YIN FRAC N

ELLIPSE\FIT\XYOUT\NPTS XIN YIN N XO YO ELLIPSE\FIT\XYOUT\FRAC XIN YIN FRAC XO YO

ELLIPSE\FIT\XYOUT\FRAC\NPTS XIN YIN FRAC N XO YO

ELLIPSE\FIT\PARAM\FRAC\NPTS XIN YIN FRAC A B CX CY ANG N

ELLIPSE\FIT\PARAM\XYOUT\FRAC\NPTS XIN YIN F A B CX CY ANG N XO YO

If the \FII qualifier is used, then the first two parameters, xin and yin, are assumed to be vectors which contain data points to which an ellipse is to be fitted.

Parameter order

The order that the qualifiers appear is irrelevant. The order that the command parameters appear is fixed: xin yin { fraction } { a b cx cy angle } { npts } { xout yout }

Method

The major axis radius and the centre are found by least squares fitting a line through the data points. The ratio of the major axis radius to the minor axis radius is found by computing the standard deviations about the major and minor axes. The minimum value of the major axis is found for each point so that the point will be inside the ellipse, then a value for the major axis is picked so as to be greater than or equal to the specified fraction of the values.

Fraction of points within ellipse

Syntax ELLIPSE\FIT\FRACTION xin yin frac

Defaults frac = 0.9

By default, the ellipse is fit to the data points so that it encloses 90% of the data points.

If the \FRACTION qualifier is used, then a scalar, frac, representing the fraction of data points to be within the ellipse will be expected, 0 < frac < 1.

Messages

By default, the major axis radius, the minor axis radius, the coordinates of the centre, and the angle of the fitted ellipse will be displayed on the monitor screen. To suppress this, use the $\backslash \text{-MESSAGES}$ qualifier.

Output parameters

Syntax ELLIPSE\FIT\PARAMETERS xin yin a b cx cy angle

If the \PARAMETERS qualifier is used, output scalar names will be expected to receive the resulting ellipse parameters:

a major axis radius

b minor axis radius

cx and cy x and y coordinates of the centre

angle angle of the major axis,

in degrees, measured counter-clockwise from the horizontal

ENABLE

Syntax ENABLE keyword

The ENABLE command allows you to re-enable features that have been disabled with the DISABLE command.

Graphics window borders

Syntax ENABLE BORDER

Default enabled

Enabling BORDER means that the rectangular borders that delimit the hardcopy page boundary and the window edges will again be drawn. These rectangles do not appear on any hardcopies, but may be considered to interfere with the graphics monitor display.

Broadcast messages

Syntax ENABLE BROADCAST

Default the terminal state when PHYSICA is invoked

The ENABLE BROADCAST command is only relevant for VMS. The ENABLE BROADCAST command is equivalent to the DCL command:

\$ SET TERMINAL/BROADCAST

This allows a broadcast message to be accepted by the terminal. If the user is not interested in receiving broadcast messages, for example, "New mail", then it is recommended that broadcast mode be turned off.

Confirmation requests

Syntax ENABLE CONFIRM

Default enabled

If CONFIRM is enabled, confirmation will be requested from the TEXT, FIGURE, LEGEND FRAME, and WINDOW commands. The CONFIRM setting can be over-ridden on a specific command by using the \nOCONFIRM (or the \CONFIRM) qualifier.

Echoing from scripts

Syntax ENABLE ECHO

Default disabled

Enabling ECHO means that commands that are entered via a command script file will be displayed on the monitor screen as they are executed.

Local effect

If the ENABLE ECHO command is encountered within a script, echoing is done only while within that script. For example, suppose you have echoing disabled at the keyboard entry level and you execute a script which has ENABLE ECHO within it. Subsequent lines that are read from that script will be echoed, but when that script is finished executing, echoing will be disabled again.

Saving a variable's history

Syntax ENABLE HISTORY

Default enabled

When HISTORY is disabled, if a variable is altered, it's history will not be updated. A variable's history is displayed with the SHOW command. This feature was included because variables altered within large DO loops can have their histories updated so many times that virtual memory limits will be exceeded. Even if HISTORY is disabled, new variables will still have initial history lines.

Journaling input and output

Syntax ENABLE JOURNAL

ENABLE JOURNAL\MACRO

Default enabled, journal file: PHYSICA. JOURNAL, script journaling disabled

ENABLE JOURNAL means that subsequent journal entries, that is, program output and user input, will be appended to the last journal file that was opened. By default, program input from script files and resultant output will not be journaled. Use the ENABLE JOURNAL to also journal script input and output. See the JOURNAL command and the DISABLE JOURNAL command for more information.

Prompting

Syntax ENABLE PROMPTING

Default enabled

Enabling PROMPTING means that commands will prompt you for input when you leave something out or enter some incorrect parameter.

Replotting

Syntax ENABLE REPLOT

Default enabled

Enabling REPLOT means that subsequent graphs and text will be stored for replotting. See the REPLOT for more information. The REPLOT setting can be over-ridden on a specific command by using the \NOREPLOT (or the \REPLOT) qualifier.

X Window graphics replay

Syntax ENABLE REPLAY

Default enabled

Enabling REPLAY means that the X Window System graphics replay storage is enabled, hence subsequent graphics will be available for replay. For example, if the graphics window size is changed, the graphics displayed in that window will be re-displayed. The virtual memory space for storing graphics vectors is allocated dynamically. Enter DISABLE REPLAY to save virtual memory space for other uses, such as large data arrays. The REPLAY keyword only applies if an X Window type monitor is being used. See the MONITOR command for more information.

Input line recall shell

Syntax ENABLE SHELL

Default enabled

Disabling SHELL means that the input line recall shell is turned off. You will not be able to recall keyboard input lines when the shell has been disabled. When the SHELL is re-enabled, the buffer of input lines will be available again. It is useful to disable the shell when reading data across a network, since the terminal I/O may become corrupted if the shell is enabled. See the BUFFER command, page 11, for more information on the input line recall shell.

Stacking commands in a file

Syntax ENABLE STACK

Default disabled

Enabling STACK means that subsequently entered commands will again be written to the specified stack file. See the STACK command for more information.

ERASEWINDOW

```
Syntax ERASEWINDOW { n }

Defaults n = current window number
```

The ERASEWINDOW command erases the graphics within a pre-defined window. It will erase the graphics within a window on the monitor screen, and on graphics hardcopy PostScript output, and on graphics hardcopy bitmap output.

This *does not apply* to any other graphics hardcopy output or to EDGR drawings, that is, any erased graphics will still appear on that hardcopy and will still be in an EDGR drawing.

The parameter $\tt n$ refers to a pre-defined window number. If $\tt n$ is not entered, it defaults to the current window number.

EXECUTE

The EXECUTE command reads program input from a file. When the end of file is reached, input will again be expected to be entered from the keyboard, or from the a calling executable file. You may have up to twenty (20) nested executable files. The 'at' sign, @, is equivalent to

'EXECUTE'.

Within script files, it is possible to have labels, GOTO statements, IF blocks, and DO loops.

Environment variables in file names

For UNIX users, it is now possible to use an environment variable in a file name, if the environment variable is preceded by a \$. For example,

```
setenv FILE dum.pcm
physica
@$FILE
```

The environment variable can be just the first part of the filename, for example,

```
setenv FILE dum
physica
@$FILE.pcm
```

Filename extensions

If the file name is entered without a filename extension, the default extension, .PCM, is automatically appended to the filename. The EXTENSION command, page 81, is used to redefine the default file extension. The default file extension applies to the EXECUTE command only.

script library

VMS: The logical name PHYSICA\$LIB can be used to point to the disk and directory for script files. For example, suppose you define:

```
$DEFINE PHYSICA$LIB dsk1:[dir1],dsk2:[dir2]
```

before running the PHYSICA program.

When you execute a script, the current disk and directory is searched for the file. If it cannot be found there, the logical search list is used to find the file. If the logical name is not defined, only the current disk and directory will be searched.

UNIX: The environment variable PHYSICA_LIB can be used to point to the disk and directory for script files. For example, suppose you define:

```
%setenv PHYSICA_LIB disk/directory
```

before running the PHYSICA program.

When you execute a script, the current disk and directory is searched for the file. If it cannot be found there, the environment variable is used to find the file. If the environment variable is not defined, only the current disk and directory will be searched.

Comments

A comment line is any line that begins with an exclamation mark, !. These lines are simply ignored, but can be useful for documentation of files. Comments can also be appended to the end of any line. Just start the comment with an exclamation mark. For example,

READ FILE.DAT X Y Z ! This is a comment

Echoing

If the ENABLE ECHO command, page 73, is entered, the commands that are read from the file will be displayed on the monitor screen. This is useful for following the progress of a command file. If ECHO is disabled globally, but is enabled within a script file, it will be enabled only while that file is executing.

Temporarily passing control to the keyboard

If the TERMINAL command, page 256, is encountered in a command file, control passes back to the terminal keyboard. The user interactively enters commands at this point, until a null line is entered. The command file then recommences execution with the command immediately after the TERMINAL command.

By default, the message 'type ¡RETURN¿ to continue' will be displayed when the TERMINAL command is encountered. You can specify the message by entering a string with the TERMINAL command.

Returning from a script

If the RETURN command, page 224, is encountered in a command file, control passes back to the calling script, if there is one, or to the keyboard, if that script was the top level script.

Aborting a script

If control-c is typed while a script is executing, the entire script stack will be aborted. That is, no matter how deeply the scripts are nested, program flow control is passed back to the keyboard.

If you type the RETURN command from the keyboard after a TERMINAL command has been encountered in a command file, execution of that command file is aborted.

Passing parameters to a script

Parameters that are entered after the file name are used in two ways.

sequential parameters the n^{th} parameter will replace the n^{th} ? that is found in the file numbered parameters the n^{th} parameter in the list will replace all ?n's found in the file

Sequential parameters must be in a one-to-one correspondence with the ?'s, and in the correct order. It is possible to mix sequential and numbered parameters in the same file, but it is not recommended as this can be very confusing.

Prompting

By default, if an incorrect parameter of a valid command is read from the file or is substituted from the parameter list, the user will be prompted to enter the correct information from the terminal keyboard, and the command will then be executed. Use the DISABLE PROMPTING command if a script is to abort when invalid command parameters are encountered.

Labels and GOTOs

A label is a string terminated with a colon, :, and with no embedded blanks. A label must be on a line by itself.

Use a GOTO to branch to a label. Do not include the colon with the label after a GOTO.

Example 1

```
GOTO A_LABEL ! branch to the label (note there is no :)
...
A_LABEL: ! this is a label (note the :)
...
```

Example 2

```
START:

IF (J>8) THEN GOTO END

...

J=J+1

GOTO START

END:
...
```

DO loops

DO loops in PHYSICA are similar to FORTRAN do loops, but *must* be closed off with an ENDDO statement. The basic form of the DO statement is:

```
DO j = x
```

where the looping variable, j, will be made into a scalar variable, and the range of the loop, x, can be any expression resulting in a vector.

Nested loops are allowed. The maximum number of DO loops in a file is fifty (50).

Example 1

```
DO J = x ! x must be a vector, to loop will execute len(x) times
... ! with J taking on the value of each element of x
ENDDO ! end of loop
...
```

Example 2

```
DO I = [2:20:4] ! the loop will execute 5 times, with I taking on the ... ! values [2;6;10;14;18]

ENDDO ! end of loop ...
```

Conditional statements

The general form of a conditional statement is:

```
IF (boolean) THEN
```

The boolean can take any form, but must be either a simple function or it must be enclosed in parentheses, and it must have a scalar result. A result of 1 is true, while anything else is false.

An IF statement can precede a single command or it can precede a block of commands. If an IF statement precedes a block of commands, it *must* be closed off with an ENDIF statement. Nested IF blocks are allowed. The maximum number of IF blocks in a file is fifty (50).

An IF statement can also precede a single command, in which case do not use the ENDIF.

Example 1

```
IF (A>B) THEN DISPLAY 'A > B'
IF (A=B) THEN DISPLAY 'A = B'
IF (A<B) THEN DISPLAY 'A < B'
```

Example 2

```
IF (A>B) THEN
...
ENDIF
...
```

Example 3

```
START2:

IF (J<=8) THEN

...

J=J+1

GOTO START2

ENDIF
...
```

EXTENSION

```
Syntax EXTENSION { 'ext' }

Examples EXTENSION 'PHYSICA'
```

The EXTENSION command is used to redefine the default file extension for executable script files. The original PHYSICA default file extension is PCM. The default file extension applies to the EXECUTE command only. If you give a file name without a file extension, the default extension is automatically appended to the file name.

If the EXTENSION command is entered without a parameter, the current default extension is displayed.

Example

If you have a script file named MACRO_FILE.PCM you can execute this file with the command: <code>@MACRO_FILE</code>

If you have a script file named MACRO_FILE.PHYSICA you can execute this file with the command: <code>@MACRO_FILE.PHYSICA</code>, or with

```
EXTENSION 'PHYSICA' @MACRO_FILE
```

FIGURE

```
FIGURE BOX { lowx lowy hix hiy }
Syntax
          FIGURE POLYGON nvert { cx cy sx sy }
          FIGURE CIRCLE radius { cx cy }
          FIGURE ARC { cx cy sx sy ex ey }
          FIGURE WEDGE { cx cy sx sy ex ey }
          FIGURE ELLIPSE a b { cx cy } angle
          FIGURE ARROW { sx sy ex ey }
Qualifiers \CONFIRM, \GRAPH, \PERCENT, \WORLD
Defaults
          \NOCONFIRM, \PERCENT
Examples FIGURE BOX 10 10 90 90
          FIGURE\NOCONFIRM\GRAPH POLY 6 -1 2 .01 .03
          FIGURE\WORLD CIRC 2
          FIGURE\WORLD WEDGE
          FIGURE\NOCONFIRM\GRAPH ARC -10.2 4.7
          FIGURE\GRAPH ELLIPSE 5.3 1.4 -2.3 3.5 45
```

The FIGURE command is used to draw geometric figures. The figure type is chosen with a keyword. See Table 2.30.

Line types

The line type used for drawing figures will be the current value of LINTYP, which can be changed with the SET LINTYP command The default value for LINTYP is 1. The line type definitions can be changed with the SET LINES command. Line type 1 defaults to a normal line. See the DISPLAY LINES command for information on viewing the line types.

Fillable figures

Most of the figures are fillable: BOX, POLYGON, WEDGE, CIRCLE, ELLIPSE, and ARROWS with

| keyword | figure type |
|---------|--------------------------------------|
| ARC | an arc of a circle |
| WEDGE | a sector of a circle (fillable) |
| CIRCLE | a circle (fillable) |
| BOX | a rectangle (fillable) |
| POLYGON | a regular polygon (fillable) |
| ELLIPSE | an ellipse (fillable) |
| ARROW | an arrow (closed heads are fillable) |

Table 2.30: Geometric figures that can be drawn with the FIGURE command

closed heads. Use the SET FILL command to change the fill type and pattern. See Table 2.55 on page 233 for a description of the interpretations of the FILL keyword.

X Windows

When running under X Windows, mouse button two toggles the continuous display of the graphics cursor location. The PHYSICA keyword CUNITS is the units type for these numbers. If CUNITS = WORLD, the numbers depend on the current units type, either CM or IN, as chosen with SET UNITS. If CUNITS = GRAPH, the numbers displayed depend on the current graph axis scales. If CUNITS = PERCENT, the numbers depend on the current window.

Units

The numeric parameters may be expressed in three types of units, which are chosen by command qualifier. The default is \PERCENT . See Table 2.31 for a listing of the qualifiers and their interpretations.

| qualifier | interpretation of the coordinates | | | | | |
|-----------|--|--|--|--|--|--|
| \PERCENT | percentages of the current window, as chosen with the WINDOW command. | | | | | |
| | Lengths are in terms of the horizontal dimension. | | | | | |
| ∖GRAPH | graph units, that is, the units defined by the minimum and maximum values for the last graph drawn. If no graph has been drawn yet, the defaults are $-1 \le x \le 1$ and $-1 \le y \le 1$ | | | | | |
| \WORLD | centimeters or inches, as chosen with the SET UNITS command | | | | | |

Table 2.31: Types of units recognized by the FIGURE command

For example, if the \PERCENT qualifier is used, then a location of (50,50) represents the centre of the current window. If the \WORLD qualifier is used, the coordinates are in units of the

world coordinate system, the plotting units. The default world coordinate system units are centimeters. See the DEVICE command for tables showing the dependence of plotting units on the graphics hardcopy output device.

Confirmation

If the FIGURE command is entered interactively, the default is that confirmation that the figure is acceptable as drawn will be requested. The figure will not be be entered into a hardcopy plot file, or into an open EDGR file, unless it is accepted. However, if CONFIRM has been disabled, with the DISABLE CONFIRM command, the default will be that no such confirmation will be requested. If the \CONFIRM qualifier is used, confirmation will be forced. If the \NOCONFIRM qualifier is used, confirmation will be suppressed.

Stack file

If a stack file is open, via the STACK command, then the (x,y) coordinates will be written to the stack file, even if they are chosen by the graphics cursor. Thus, when this stack file is executed, using the EXECUTE command, the graphics cursor will not be used. If confirmation is requested and the figure is not \mathtt{OK} , then the command is *not* written to the stack file.

Circle

```
Syntax FIGURE CIRCLE r { cx cy }
```

The FIGURE CIRCLE command draws a circle, centred at the point (cx,cy), with radius r. The parameter r is not optional, and is in terms of the horizontal dimension. The graphics cursor is used if either cx, or cy is not entered.

Arc

```
Syntax FIGURE ARC { cx cy sx sy ex ey }
```

The FIGURE ARC command draws an arc of a circle, centred at the point (cx,cy), starting at the point (sx,sy) and finishing on the line through the points (ex,ey) and (cx,cy). The graphics cursor is used if any of cx, cy, sx, sy, ex, or ey are not entered.

The first two points determine the radius and the starting azimuth, while the final point determines the final azimuth only. If the final point is the same as the first point, the centre, then a complete circle will be drawn.

Wedge

```
Syntax FIGURE WEDGE { cx cy sx sy ex ey }
```

The FIGURE WEDGE command draws a sector of a circle, centred at the point (cx,cy), starting at the point (sx,sy) and finishing on the line through the points (ex,ey) and (cx,cy). The endpoints of the arc are joined to the centre of arc. The graphics cursor is used if any of cx, cy, sx, sy, ex, or ey are not entered.

The first two points determine the radius and the starting azimuth, while the final point determines the final azimuth only. If the final point is the same as the first point, the centre, then a complete circle will be drawn.

Box

```
Syntax FIGURE BOX { lowx lowy hix hiy }
```

The FIGURE WEDGE command draws a box, or rectangle, with lower left hand corner at (lowx,lowy), and upper right hand corner at (hix,hiy). The graphics cursor is used if any of lowx, lowy, hix, or hiy are not entered.

Polygon

```
Syntax FIGURE POLYGON n { cx cy sx xy }
```

The FIGURE POLYGON command draws a regular polygon, with nvert vertices, centred at (cx,cy), and with the first vertex at (sx,sy). The graphics cursor is used if any of cx, cy, sx, or sy are not entered.

Ellipse

```
Syntax FIGURE ELLIPSE a b { cx cy } ang
```

The FIGURE ELLIPSE command draws an ellipse, with a being the major axis radius, b the minor axis radius, centred at point (cx,cy), and with the major axis at an angle of ang degrees, measured counter-clockwise from the horizontal. The graphics cursor is used if either cx or cy are not entered.

Arrow

```
Syntax FIGURE ARROW { sx sy ex ey }
```

The FIGURE ARROW command draws an arrow with base at (sx,sy) and end point at (ex,ey). The graphics cursor is used if any of sx, sy, ex, or ey are not entered.

Styles

The arrow style is chosen with the SET ARROTYP command. The default is value of ARROTYP is 0. See Table 2.54 on page 231 and Figure 2.24 on page 230.

Head width

The width of the arrowhead is chosen with the SET ARROWID command. ARROWID is the arrow head width as a fraction of the total arrow length. The default value of ARROWID is 0.15.

Length

The length of the arrowhead is chosen with the SET ARROLEN command. ARROLEN is the arrow head length as a fraction of the total arrow length. The default value of ARROLEN is 0.20.

FILTER

```
Syntax FILTER\MEDIAN x f npt
FILTER\MEAN x f npt
FILTER\NONRECURSIVE x f c
FILTER\RECURSIVE x f c d

Qualifiers \MEDIAN, \MEAN, \NONRECURSIVE, \RECURSIVE

Default \MEDIAN

Examples FILTER\MEDIAN X XF 5
FILTER\NONRECURSIVE X XF [1;-2;1]
FILTER\MEAN X XF -5
FILTER\RECURSIVE X XF [.3584;1.2832;.3584;0;0] [0;1]
```

A digital filter is a linear combination of the input data, x, and possibly the output data, f. The input data is assumed to be *equally spaced* samples of some continuously varying quantity; and any error or noise is in the measurements. In the PHYSICA implementation of filters, the input data is assumed to have unit spacing, so a scale factor may have to be applied to produce the correctly scaled output data.

The simplest kinds of filters are the nonrecursive filters defined by the convolution formula:

$$f_n = \sum_{k=-N}^{N} c_k x_{n-k}$$

The coefficients c_k are the constants of the filter, the x_{n-k} are the input data, and the f_n are the outputs. When values of the output as well as the data values are used to compute the output values, the filter is called a recursive filter. It is usual to limit the range of

nonzero coefficients to current and past values of the data x_n and to only past values of the output f_n . This type of filter is called causal recursive and can be defined by the convolution formula:

$$f_n = \sum_{k=0}^{N} c_k x_{n-k} + \sum_{k=1}^{M} d_k f_{n-k}$$

Nonrecursive or recursive filters using constant coefficients q_k and d_k are called time-invariant filters. Users are urged to refer to textbooks dealing with digital filters, such as <u>Digital Filters</u> by R.W. Hamming, Prentice-Hall 1977, or <u>Digital Signal Analysis</u> by Samual D. Stearns, Hayden Book Co. Inc.

Noise amplification caused by filtering

It can be shown (Hamming, p.17) that the sum of the squares of the filter coefficients measures the noise amplification of the filtering process. Thus, the variance, σ^2 , will be amplified by $\sum \sigma c_i^2$.

Median filter

The default is \MEDIAN, that is, to use a running median filter. The data array, x, is filtered through a window npt points in width. npt must be ≥ 2 .

The median filter is particularly good at removing 'spikes' from data. The median filter moves a window over the data and outputs the median value of the data points within each window placement. The window butts up against the ends. When npt is even, the filter is applied twice, first skewed left then skewed right, and the results are averaged.

Mean filter

```
Syntax FILTER\MEAN xin xout npt
```

If the \mean qualifier is used, the filter will be the running mean, or average, filter. This filter method is sensitive to large spikes in the data. Any large spikes, for example, > 1000 times normal value, should first be removed, by, for example, the median filter.

There are two versions of the running mean filter, which are chosen by whether npt is positive or negative. The window width is always |npt|.

If npt > 0, the average value of each window placement is calculated by summing the rele-

vant points and dividing by npt. The averaging window butts up against the end.

If npt < 0, a much faster method is used which adds a new point to the right and drops an old one from the left. The window runs off half way from each end, but pseudo points outside the range are set to the appropriate end point values.

Nonrecursive filters

```
Syntax FILTER\NONRECURSIVE x f c
```

If the \NONRECURSIVE qualifier is used, the third parameter, c, must be a vector. The data array, x, is processed through a nonrecursive filter using the values of c as the data coefficients:

$$f[n] = \sum_{k=1}^{N} c[k] x[n+k-(\frac{N}{2}+1)]$$

where N is the length of vector c. Note that when c has an even number of elements, the filter will be applied to the n^{th} point by application to points from n-N/2 to n-1+N/2. For example, when N is two, the weightings will be applied to the previous point and to the current point.

Differentiating nonrecursive filters

Remember, that the x's must be equally spaced, and are actually assumed by the FILTER command to have unit spacing. Thus, to obtain the correct output scaling, multiply f by $k!/(N-1)!h^k$, where k is the order of the derivative, N is the length of vector c, and h is the spacing of x, that is, h = x[i+1]-x[i]. For example:

```
FILTER\NONRECURSIVE X XOUT [2;-16;0;16;-2] ! 1st deriv. nonrecursive filter XOUT=XOUT/(24*(X[2]-X[1])) ! use scale factor 1/(h*4!)
```

See Table 2.32 for various first derivative nonrecursive filter data coefficients. See Table 2.33 for various second derivative nonrecursive filter data coefficients. See Table 2.34 for various third derivative nonrecursive filter data coefficients.

| type | data coefficients | scale factor |
|---------|------------------------------|--------------|
| 3 point | [1;0;-1] | 1/2h |
| 4 point | [1; -6; 3; 2] | 1/6h |
| 5 point | [2; -16; 0; 16; -2] | 1/24h |
| 6 point | [-4; 30; -120; 40; 60; -6] | 1/120h |

Table 2.32: Various 1^{st} derivative nonrecursive filters

| type | data coefficients | scale factor |
|---------|---|--------------|
| 3 point | [1; -2; 1] | $1/h^{2}$ |
| 4 point | [0; 3; -6; 3] | $1/3h^{2}$ |
| 5 point | [-1; 16; -30; 16; -1] | $1/12h^2$ |
| 6 point | [-1; 16; -30; 16; -1] [0; 5; 80; -150; 80; -5] | $1/60h^2$ |

Table 2.33: Various 2nd derivative nonrecursive filters

| type | data coefficients | scale factor |
|---------|---------------------------|--------------|
| 4 point | [-1; 3; -3; 1] | $1/h^{3}$ |
| 5 point | [-2; 4; 0; -4; 2] | $1/4h^{3}$ |
| 6 point | [5; -35; 70; -50; 5; 5] | $1/20h^{3}$ |

Table 2.34: Various 3^{rd} derivative nonrecursive filters

Smoothing nonrecursive filters

See Table 2.35 for various quadratic smoothing nonrecursive filter data coefficients. See Table 2.36 for various quartic smoothing nonrecursive filter data coefficients. See Table 2.37 for Spencer's formulae smoothing nonrecursive filter data coefficients.

| type | data coefficients | scale factor |
|----------|--|--------------|
| 5 point | [-3; 12; 17; 12; -3] | 1/35 |
| 7 point | [-2; 3; 6; 7; 6; 3; -2] | 1/21 |
| 9 point | [-21; 14; 39; 54; 59; 54; 39; 14; -21] | 1/231 |
| 11 point | [-36; 9; 44; 69; 84; 89; 84; 69; 44; 9; -36] | 1/429 |

Table 2.35: Smoothing nonrecursive filters (quadratic)

Interpolating nonrecursive filters

Suppose we have points in a vector which are "bad" and need to be replaced. Assuming one can fit the data with an odd degree polynomial. The next higher order difference equation, when set to zero, can be used to give the desired filter coefficients. For example, if the data can be fit with a 5^{th} order polynomial, the fourth difference set to zero gives:

$$y_{n-2} - 4y_{n-1} + 6y_n - 4y_{n+1} + y_{n+2} = 0$$

and solving for y_n gives:

$$y_n = \frac{1}{6}(-y_{n-2} + 4y_{n-1} + 4y_{n+1} - y_{n+1})$$

so the data filter coefficients are [$-\frac{1}{6}$; $\frac{2}{3}$; 0; $\frac{2}{3}$; $-\frac{1}{6}$].

| type | data coefficients | scale factor |
|----------|---|--------------|
| 7 point | [5; -30; 75; 131; 75; -30; 5] | 1/231 |
| 9 point | [15; -55; 30; 135; 179; 135; 30; -55; 15] | 1/429 |
| 11 point | [18; -45; -10; 60; 120; 143; 120; 60; -10; -45; 18] | 1/429 |
| 13 point | [110; -198; -135; 110; 390; 600; 677; 600; 390; 110; -135; -198; 110] | 1/2431 |

Table 2.36: Smoothing nonrecursive filters (quartic)

| type | data coefficients | scale factor |
|----------|--|--------------|
| 15 point | [-3; -6; -5; 3; 21; 46; 67; 74; 67; 46; 21; 3; -5; -6; -3] | 1/320 |
| 21 point | [-1; -3; -5; -5; -2; 6; 18; 33; 47; 57; 60; 57; 47; 33; 18; 6; -2; -5; -5; -3; -1] | 1/350 |

Table 2.37: Smoothing nonrecursive filters (Spencer's formulae)

Recursive filters

Syntax FILTER\RECURSIVE x f c d

If the \RECURSIVE qualifier is used, the third parameter, c, must be a vector, and the fourth parameter, d, must also be a vector. The data array, x, is processed through a recursive filter. This allows for the specification of a completely general recursive filter of arbitrary length. The values of c are the filter coefficients which operate on the data. The values of d are the filter coefficients which operate on the previously made output.

$$\mathbf{f}[n] = \sum_{k=1}^{N} \mathbf{c}[k] \; \mathbf{x}[n+k-(\frac{N}{2}+1)] + \sum_{k=1}^{M} \mathbf{d}[k] \; \mathbf{f}[n-k]$$

where N is the length of vector c and M is the length of vector d.

Integrating recursive filters

The trapezoidal rule integration filter:

$$G_{n+1} = 0.5(x_{n+1} + x_n) + G_n$$

The Leo Tick formula for integration:

$$G_{n+1} = h(0.3584x_{n+1} + 1.2832x_n + 0.3584x_{n-1}) + G_{n-1}$$

See Table 2.38 for the trapezoidal rule and the Leo Tick formula integrating recursive filter coefficients.

| | data coefficients | output coefficients |
|------------------|----------------------------|---------------------|
| Trapezoidal rule | | [1] |
| Leo Tick formula | [0.3584; 1.2832; 0.3584] | [0; 1] |

Table 2.38: Integrating recursive filters

Examples

The following script demonstrates how you can use the FILTER command to smooth data. See Figure 2.10.

```
X = [0:4:.05]
Y=X^2-3*X+3+SIN(X*3)*RAN(X)
WINDOW 15
SET PCHAR -1
LABEL\XAXIS 'original data'
GRAPH X Y
WINDOW 16
LABEL\XAXIS '11 point quadratic formula'
FILTER\NONRECURSIVE Y YF [-36;9;44;69;84;89;84;69;44;9;-36]
SET PCHAR O
GRAPH X YF/429
WINDOW 17
LABEL\XAXIS '11 point quartic formula'
FILTER\NONRECURSIVE Y YF [18;-45;-10;60;120;143;120;60;-10;-45;18]
GRAPH X YF/429
WINDOW 18
LABEL\XAXIS '15 point Spencer'//CHAR(39)//'s formula'
FILTER\NONRECURSIVE Y YF [-3;-6;-5;3;21;46;67;74;67;46;21;3;-5;-6;-3]
GRAPH X YF/320
```

The following script demonstrates how you can use the FILTER command to differentiate data. See Figure 2.11.

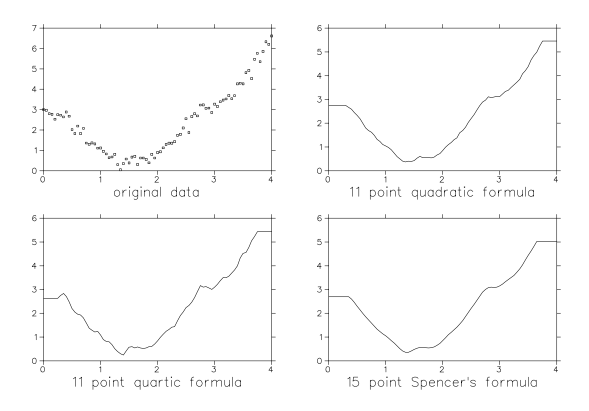


Figure 2.10: A FILTER example showing data smoothing

```
X = [0:4:.2]
H=X[2]-X[1]
Y=X^2-3*X+3
WINDOW 5
SET PCHAR -1
LABEL\XAXIS 'original data'
SET %XLABSZ 5
GRAPH X Y
WINDOW 6
SET PCHAR O
LABEL\XAXIS '1<^>st<_> derivatives'
FILTER\NONRECURSIVE Y YF [-4;30;-120;40;60;-6]
SCALE 0 4 0 -3 5 0
SET PCHAR -2
GRAPH X YF/(120*H)
SET PCHAR O
GRAPH\NOAXES X 2*X-3
```

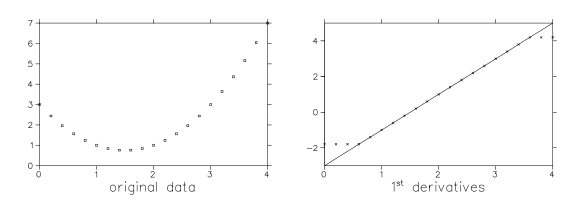


Figure 2.11: A FILTER example showing 1^{st} derivative

FIT

Syntax FIT y = expression
FIT\UPDATE yout

Qualifiers \ \NORMAL, \POISSON, \UPDATE, \ITMAX, \WEIGHTS, \ZEROS, \TOLERANCE, \CHISQ, \CL, \CORRMAT, \COVMAT, \E1, \E2, \VARNAMES, \FREE, \RESET, \MESSAGES

Defaults \ \NORMAL, \-UPDATE, \-ITMAX, \-WEIGHTS, \ZEROS, \-TOLERANCE, \-CHISQ, \-CL, \-CORRMAT, \-COVMAT, \-E1, \-E2, \-VARNAMES, \-FREE, \-RESET, \MESSAGES

Examples FIT Y=A*X+B
FIT\WEIGHTS\CHISQ\CL\ITMAX W 3 Y=A*EXP(-B*X)+C
FIT\CORR\NOMESS Y=A*X+B
FIT\POISSON Y=A*EXP(-B*X)+C
FIT\UPDATE YF

By default, or if the \NORMAL qualifier is used, it is assumed that each data point has an error that is distributed as a normal distribution,

$$\begin{split} N(x;\mu,\sigma) &\equiv \frac{1}{\sigma\sqrt{2\pi}} e^{-\frac{1}{2}(\frac{x-\mu}{\sigma})^2} \quad \text{for} \quad -\infty < x < \infty \\ &= \sqrt{w/(2\pi)} e^{-\frac{w}{2}(x-\mu)^2} \end{split}$$

where μ is the mean and σ is the standard deviation of the distribution. The weight w is defined as: $w = \frac{1}{\sigma^2}$.

If the \POISSON qualifier is used, the data errors are assumed to be distributed as a Poisson distribution,

$$P(x;\lambda) \equiv \frac{\lambda^x e^{-\lambda}}{x!}$$
 for $x = 0, 1, 2, ...$

where λ is the mean and the variance of the distribution.

Expression and parameters

The expression must result in a vector with the same length as the data vector, y. A maximum of twenty-five (25) fitting parameters are allowed in the expression. The fitting parameter values are altered during the fit. Fit parameters are created with the SCALAR\VARY command, and can be converted to fixed value scalars with the SCALAR command. If you use the \RESET qualifier, the fitting parameters will be reset to their original values after an unsuccessful fit, or a control-c abort.

If the \VARNAMES qualifier is used with the FIT command, a string array variable named

FIT\$VAR will be made which will contain the names of the fitting parameter variables. The array length of FIT\$VAR will be equal to the number of fit parameters.

Method

Suppose that you have N data points, y_k , for $k=1,\ldots,N$, and the function to be fitted is f(x,p), where p represents the M parameters $< p_1,p_2,\ldots,p_M>$. Define the likelihood of the parameters, given the data, as the probability of the data, given the parameters. We fit for the parameters, p, by finding those values, p_{\min} that maximize this likelihood. This form of parameter estimation is known as maximum likelihood estimation.

Some good references are:

- Practical Methods of Optimization, by R. Fletcher, 1980;
- Methods for Unconstrained Optimization Problems by J. Kowalik and M.R. Osborne, 1968;
- Statistical Methods in Experimental Physics, by W.T. Eadie, et.al., 1971;
- Mathematical Statistics, by John E. Freund, 1971;
- Formulae and Methods in Experimental Data Evaluation, Volume 3,
 Elements of Probability and Statistics, by Siegmund Brandt, 1984;
- Numerical Recipes The Art of Scientific Computing, by W.H. Press, et.al. 1986.

Consider the likelihood function $\mathcal{L}(p) \equiv \prod_{k=1}^N P(x_k,p)$ where P is the probability density, which depends on the random variable x and the parameters p. \mathcal{L} is a measure for the probability to observe just the particular sample we have, and is called an a-posteriori probability since it is computed after the sampling is done. The best estimates for p are the values which maximize \mathcal{L} . But maximizing the logarithm of \mathcal{L} also maximizes \mathcal{L} , and maximizing $\ln(\mathcal{L})$ is equivalent to minimizing $-\ln(\mathcal{L})$. So, the goal becomes minimizing the log likelihood function:

$$-L(\boldsymbol{p}) \equiv -\ln \mathcal{L}(\boldsymbol{p}) = -\sum_{k=1}^{N} \ln P(x_k, \boldsymbol{p})$$

Let p^0 be the initial values given for p. The goal is to find a ∇p so that $p^1 = p^0 + \nabla p$ is a better approximation to the data. We use the iterative Gauss-Newton method, and the series p^1, p^2, p^3, \ldots will hopefully converge to the minimum, p_{\min} .

Generally, the Gauss-Newton method is locally convergent when χ^2 is zero at the minimum. Serious difficulties arise when f is sufficiently nonlinear and χ^2 is large at the minimum.

The Gauss-Newton method has the advantage that linear least squares problems are solved in one iteration.

Consider the Taylor expansion of L(p):

$$L(\boldsymbol{p}) = L(\boldsymbol{p}^0) + \sum_{j=1}^{M} \frac{\partial L}{\partial p_j} \bigg|_{\boldsymbol{p}^0} p_j + \frac{1}{2} \sum_{i=1}^{M} \sum_{j=1}^{M} \frac{\partial^2 L}{\partial p_i \partial p_j} \bigg|_{\boldsymbol{p}^0} p_i p_j + \cdots$$

Define the arrays *b*, *B* and *c*:

$$[m{b}]_i \equiv -rac{\partial L}{\partial p_i}igg|_{m{p}^0} \quad ext{for} \quad i=1,2,\ldots,M$$
 $[m{B}]_{ij} \equiv rac{\partial^2 L}{\partial p_i \partial p_j}igg|_{m{p}^0} \quad ext{for} \quad i,j=1,2,\ldots,M$ $m{c} \equiv L(m{p}^0)$

If we linearize, that is, assume that $\frac{\partial^2 \ln P}{\partial p_i \partial p_j} \to 0$, then $L(p) \approx c - b \cdot p + \frac{1}{2} p \cdot B \cdot p$, and so $\nabla L = B \cdot p - b$. The problem has reduced to solving the matrix equation $B \cdot \nabla p = b$.

Note: The partial derivatives are approximated numerically using a central difference approximation:

$$\frac{\partial f(x_k, \mathbf{p})}{\partial p_i} = \frac{f(x_k, \mathbf{p} + \nabla \mathbf{p}_i) - f(x_k, \vec{\mathbf{p}} - \nabla \mathbf{p}_i)}{2\nabla \mathbf{p}_i}$$

Tolerance

Syntax FIT\TOLERANCE eps y=expression
FIT\WEIGHTS\TOLERANCE w eps y=expression
FIT\ITMAX\TOLERANCE n eps y=expression
FIT\WEIGHTS\ITMAX\TOLERANCE w n eps y=expression

The \text{TOLERANCE} qualifier allows the user to specify the fitting tolerance, which has a default value of 0.00001. This value is used in calculating the central difference formula for the partial derivatives, that is, $\frac{\partial f(x,p)}{\partial p}$ is approximated by

$$\frac{f(x, p(1 + eps)) - f(x, p(1 - eps))}{(2p \times eps)}$$

This value is also used to determine when the fit is successful.

Normal distribution

Assume that each data point, y_k , has an error that is independently random and distributed as a normal distribution, that is,

$$P(x_k, \mathbf{p}) = \frac{1}{\sigma_k \sqrt{2\pi}} e^{-\frac{1}{2} \left[\frac{y_k - f(x_k, \mathbf{p})}{\sigma_k} \right]^2}$$

where σ^2 is the variance, and $f(x_k, p)$ is the expression that we want to fit.

$$L(\boldsymbol{p}) = \sum_{k=1}^{N} \ln P(x_k, \boldsymbol{p}) = -\frac{1}{2} \sum_{k=1}^{N} \left[\frac{y_k - f(x_k, \boldsymbol{p})}{\sigma_k} \right]^2 + \mathbf{constant}$$

The goal is to minimize the χ^2 function:

$$\chi^{2}(\boldsymbol{p}) \equiv \sum_{k=1}^{N} \left[\frac{y_{k} - f(x_{k}, \boldsymbol{p})}{\sigma_{k}} \right]^{2} = \sum_{k=1}^{N} w_{k} \left[y_{k} - f(x_{k}, \boldsymbol{p}) \right]^{2}$$

where the weights, w_k , are defined as: $w_k \equiv 1/\sigma_k^2$. Consider the Taylor expansion of χ^2 :

$$\chi^{2}(\boldsymbol{p}) = \chi^{2}(\boldsymbol{p}^{0}) + \sum_{j=1}^{M} \frac{\partial \chi^{2}}{\partial p_{j}} \Big|_{\boldsymbol{p}^{0}} p_{j} + \frac{1}{2} \sum_{i=1}^{M} \sum_{j=1}^{M} \frac{\partial^{2} \chi^{2}}{\partial p_{i} \partial p_{j}} \Big|_{\boldsymbol{p}^{0}} p_{i} p_{j} + \cdots$$

Define the arrays b, B and c:

$$[\boldsymbol{b}]_{i} \equiv -\frac{\partial \chi^{2}}{\partial p_{i}}\Big|_{\boldsymbol{p}^{0}} = 2\sum_{k=1}^{N} w_{k} \left[y_{k} - f(x_{k}, \boldsymbol{p})\right] \frac{\partial f(x_{k}, \boldsymbol{p})}{\partial p_{i}}\Big|_{\boldsymbol{p}^{0}} \quad \text{for} \quad i = 1, 2, \dots, M$$

$$[\boldsymbol{B}]_{ij} \equiv \frac{\partial^{2} \chi^{2}}{\partial p_{i} \partial p_{j}}\Big|_{\boldsymbol{p}^{0}} = 2\sum_{k=1}^{N} w_{k} \frac{\partial f(x_{k}, \boldsymbol{p})}{\partial p_{i}}\Big|_{\boldsymbol{p}^{0}} \frac{\partial f(x_{k}, \boldsymbol{p})}{\partial p_{j}}\Big|_{\boldsymbol{p}^{0}} \quad \text{for} \quad i, j = 1, 2, \dots, M$$

$$\boldsymbol{c} \equiv \chi^{2}(\boldsymbol{p}^{0})$$

Linearize and the problem reduces to solving the matrix equation $B \cdot \nabla p = b$.

 χ^2 and weights

The weight at each point defaults to one (1), if a weight vector is not entered. Weights only make sense with a normal distribution, and are ignored when used with the \POISSON qualifier.

To make use of a weight array, the \WEIGHTS qualifier *must* be entered. If the \WEIGHTS qualifier is used, the weight vector, w, will then be expected. The weights are assigned to

the dependent variable in a one-to-one fashion, that is, the weight vector must be the same length as the data vector, y. If the \ITMAX qualifier is used, the weight comes before the iteration maximum in the command parameter list. If the \TOLERANCE qualifier is used, the iteration maximum comes before the tolerance in the command parameter list.

By default, the zero elements of the weight vector are used when calculating the number of degrees of freedom. If the \-ZEROS qualifier is used with the \WEIGHTS qualifier, then the zero elements of the weight vector will not be used when calculating the number of degrees of freedom. This could have an affect on the calculation of the confidence level, the χ^2 per degrees of freedom, and E2, the root mean square total errors of estimate.

If the \CHISQ qualifier is used, a new scalar, named FIT\$CHISQ, will be made with value equal to the total $\chi^2 = \sum w_k \left[y_k - f(x_k, p_{\min}) \right]^2$ where w_k represents the optional weight at each data point y_k , f is the expression to be fitted, and p_{\min} are the best values of the p parameters.

Hint for physicists

Very often, the data to be fitted is a histogram of physical events. In that case, since each bin would follow a multinomial distribution, the error is equal to \sqrt{f} , where f is the expression you are trying to fit. Of course, since you don't know the parameter values yet, you don't actually know f, so you approximate by using the y data values. In the limit, these results are the same. In the case of a large number of bins, the variance can be approximated by \sqrt{y} . Hence, the correct weighting factor that will give properly normalized errors is w=1/y, and the corresponding one standard deviation error, $\sigma=E2/\sqrt{\chi^2/n}$, where E2 is the standard error and n is the number of degrees of freedom, usually equal to the number of data points minus the number of parameters, (N-M).

Degrees of freedom

If the \FREE qualifier is used, then the number of degrees of freedom for the fit is output into an automatically created scalar named FIT\$FREE. The number of degrees of freedom is either the number of data values minus the number of parameters, or, if the \FREE qualifier is also used, the number of non-zero weights minus the number of parameters.

Poisson distribution

Assume that each data point has an error that is independently random and distributed as a Poisson distribution. The log likelihood function, L(p), as a function of the fit parameters, p, is minimized using a Gauss-Newton method. Since logarithms are involved, a good first approximation is required before starting the Poisson fit, so try a normal fit first, and use the resultant parameter values to start off the Poisson fit.

Weights do not have meaning, and so are not used, in a Poisson fit.

Assume that each data point, y_k , has an error that is independently random and distributed as a Poisson distribution, that is, $P(x_k, \mathbf{p}) = f(x_k, \mathbf{p})^{y_k} e^{-f(x_k, \mathbf{p})}/y_k!$. We want to minimize:

$$-L = \sum_{k=1}^{N} \ln \left[\frac{f(x_k, \boldsymbol{p})^{y_k} e^{-f(x_k, \boldsymbol{p})}}{y_k!} \right] = \sum_{k=1}^{N} \left[y_k \ln \left(f(x_k, \boldsymbol{p}) \right) - f(x_k, \boldsymbol{p}) - \ln \left(y_k! \right) \right]$$

but $\sum \ln (y_k!)$ is a constant. So, the goal is to minimize

$$\mathcal{P}(\boldsymbol{p}) \equiv \sum_{k=1}^{N} [y_k \ln (f(x_k, \boldsymbol{p})) - f(x_k, \boldsymbol{p})]$$

Consider the Taylor expansion of \mathcal{P} :

$$\mathcal{P}(\boldsymbol{p}) = \mathcal{P}(\boldsymbol{p}^0) + \sum_{j=1}^{M} \frac{\partial \mathcal{P}}{\partial p_j} \bigg|_{\boldsymbol{p}^0} p_j + \frac{1}{2} \sum_{i=1}^{M} \sum_{j=1}^{M} \frac{\partial^2 \mathcal{P}}{\partial p_i \partial p_j} \bigg|_{\boldsymbol{p}^0} p_i p_j + \cdots$$

Define:

$$\begin{aligned} [\boldsymbol{b}]_i &\equiv -\frac{\partial \mathcal{P}}{\partial p_i} \bigg|_{\boldsymbol{p}^0} = -\sum_{k=1}^N \left[\frac{y_k}{f(x_k, \boldsymbol{p})} \frac{\partial f(x_k, \boldsymbol{p})}{\partial p_i} - \frac{\partial f(x_k, \boldsymbol{p})}{\partial p_i} \right] \quad \text{for} \quad i = 1, 2, \dots, M \\ [\boldsymbol{B}]_{ij} &\equiv \frac{\partial^2 \mathcal{P}}{\partial p_i \partial p_j} \bigg|_{\boldsymbol{p}^0} = -\sum_{k=1}^N \frac{y_k}{f^2(x_k, \boldsymbol{p})} \frac{\partial f(x_k, \boldsymbol{p})}{\partial p_i} \frac{\partial f(x_k, \boldsymbol{p})}{\partial p_j} \quad \text{for} \quad i, j = 1, 2, \dots, M \\ \boldsymbol{c} &\equiv \mathcal{P}(\boldsymbol{p}^0) \end{aligned}$$

Then: $\mathcal{P}(p) \approx c - b \cdot p + \frac{1}{2} p \cdot B \cdot p$. Linearize, and the problem reduces to solving the matrix equation $B \cdot \nabla p = b$.

 χ^2 of the fit

If the \CHISQ qualifier is used, a new scalar, named FIT\$CHISQ, will be made with value equal to the total $\chi^2=2\sum\left[f(x_k,p_{\min})-y_k+y_k\ln\frac{y_k}{f(x_k,p_{\min})}\right]$ where f is the expression to be fitted, and p_{\min} are the best values of the parameters p. This assumes that y_k is the outcome of a Poisson process.

Correlation and covariance

An indication of the accuracy of the fit is displayed in the output under the names E1 and E2.

$$[\boldsymbol{E1}]_i \equiv \sqrt{\left[\boldsymbol{B}^{-1}\right]_{ii}} \quad \text{for } i = 1, \dots, M$$

$$[\boldsymbol{E2}]_i \equiv \left[\boldsymbol{E1}\right]_i \sqrt{\frac{1}{n} \sum_{k=1}^N w_k [y_k - f(x_k, \boldsymbol{p})]^2} \quad \text{for } i = 1, \dots, M$$

where n is the number of degrees of freedom, and where $[B^{-1}]_{ii}$ are the diagonal elements of the inverse of the matrix B. B^{-1} is called the covariance matrix. The $[E1]_i$ are called the root mean square statistical errors of estimate, while the $[E2]_i$ are called the root mean square total errors of estimate, or standard errors.

The accuracy of the parameters in a linear fit is $p_i \pm [\mathbf{E2}]_i$ for i = 1, ..., M. In the linear case, for the standard error $\mathbf{E2}$ to be correct, the weights w_k must be proportional to $1/\sigma_k^2$, where σ_k is the standard deviation of the probability distribution of y_k . In the nonlinear case, $\mathbf{E2}$ does not have the same statistical significance.

If the \COVMAT qualifier is used, a matrix called FIT\$COVM will be created which will contain B^{-1} . If the \CORRMAT qualifier is used, a matrix with the name FIT\$CORR will be created which will contain the correlation matrix for the fit. The size of these matrices will be M by M. If the \E1 qualifier is used, then the root mean square statistical error for each fit parameter are output into an automatically created vector named FIT\$E1. If the \E2 qualifier is used, then the root mean square total error of estimate for each parameter are output into an automatically created vector named FIT\$E2. The values are stored in these vectors in the order corresponding to the order in which the parameters appeared in the expression. The length of these vectors will be equal to the number of parameters in the fit expression.

Confidence level of the fit

If the \CL qualifier is used, a new scalar, named FIT\$CL, will be made with value equal to the confidence level:

$$CL(\chi^2) = \frac{1}{2^{n/2}\Gamma(n/2)} \int_{\chi^2}^{\infty} t^{n/2-1} e^{-t/2} dt$$

where n is the degrees of freedom, usually equal to the number of data points minus the number of parameters, (N-M). The confidence level is the probability that a random repeat of the given experiment would observe a worse χ^2 , assuming the correctness of the model.

Number of iterations

The \ITMAX qualifier allows the user to specify the maximum number of iteration steps for the fit. When this maximum number is reached, the fit will stop, and the variable parameters will be updated to their last values. The fit will also stop if the fit is successful before this maximum iteration number is reached. If the \WEIGHT qualifier is also used, the weight array comes before the iteration number in the command parameter list.

Informational messages

By default, information on the progress of the fit, as well as the results, are displayed on the monitor screen. If the \NOMESSAGES qualifier is used, these informational messages will be suppressed.

Update after a fit

```
Syntax FIT\UPDATE yout
```

The FIT\UPDATE command evaluates the previously fitted expression, that is, the expression in the last FIT command, for the current parameter values, and stores this result in the vector yout.

This is exactly equivalent to entering: yout=previously_fitted_expression and is provided only to obviate the necessity of re-entering a complicated expression.

Note: The vector name yout usually differs from the name of the vector being fitted to avoid destroying the original data.

FMIN

| Syntax | FMIN x y xlo xhi expression |
|-----------|--|
| Qualifier | \MESSAGES |
| Defaults | \NOMESSAGES |
| Example | FMIN\-MESSAGES X Y -10 10 $2*X^2-10*X+5$ |

The FMIN command returns the location and value of the local minimum in the range xlo to xhi of the specified expression. This expression must be a function of the independent variable x, which must be a scalar.

The value of x is interpreted as the user's initial guess for the location of the local minimum. If the value of x is outside the range xlo to xhi, then the midpoint, (xlo+xhi)/2 is chosen as this initial guess.

On output, the scalar x will contain the location of the local minimum, and scalar y will contain the value of this minimum.

The expression must contain the variable x, and may contain other scalars, but must *not* contain any other non-scalar variables.

Informational messages

By default, the value of the local minimum, its location, and the upper limit on the error are displayed on the monitor screen. If the \NOMESSAGES qualifier is used, this informational message will be suppressed.

Example

The following script demonstrates how you can use the FMIN command to find a local minimum. See Figure 2.12.

```
XMIN=-10
XMAX=10
XD=[XMIN:XMAX:.1]
GRAPH XD 2*XD^2-10*XD+5
GET
    YMIN YMIN
    YMAX YMAX

X=0
FMIN X Y -10 10 2*X^2-10*X+5
GRAPH\NOAXES [XMIN;XMAX] [Y;Y]
GRAPH\NOAXES [X;X] [YMIN;YMAX]
TEXT 'y=2x<^>2<_>-10x+5'
TEXT 'minimum value= '//rchar(y)//' at x='//rchar(x)
```

FZERO

```
Syntax FZERO x expression

Qualifier \messages

Defaults \nomessages \text{NOMESSAGES}

Example FZERO\-messages x SIN(X)/X
```

The FZERO command returns the zeros, or roots, of expression. This expression must be a function of the independent variable x, which must be a vector.

If the length of vector \mathbf{x} is N, then a maximum of N roots of the expression will be found. On output, the vector \mathbf{x} will contain these N roots.

The expression must contain the variable x, and may contain scalars, but must *not* contain any other non-scalar variables.

Muller's method

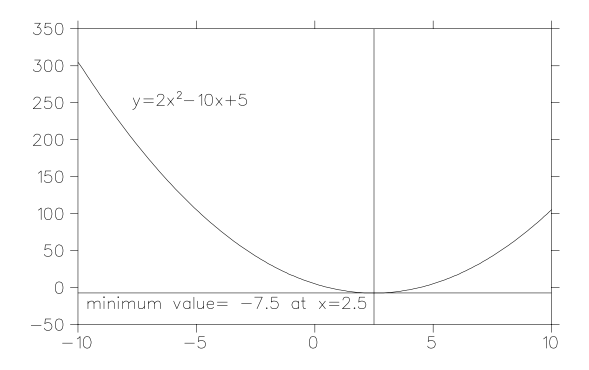


Figure 2.12: Finding a local minimum with the FMIN command

If z_{i-2} , z_{i-1} and z_i are three approximations to a root, the next approximation to the root, z_{i+1} , is taken as a zero of the quadratic that passes through $f(z_{i-2})$, $f(z_{i-1})$ and $f(z_i)$. The iteration continues by dropping z_{i-2} and repeating the quadratic fit for z_{i-1} , z_i and z_{i+1} and associated function values.

If (r-1) roots have been found, the r^{th} root is found by deflating f(z) and solving the equation $f_r(z)$ where

$$f_r(z) = f(z) / \prod_{i=1}^{r-1} (z - z_i)$$

where the z_i are the previously found roots. The roots are found one at a time in approximately increasing order.

The iteration stops when either

$$|(z_{i+1}-z_i)/z_{i+1}| < e_1$$
 or $f_r(z_{i+1})$ and $f(z_{i+1}) < e_2$

where e_1 and e_2 are error tolerances. For more information refer to A Method for Solving Algebraic Equations Using an Automatic Computer by D.E. Muller, M.T.A.C. 10, 1956, pages 208-215.

In the PHYSICA implementation of Muller's method, $e_1=10^{-7}$ and $e_2=10^{-20}$. The values

in vector x are interpreted as the user's initial guesses for the location of the roots. If x[i] = 0, the starting approximations for the i^{th} root are taken as: -1, 1, and 0. If $x[i] \neq 0$, the starting approximations for the i^{th} root are taken as: 0.9x[i], 1.1x[i], and x[i]. On each iteration, if $|r_t - r_p| < e_3$, where r_t is the approximation to a root, and r_p is a previously found root, then r_t is replaced by $r_t + e_4$. In PHYSICA, $e_3 = 10^{-20}$ and $e_4 = 10^{-4}$. If a root is not found in 60 iterations, the search is terminated.

Informational messages

By default, the values of the roots are displayed on the monitor screen. If the \NOMESSAGES qualifier is used, informational messages will be suppressed.

Example

The following script demonstrates how you can use the FZERO command to find roots. See Figure 2.13.

```
XMIN=-30
XMAX = 10
XD = [XMIN: XMAX: .1]
SET %XLABSZ 5
LABEL\XAXIS 'y=e<^>0.4x<_>-0.4x-9'
GRAPH XD EXP(0.4*XD)-0.4*XD-9
GET
YMIN YMIN
YMAX YMAX
NROOTS=2
X[1:NROOTS]=0
FZERO X EXP(0.4*X)-0.4*X-9
DO J = [1:NROOTS]
  GRAPH\NOAXES [X[J];X[J]] [YMIN;YMAX]
 DISPLAY 'please position the string: root at '//rchar(x[j])
 TEXT 'root at '//rchar(x[j])
ENDDO
ZEROLINE\HORIZONTAL
```

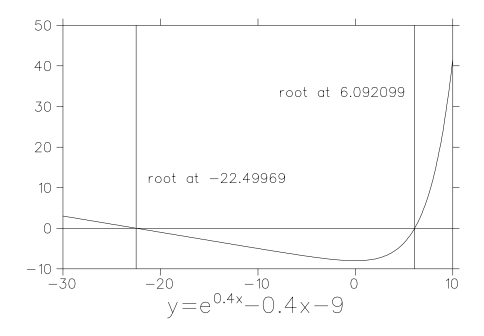


Figure 2.13: Finding roots with the FZERO command

GENERATE

| Syntax | GENERATE x min inc ,, npts |
|-----------|--------------------------------|
| | GENERATE x min ,, max npts |
| | GENERATE x min inc max |
| | GENERATE\RANDOM x min max npts |
| Qualifier | \RANDOM |
| Default | \-RANDOM |

The GENERATE command creates a new vector, x.

By default, the new vector, x, will be generated according to the formula: x[i] = min + (i-1) inc for $1 \le i \le npts$.

The minimum value, min, must be given. Two other values are also required: the increment and the number of points, or the maximum and the number of points, or the increment and the maximum value.

Increment and number of points given

```
Syntax GENERATE x min inc ,, npts
```

If the increment and the number of points are given, the above formula is applied directly.

Example

```
After the command: GENERATE X -1 .5 ,, 4 the vector X = [-1; -0.5; 0; 0.5]
```

Maximum and number of points given

```
Syntax GENERATE x min ,, max npts
```

If maximum value and the number of points are given, the increment is calculated: inc = (max - min)/(npts - 1) and then used in the usual formula.

Example

```
After the command: GENERATE X -1 ,, 2 4 the vector X = [-1; 0; 1; 2]
```

Increment and maximum given

```
Syntax GENERATE x min inc max
```

If the increment and the maximum value are given, the number of points will be ignored if entered. The usual formula will be applied until the next value would be greater than \max , the maximum. If the calculated maximum is different than the given maximum, a warning message will be displayed on the monitor screen. The calculated maximum will be the last value stored in the vector.

Example

After the command GENERATE X -1 .41 1 the warning message:

```
GENERATE warning: calculated maximum = 0.63999999E+00 given maximum = 0.10000000E+01
```

will be displayed on the monitor screen and the vector X = [-1; -0.59; -0.18; 0.23; 0.64]

Random numbers

```
Syntax GENERATE\RANDOM x min max npts
```

If the \RANDOM qualifier is used, the vector x will be filled with npts random numbers that fall

between min and max. No increment should be given.

The initial value for the random number seed is 12345. Every time a random number is requested, either from the GENERATE\RANDOM command or from the RAN function, the seed is updated. You can change the seed value with the SET SEED command.

Example

```
After the command GENERATE\RANDOM X 1 2 5
the vector X = [1.198525; 1.897874; 1.238289; 1.367985; 1.381705]
```

GET

```
Syntax GET { keyword { value }}

Examples GET %XLAXIS XLX
GET
GET NSXINC
```

The GET command gets the values of the GPLOT plot characteristic keywords as well as the PHYSICA specific keywords. Use the SET command to change the values of these keywords.

If the GET command is entered with no parameters, more than one keyword value can be obtained without re-entering the GET command. Other keywords and values will be requested, until a blank line is entered, at which time the user is put back into command line entry mode. If the GET command is used in this way in a script file, the blank line is *necessary* to indicate that the GET command is finished.

If a keyword is entered with the GET command, then only that one keyword's value can be obtained with that command.

If an output variable is not entered after the keyword, the current value of that keyword will be displayed on the terminal screen. If an output variable is entered, a variable will be created and the current value of that keyword assigned to that variable.

Note: The keywords FONT, CUNITS, UNITS, VERSION, VERSIONDATE, and AUTOSCALE return a string instead of a numeric value.

Examples

To display the current value of XMIN, enter: GET XMIN

To obtain the current value of XMIN and then change it to XMIN -10, and to set the value of XMAX to XMIN+100, enter:

```
GET XMIN A ! makes scalar A

SET

XMIN A-10

XMAX A+100 ! don't forget the blank line
```

To display the current graphics font name, enter GET FONT

The command: GET FONT TXT[3] places the current font name into the 3^{rd} element of the array text variable TXT.

The GPLOT keywords

See **Appendix A** for descriptions of all the GPLOT plotting characteristic keywords. The tables produced by the DISPLAY MENU contain most of the keywords that can be accessed with the SET and GET commands, along with their current values.

The GPLOT keywords: MASK, ALIAS, PMODE, PTYPE, and ERRBAR, should not be changed in PHYSICA, as these are internally adjusted and used by various commands.

The PHYSICA keywords

ARROLEN

ARROLEN is the arrow head length as a fraction of the total arrow shaft length. It is used for arrows drawn with the FIGURE command.

ARROTYP

ARROTYP controls the type of arrow drawn with the FIGURE command. See Table 2.54 on page 231 and Figure 2.24 on page 230.

ARROWID

ARROWID is the arrow head width as a fraction of the total arrow shaft length. It is used for arrows drawn with the FIGURE command.

PCHAR

```
Optional parameters: symbol { size { colour { angle }}}
```

PCHAR controls the plotting symbols, or the appearance of the histogram bars, when the GRAPH command is entered. GET PCHAR obtains the plotting symbol values or arrays, as set with the SET PCHAR command. It is not necessary to set the size, colour or angle with the SET PCHAR command, but if these are not set, you cannot request their values. The type of output variable that will be made for each parameter depends on the type of variable used in the SET PCHAR command.

AUTOSCALE

Note: The value of the AUTOSCALE keyword is a string instead of a numeric value. So, if you enter: GET AUTOSCALE X, the variable X will be a string variable.

The AUTOSCALE keyword controls autoscaling for graph axes. Autoscaling remains in effect until either the command SET AUTOSCALE OFF is entered, or the SCALES command is entered. Autoscaling affects commands that draw axes, for example, the commands GRAPH, CONTOUR, DENSITY, REPLOT, and SLICES.

Autoscaling means to automatically choose the minimum and maximum values for the axes, as well as the number of large, numbered, tic marks for the axes. The type of autoscaling that is done depends on the keyword that is used with the command.

| keyword | result |
|--------------|--|
| ON | Autoscale the horizontal and the vertical axes |
| OFF | turn off all autoscaling, |
| | the axes will appear as they are currently set |
| COMMENSURATE | Autoscale the horizontal and vertical axes and |
| | change the lengths of the axes so that they will be commensurate |
| XAXIS | Autoscale the horizontal axis only, |
| | the vertical axis will remain as currently set |
| YAXIS | Autoscale the vertical axis only, |
| | the horizontal axis will remain as currently set |

When the \VIRTUAL qualifier is used, the virtual minima and maxima for the axes will be determined, so that the axes may not begin or end at a large tic mark. If the keyword ON is used, both x- and y-axes will have virtual minima and maxima. If the keyword YAXIS is used, only the x-axis will have virtual minimum and maximum. If the keyword YAXIS is used, only the y-axis will have virtual minimum and maximum.

CNTSEP

CNTSEP, or %CNTSEP, is the separation between contour labels in the CONTOUR command. %CNTSEP is the separation as a percentage of the height of the window, that is, YUWIND-YLWIND, while CNTSEP is the separation expressed in centimeters or inches, depending on the units type as set with the SET UNITS command.

LABSIZ

LABSIZ, or %LABSIZ, is the size of the contour labels in the CONTOUR command. %LABSIZ is the size as a percentage of the height of the window, that is, YUWIND-YLWIND, while LABSIZ is the size expressed in centimeters or inches, depending on the units type as set with the SET UNITS command.

LEGSI7

LEGSIZ, or %LEGSIZ, is the size of the contour plot and density plot legend entries. %LEGSIZ is the size as a percentage of the height of the window, that is, YUWIND-YLWIND, while LEGSIZ is the size expressed in centimeters or inches, depending on the units type as set with the SET UNITS command.

LEGFRMT

Note: The value of the LEGFRMT keyword is a string instead of a numeric value. So, if you enter: GET LEGFRMT X, the variable X will be a string variable.

The numeric legend entries drawn by the DENSITY and CONTOUR commands are written using the LEGFRMT format.

ERRFILL

If the \ERRFILL qualifier is used with the READ\VECTOR command, an invalid field in the data file causes either the entire record to be filled with the value of ERRFILL if a format is used, or only that invalid field will be filled with ERRFILL if no format is used.

FILL

FILL is used in the FIGURE command, with the fillable figures: BOX, POLYGON, WEDGE, CIRCLE, ELLIPSE, and ARROWS with closed heads. It is also used for filling the boxes with the DENSITY\BOXES command. See Table 2.55 on page 233 for a description of the interpretations of the FILL keyword.

See the SET HATCH command for information on changing the hatch pattern definitions. See the DISPLAY HATCH command for information on how to display examples of the hatch

patterns.

HATCH

Optional parameters: n { v a }

The GET HATCH command is used for obtaining the hatch pattern definitions that are used for text bolding, for filling areas under curves or histograms, and for use by the TILE, PIEGRAPH, and FIGURE commands.

The SET HATCH command is used for changing the hatch pattern definitions. The SET HATCH command does *not* choose the hatch pattern to be used by other commands. It only alters the definition of a hatch pattern.

If just the keyword HATCH is entered, a table of the spacings and angles for all ten hatch patterns is displayed. If the hatch pattern number, n, is entered, then only pattern n will be displayed. If the hatch pattern number, an output vector, v, and an output scalar, a, are entered, then v will contain the spacings and a will contain the angle for that pattern number.

The hatch pattern number, n, should be between one and ten. A hatch pattern is composed of an angle and from one to ten spacings. The default spacings and angles are listed in Table 2.56 on page 235. The angles are in degrees and the spacing lengths, by default, are expressed in centimeters, but if the units are changed to inches, with the SET UNITS command, the lengths will be converted to inches. See Figure 2.7 on page 61 for examples of the hatch patterns. See the DISPLAY command, page 58, for information on how to display examples of the hatch patterns.

When an object is being filled, a line is drawn inside the object at the specified angle, then a parallel line is drawn at the first spacing, and so on for the number of spacings in that pattern. This process is repeated until the object is filled.

LINE

Optional parameters: n { v }

The GET LINE command is used for obtaining the definition of the line types that are used by the commands: GRAPH, LINE, PICK, ELLIPSE, FIGURE, and ZEROLINES.

The SET LINE command is used for changing the definition of the line types. This command does *not* choose the line type to be used by other commands. It only alters the definition of a line type. To choose a line type, use the SET LINTYP command.

If just the keyword LINE is entered, a table of the spacings for all ten line types is displayed. If the line type number, n, and an output vector, v, are entered, then nothing will be displayed. For example, to get line type 2 into vector X2, enter:

GET LINE 2 X2

See the SET LINE command for information on how the line types are defined. There are ten line types available. The defaults are listed in Table 2.58 on page 236. The lengths are expressed in centimeters, the default, but if the units are changed to inches, with the SET UNITS command, the lengths will be converted to inches. See Figure 2.8 on page 62 for examples of the default line types. See the DISPLAY command, page 58, for information on how to display examples of the line types.

TENSION

TENSION controls the spline tension for the functions using cubic splines:

DERIV, INTEGRAL, INTERP, SMOOTH, SPLINTERP, and SPLSMOOTH.

SEED

SEED is the random number seed value. This seed is updated whenever the GENERATE\RANDOM command is entered, or the RAN is used.

POSTRES

POSTRES controls the PostScript graphics output resolution, in dots per inch. This applies to dot filled text characters and dot types of DENSITY plots. The resolution can be changed at any time, so different parts of a single drawing can be drawn with different resolutions.

SPEED

SPEED controls the pen plotter speed. This applies to Hewlett-Packard, Houston, and Roland RDGL || pen plotters. The speed can be changed at any time, so different parts of a single drawing can be drawn at different speeds.

WIDTH

WIDTH controls the character width of the alphanumeric monitor screen. The value for WIDTH should be between 2 and 132.

XPREV

XPREV is the last world x-coordinate that was drawn by any graphics command. The value of this keyword is automatically updated.

YPREV

YPREV is the last world y-coordinate that was drawn by any graphics command. The value of this keyword is automatically updated.

NCURVES

NCURVES is the total number of data curves that have been drawn, using the GRAPH command, since the last CLEAR command. The value of this keyword is automatically updated.

UNITS

Note: The value of the UNITS keyword is a string instead of a numeric value. So, if you enter: GET UNITS X, the variable X will be a string variable.

UNITS controls the plotting units type, either centimeters, CM, the default, or inches, IN.

CUNITS

Note: The value of the CUNITS keyword is a string instead of a numeric value. So, if you enter: GET CUNITS X, the variable X will be a string variable.

CUNITS is the units type for the graphics cursor readout when the graphics cursor is invoked by the PICK, PEAK, LINE, or FIGURE command when running under X Windows and mouse button two is pressed. If WORLD is chosen, the numbers displayed depend on the current units type, either centimeters or inches, as chosen with SET UNITS. If GRAPH is chosen, the numbers displayed depend on the current graph axis scales.

FONT

Note: The value of the FONT keyword is a string instead of a numeric value. So, if you enter: GET FONT X, the variable X will be a string variable.

FONT controls the graphics font. For a list of the font names, see Table 2.60 on page 243.

The DISPLAY FONT command will draw a font table for any font.

VERSION

Note: The value of the VERSION keyword is a string instead of a numeric value. So, if you enter: GET VERSION X, the variable X will be a string variable.

This is the current program's version number. It is character valued, with a length of 5.

VERSIONDATE

Note: The value of the VERSIONDATE keyword is a string instead of a numeric value. So, if you enter: GET VERSIONDATE X, the variable X will be a string variable.

This is the current program's version date. It is character valued, with a length of 20.

SHOWHISTORY

SHOWHISTORY controls how many lines of history to display for each numeric variable as a result of the SHOW command.

SHOWHISTORY

```
n < 0 \rightarrow \text{all stored history lines will be displayed} n = 0, 1 \rightarrow \text{only the latest history line will be displayed} n > 0 \rightarrow \text{a maximum of n lines of history will be displayed for each variable}
```

MAXHISTORY

MAXHISTORY is the maximum number of history lines to store for each numeric variable. MAXHISTORY was added because if a variable had its value changed within a large DO loop, a new history line was added each time the loop was processed, which could lead to virtual memory problems.

WRAP

If WRAP = 0, history lines and string variable contents lines are not wrapped when displayed with the SHOW command. If WRAP is non-zero, these lines are wrapped.

DEVICE

Note: The value of the DEVICE keyword is a string instead of a numeric value. So, if you enter: GET DEVICE X, the variable X will be a string variable.

This is the current hardcopy device as chosen with the DEVICE command.

GLOBALS

Syntax GLOBALS

This command only works under VAX/VMS.

The GLOBALS command displays the names of global sections to which you have access. This command is meant to be used in conjunction with the MAP\FIOWA or the MAP\FIOWABIG commands.

GRAPH

```
Syntax GRAPH { 'legendtext' } x y { ye1 { xe1 { ye2 { xe2 }}}}

Qualifiers \AXESONLY, \NOAXES, \POLAR, \REPLOT, \HISTOGRAM

Defaults axes drawn, \REPLOT, \NOPOLAR, \NOHISTOGRAM, legendtext ignored

Examples GRAPH X Y

GRAPH 'legend entry' X Y YERR XERR

GRAPH\NOAXES X Y

GRAPH\HISTOGRAM X Y

GRAPH\POLAR RAD THETA
```

The GRAPH command draws: data with axes GRAPH

just the data GRAPH\NOAXES

just the axes GRAPH\AXESONLY

The data curve may be a histogram. The parameters must be vectors, but can be vectors of length one. The input vectors can have different lengths as the minimum length of all of the input vectors will be used.

Plotting symbols

The SET PCHAR command controls the plotting symbols, or the appearance of the histogram bars. For information on how to set the plotting symbol type, size, colour, and angle; as well as how to set hatch fill patterns, colours, and relative bar size for histograms, refer to the SET PCHAR command section.

Axis scaling

Autoscaling of the axes may apply if the two vectors, x and y, are entered. Refer to the SET AUTOSCALE command for information on how to set up autoscaling for the axes.

Use the SCALES command, page 226, to manually set the scales for the axes.

Graph legend

If LEGEND is ON, a legend entry is drawn into a legend frame box. A legend entry consists of a short line segment, with plotting symbol(s), and a string. The legend entry is drawn when the GRAPH command is entered. The string portion of the legend entry is expected as the first parameter of the GRAPH command, for example:

```
GRAPH 'legend entry' X Y
```

If LEGEND is OFF, a string entered as a first parameter with the GRAPH command is ignored. Refer to the LEGEND command, page 137, for more information on a graph legend.

Plotting data and axes

By default, if the GRAPH command is entered with *neither* the \AXESONLY qualifier *nor* the \NOAXES qualifier is used, then axes will be drawn as well as the data curve or histogram. Autoscaling will apply if it is on. For example:

```
GRAPH X Y ! plots axes and data curve GRAPH\HISTOGRAM X Y ! plots axes and histogram
```

Plotting axes only

If the GRAPH\AXESONLY command is entered with no parameters, autoscaling does not apply. The minima, maxima, and number of increments will be the same as the last set of axes drawn. The axes scales can be set up before entering the GRAPH\AXESONLY command, using the SCALES command. Autoscaling may apply if two parameters, x and y, are entered. For example:

Plotting data only

The GRAPH\NOAXES command plots the data, but does not draw axes. Autoscaling will not apply. The GRAPH\NOAXES command overlays on an existing set of axes. For example:

```
GRAPH\NOAXES X Y ! overlay data curve on current axes
GRAPH\HIST\NOAX X Y ! overlay histogram on current axes
```

If the LEGEND is ON then the data variables are not necessary. That is, you can enter:

```
GRAPH\NOAXES 'legendentry'
```

and an entry is made to the legend, but no curves are plotted. The plotting character, as set by the SET PCHAR command, will be used in the legend.

Replotting data on a common scale

By default, the graph will be stored for replotting. If the \NOREPLOT qualifier is used, the graph will not be saved for replotting. The default is \REPLOT .

To redraw a graph with multiple data sets so that all the data will appear within the axis boundaries, use the REPLOT command in association with the SET AUTOSCALE command. For example:

```
SET AUTOSCALE ON ! turn on autoscaling

SET PCHAR -1 ! set plotting character to 'box'

GRAPH X Y ! plot data with axes

SET PCHAR -2 ! set plotting character to 'cross'

GRAPH\NOAXES U V ! overlay another data curve

CLEAR\NOREPLOT ! clears graphics but not the replot buffers

REPLOT ! replot both data sets on common scale
```

Refer to the REPLOT command, page 208, for more information.

Histograms

It is possible to draw four types of histograms using the SET HISTYP approach, or you can use the \HISTOGRAM qualifier to plot a histogram with tails to y=0 and profile along the x-axis.

Using the HISTYP keyword

Table 2.39 on page 118 shows the histogram type that will be produced depending on the value of HISTYP.

Using the \HISTOGRAM qualifier

| HISTYP | Result |
|--------|--|
| 0 | (default value) line graph, not a histogram |
| 1 | histogram with no tails and profile along the x -axis. You may control the width and colour of each individual bar. |
| 2 | histogram with tails to $y=0$ and profile along the x -axis. You may control the filling pattern, width and colour of each individual bar |
| 3 | histogram without tails and profile along the y -axis. You may control the height and colour of each individual bar. |
| 4 | histogram with tails to $x=0$ and profile along the y -axis. You may control the filling pattern, height and colour of each individual bar |

Table 2.39: The HISTYP keyword

The \HISTOGRAM qualifier is inconsistent when used in conjunction with the \AXESONLY qualifier.

Using the \HISTOGRAM qualifier is equivalent to using a HISTYP setting of 2. A histogram with tails to y = 0 and profile along the x-axis will be plotted. The following three commands:

```
SET HISTYP 2 ! force histogram plotting
GRAPH X Y ! plot axes and histogram
SET HISTYP 0 ! reset to default value
```

are equivalent to the single command:

```
GRAPH\HIST X Y
```

Filling

To fill the area under a histogram, you can use the SET PCHAR command. The SET PCHAR command allows you to fill each histogram bar with a different fill pattern. This only applies to histograms with tails, ${\tt HISTYP}=2$ or 4.

See the SET PCHAR command for more information on filling, colours, and relative bar size for histograms.

See the SET HATCH command, page 228, for information on changing the hatch pattern defi-

nitions.

See the DISPLAY command for information on how to display examples of the hatch patterns.

Polar coordinates

If the \POLAR qualifier is used, the input vectors, x and y, are assumed to represent polar coordinates, where x contains the radial components and y contains the angular components, in degrees. The polar coordinates are transformed to rectangular coordinates before plotting, but the vectors x and y are returned unchanged.

Error bars

```
Syntax GRAPH { 'legendtext' } x y { ye1 { xe1 { ye2 { xe2 }}}}
```

The optional vectors ye1, xe1, ye2, and xe2 are interpreted as errors for drawing error bars. You can have symmetric or asymmetric error bars.

Symmetric error bars

For symmetric error bars, the error variable should contain one half of the total error. See Table 2.40.

| parameters | |
|-----------------|---|
| present | Result |
| ye1 but not ye2 | symmetric vertical error bars will be drawn at the point (x[j],y[j]), the |
| | error bar is drawn from y[j]-ye1[j] to y[j]+ye1[j] |
| xe1 but not xe2 | symmetric horizontal error bars will be drawn at the point $(x[j],y[j])$, the error bar is drawn from $x[j]-xe1[j]$ to $x[j]+xe1[j]$ |

Table 2.40: Symmetric error bars

Asymmetric error bars

For asymmetric error bars, the first error variable contains the lower error and the second error variable contains the upper error. See Table 2.41.

Error bar shape

The error bars will have "feet", that is, short line segments, one at each end of the error bar, which are perpendicular to the error bar. The size of the foot is the same as the size of the plotting symbol, which can be changed using the SET %CHARSZ command or by

| parameters | |
|-------------|--|
| present | Result |
| ye1 and ye2 | asymmetric vertical error bars are drawn at the point $(x[j],y[j])$, the |
| | error bar is drawn from y[j]-ye1[j] to y[j]+ye2[j] |
| xe1 and xe2 | asymmetric horizontal error bars are drawn at the point $(x[j],y[j])$, the error bar is drawn from $x[j]-xe1[j]$ to $x[j]+xe2[j]$ |

Table 2.41: Asymmetric error bars

entering a relative size vector with the SET PCHAR command. The error bar will be clipped at the boundaries of the plotting symbol if the symbol is symmetric under 90° rotations, for example, a box (symbol number 1).

Filling

To fill the area under a curve, use the SET LINTYP command. See also the histogram discussion above, for specifics on filling histograms. If $101 \le \texttt{LINTYP} \le 110$, then the hatch pattern LINTYP-100 is chosen. If $211 \le \texttt{LINTYP} \le 299$, then the dot fill pattern LINTYP-200 is chosen. The polygonal region defined by the (x,y) coordinate pairs, with the last point connected to the first, will be filled with the chosen hatch or dot pattern.

Hatch patterns

 $101 \le |\texttt{LINTYP}| \le 110$ means to fill using a hatch pattern. The filling will be done with hatch pattern |LINTYP|-100. For example, if LINTYP = 108, then hatch pattern number 8 will be used.

A hatch pattern is composed of an angle and one to ten spacings. These spacings are simply cycled through as the region is being filled, that is, a line is drawn inside the region at the specified angle, then a parallel line is drawn at the first spacing, then another parallel line is drawn at the second spacing, and so on for the number of spacings in that pattern. This process is repeated until the region is filled. The hatch patterns can be redefined with the SET HATCH command and displayed with the DISPLAY FILL command. There are ten hatch patterns available.

Dot fill patterns

 $211 \le |\mathtt{LINTYP}| \le 299$ means to fill using a dot pattern. The filling will be done with dot pattern $|\mathtt{LINTYP}| - 200$. For example, if $\mathtt{LINTYP} = 234$, then dot pattern 34 will be used. If $\mathtt{LINTYP} < 0$, then the dots are erased instead of turned on.

A dot pattern is of the form: uv, where the digit u is the increment number of dots to light up horizontally, $1 \le u \le 9$, and the digit v is the increment number of dots to light up vertically, $1 \le v \le 9$. For example, a dot pattern of 34 means to light up every third dot horizontally and every fourth dot vertically. If uv is negative, then the dots are erased instead of turned on. Note that 200 is interpreted the same as 211, that is, every dot is lit.

PostScript output

For PostScript output, set the POSTRES keyword to the appropriate resolution for your hard-copy device, using the SET command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark inkjet printer has a resolution of 360 dpi. A "good" picture can be obtained with POSTRES = 180 and LINTHK = 2.

Examples

The following script demonstrates the plotting of error bars. See Figure 2.14.

```
X = [0:20]
                     ! generate some "data"
Y=X^2-20*X+50
YEL=10*RAN(X)
                     ! lower error
YEU=10*RAN(X)
                     ! upper error
YES=20*RAN(X)
                     ! symmetric error
SET PCHAR -1
                     ! set plotting symbol
WINDOW 5
                     ! set window
LABEL\X 'Asymmetric errors'
GRAPH X Y YEL,, YEU ! plot with asymmetric errors
WINDOW 6
                     ! change windows
LABEL\X 'Symmetric errors'
GRAPH X Y YES
                     ! plot with symmetric errors
```

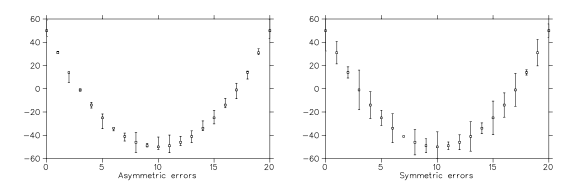


Figure 2.14: Plotting error bars with the GRAPH command

-20

-40

-60 |

10

Dot fill

15

The following script demonstrates filling under area under a curve. See Figure 2.15.

```
X = [0:20]
                      ! generate some "data"
Y=X^2-20*X+50
XX[1]=X[1]
                      ! fix up the data so it starts at (x[1],0)
XX[2:LEN(X)+1]=X
                     ! and ends at (x[#],0)
XX[LEN(X)+2:LEN(X)+2]=X[#]
YY[1]=0
YY[2:LEN(Y)+1]=Y
YY[LEN(Y)+2:LEN(Y)+2]=0
SET PCHAR O
                     ! no plotting symbol
SET LINTYP 233
                     ! dot pattern (every 3rd dot both directions)
WINDOW 15
                     ! set window
LABEL\X 'Dot fill'
                     ! label the x axis
GRAPH XX YY
ZEROLINE\HORIZONTAL ! draw horizontal line through (0,0)
SET LINTYP 108
                     ! hatch pattern number 6
WINDOW 16
                      ! change windows
LABEL\X 'Hatch fill' ! label the x axis
GRAPH XX YY
ZEROLINE\HORIZONTAL ! draw horizontal line through (0,0)
                                              40
        40
        20
                                              20
        0
                                              0
```

Figure 2.15: Filling the area under a curve drawn with the GRAPH command

The following script demonstrates plotting histograms and filled histograms. See Figure 2.16.

-20

-40

-60 |

10

Hatch fill

X = [0:24:3]! generate some "data" Y=X^2-20*X+50 ! histogram with no tails SET HISTYP 1 WINDOW 5 ! set window LABEL\X 'HISTYP = 1' ! label the x axis GRAPH X Y ! plot the histogram WINDOW 6 ! set window LABEL\X 'Narrow bars' ! label the x axis SET PCHAR 0 .8 GRAPH X Y ! plot the histogram WINDOW 7 ! set window SET HISTYP 2 ! histogram with tails LABEL\X 'Hatch pattern #8' SET PCHAR 8 .8 ! hatch pattern #8 GRAPH X Y ! plot the histogram ZEROLINES\HORIZONTAL ! draw horizontal line thru (0,0) WINDOW 8 ! set window LABEL\X 'Individual bar filling' SET PCHAR [11:99:11] .8 ! each bar filled GRAPH X Y ! plot the histogram ZEROLINES\HORIZONTAL ! draw horizontal line thru (0,0) 150 150

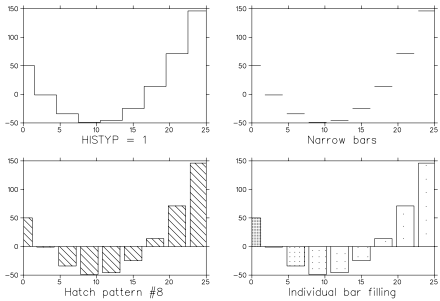


Figure 2.16: Histogram examples drawn with the GRAPH command

GRID

The GRID command creates a regular matrix from scattered data points. The three vectors, x, y and z, are assumed to represent scattered points, where z[i] is the altitude corresponding to the coordinates (x[i],y[i]). Suppose l = min(len(x),len(y),len(z)). By default, a square matrix, m, is interpolated, with row and column dimensions equal to $5 \times \sqrt{l}$.

The coordinates of element m[i,j] of the output matrix will be (xout[j],yout[i]). The columns of m are of constant x, and the rows are of constant y.

Polar coordinates

The \NOINTERPOLATE qualifier cannot be used with the \POLAR qualifier.

If the \POLAR qualifier is used, x is assumed to contain the radial components and y is assumed to contain the angular components, in degrees. The output matrix, m, will be regular in polar coordinates, with the columns of constant radius and the rows of constant angle.

Duplicate points

By default, duplicate (x,y) locations are not checked for before the matrix is made. If you want duplicate points to be ignored, use the \CHECKDUP qualifier.

Interpolated grid

```
Syntax GRID x y z m

Qualifiers \POLAR, \SIZE, \XYOUT, \BOUNDS, \CHECKDUP

Defaults \NOPOLAR, \NOSIZE, \NOXYOUT, \NOBOUNDS, \NOCHECKDUP
```

By default, the set of scattered data points is used to construct a Thiessen triangulation of the plane and a regular matrix, m, is interpolated.

Output matrix size

```
Syntax GRID\SIZE s x y z m
```

Suppose l = min(len(x),len(y),len(z)). By default, the interpolated matrix will be square, with row and column dimensions both equal to $5 \times \sqrt{l}$. If another size, s, is desired, you must use the \SIZE qualifier, and the row and column dimensions will be both equal to s.

Output vectors

```
Syntax GRID\XYOUT x y z m xout yout
```

If output vectors, xout and yout, are desired, you must use the \XYOUT qualifier. The coordinates of output matrix element m[i,j] will be (xout[j], yout[i]), where xout contains the x-coordinates of each column and yout contains the y-coordinates of each row.

Range of interpolation

```
Syntax GRID\BOUNDS x y z m minx maxx miny maxy
GRID\BOUNDS\XYOUT x y z m xout yout minx maxx miny maxy

Defaults \NOBOUNDS, interpolation range = range of x and y
```

By default, the range of the grid interpolation is the range of values of the vectors x and y. If the \BOUNDS qualifier is used, this range is specified by the final four numbers, minx, maxx, miny, and maxy.

Non-interpolated grid

```
Syntax GRID\PATTERN x y z m

Qualifiers \XYOUT, \CHECKDUP

Defaults \NOXYOUT, \NOCHECKDUP
```

Suppose the vectors x and y have length h, and suppose that for some n1 and n2, x and y have the following pattern:

$$x_1$$
 = x_2 = \cdots = x_{n2}
 x_{n2+1} = x_{n2+2} = \cdots = x_{n2+n2}
 \vdots \vdots
 $x_{(n1-1)n2+1}$ = $x_{(n1-1)n2+2}$ = \cdots = $x_{n1\cdot n2}$

```
y_1 = y_{n2+1} = \cdots = y_{(n1-1)n2+1}

y_2 = y_{n2+2} = \cdots = y_{(n1-1)n2+2}

\vdots

\vdots

y_{n2} = y_{n2+n2} = \cdots = y_{n1,n2}
```

where $h=n1\cdot n2$. If the x and y vectors have this form, it is possible to construct a matrix, without interpolation, with n2 rows and n1 columns, that is, $m_{i,j}=z_k$ where k=j+(i-1)n1 for $i=1,2,\ldots,n2$ and for $j=1,2,\ldots,n1$.

Output vectors

```
Syntax GRID\PATTERN\XYOUT x y z m xout yout
```

If output vectors, xout and yout, are desired, you must use the \XYOUT qualifier. The coordinates of output matrix element m[i,j] will be (xout[j],yout[i]), where xout contains the x-coordinates of each column and yout contains the y-coordinates of each row. If the output matrix has n1 columns and n2 rows, then the length of xout will be n1 and the length of yout will be n2.

```
\mathbf{xout} = [x_1; x_{n2+1}; \cdots; x_{(n1-1)n2+1}]
\mathbf{yout} = [y_1; y_2; \cdots; y_{n2}]
```

Example

The vectors:

```
X = [1; 1; 1; 1; 2; 2; 2; 2; 3; 3; 3; 3]

Y = [1; 2; 3; 4; 1; 2; 3; 4; 1; 2; 3; 4]

Z = [1; 2; 3; 4; 5; 6; 7; 8; 9; 10; 11; 12]
```

have the proper form, with n2=4 and n1=3. If you entered the command <code>GRID\PATTERN\XYOUT X Y Z M XO YO</code> the resultant variables would be:

$$\mathbf{M} = \begin{pmatrix} 1 & 5 & 9 \\ 2 & 6 & 10 \\ 3 & 7 & 11 \\ 4 & 8 & 12 \end{pmatrix} \mathbf{XO} = [1;2;3], \ \mathbf{YO} = [1;2;3;4]$$

Matrix from sparse data

```
Syntax GRID\INDICES x y z m

Qualifiers \XYOUT, \CHECKDUP

Defaults \NOXYOUT, \NOCHECKDUP
```

The vectors \mathbf{x} and \mathbf{y} are assumed to contain index locations for the \mathbf{z} data values.

Suppose that h is the minimum length of x, y, and z; and nc = max(x[i]), nr = max(y[i]) for i = 1, ..., h.

Then m[i,j] = 0 for i = 1, ..., nr; j = 1, ..., nc except m[y[i],x[i]] = z[i] for i = 1, ..., h. m will have nr rows and nc columns.

Output vectors

```
Syntax GRID\INDICES\XYOUT x y z m xout yout
```

If output vectors, xout and yout, are desired, you must use the \XYOUT qualifier. The coordinates of output matrix element m[i,j] will be (xout[j], yout[i]), where xout contains the x-coordinates of each column and yout contains the y-coordinates of each row. If the output matrix has no columns and no rows, then xout = [1:nc] and yout = [1:nr].

Example

Suppose

```
X = [1; 4; 1; 3; 5]

Y = [2; 1; 6; 4; 6]

Z = [10; 15; 20; 25; 30]
```

after the command: GRID\INDICES\XYOUT X Y Z M XO YO

the resultant variables will be:

HARDCOPY

```
Syntax HARDCOPY keyword { queue }
HARDCOPY keyword { file }
```

The HARDCOPY command is used for obtaining graphics hardcopies for the currently active device type. The initial default graphics hardcopy type is an HP LaserJet bitmap at 150 dpi. Use the DEVICE command to choose a different graphics hardcopy device type.

Printing and saving

The first command parameter is a keyword which refers to the action of printing or saving the graphics. This depends on which graphics hardcopy device is currently active. Refer to Table 2.42 for a listing of the appropriate hardcopy print and save codes for the bitmap device types. Refer to Table 2.43 for a listing of the appropriate hardcopy print and save codes for the non-bitmap device types.

| device | action | | | |
|------------|--------|---------------------------|--|--------------|
| keyword | code | parameter | description | default |
| | P | { queue } | print on a queue | HP\$LASER |
| | PC | $\{$ queue $\}$ | print on a queue -compressed- | HP\$LASER |
| | S | $\{ \ \mathtt{file} \ \}$ | save in a file | HPLASER.PLT |
| HPLASER | Α | | auxiliary port output | |
| | T | $\{ \ \mathtt{file} \ \}$ | T _E Xoutput file file | HPTEX.PLT |
| | TC | $\{ \ \mathtt{file} \ \}$ | T _E Xoutput file -compressed- | HPTEX.PLT |
| | TJ | $\{ \ \mathtt{file} \ \}$ | T _E Xoutput file -justified- | HPTEX.PLT |
| | TJC | $\{ \ \mathtt{file} \ \}$ | TEXoutput file -compressed-justified- | HPTEX.PLT |
| | P | { queue } | print on a queue | INK_JET |
| | PT | $\{$ queue $\}$ | print on a queue -transparency- | INK_JET |
| INKJET | S | $\{ \ \mathtt{file} \ \}$ | save in a file | INKJET.PLT |
| | Α | | auxiliary port output | |
| | P | { queue } | print on a queue | HP\$THINKJET |
| HPTHINKJET | S | $\{ \ \mathtt{file} \ \}$ | save in a file | HPTHINK.PLT |
| | Α | | auxiliary port output | |
| | P | { queue } | print on a queue | LNPTR |
| PRINTRONIX | S | $\{ \ \mathtt{file} \ \}$ | save in a file | PX.PLT |
| | A | | auxiliary port output | |

Table 2.42: HARDCOPY command print and save codes for bitmap devices

Note: Compressed format can speed up the printing of a drawing by as much as a factor of 5. Compressed format is recognized *only* by the LaserJet IIP, the LaserJet III, and later model

| device | action | | | |
|------------|--------|---------------------------|-----------------------|----------------|
| keyword | code | parameter | description | default |
| | Р | { queue } | print on a queue | POST\$SCRIPT |
| POSTSCRIPT | S | $\{ \ \mathtt{file} \ \}$ | save in a file | POSTSCRIPT.PLT |
| | A | | auxiliary port output | |
| | P | { queue } | print on a queue | HPLTR |
| HPPLOTTER | S | $\{ \ \mathtt{file} \ \}$ | save in a file | HPP.PLT |
| | Α | | auxiliary port output | |
| GKS | S | $\{ \text{ file } \}$ | save file | GKS.PLT |
| | P | { queue } | print on a queue | HOUSTON |
| HOUSTON | S | $\{ \ \mathtt{file} \ \}$ | save in a file | HOUSTON.PLT |
| | A | | auxiliary port output | |
| | P | { queue } | print on a queue | IMAGEN |
| IMAGEN | S | $\{ \ \mathtt{file} \ \}$ | save in a file | IMAGEN.PLT |
| | Α | | auxiliary port output | |
| | Р | { queue } | print on a queue | LA100 |
| LA100 | S | $\{ \ \mathtt{file} \ \}$ | save in a file | LA100.PLT |
| | Α | | auxiliary port output | |
| | Р | { queue } | print on a queue | LN03 |
| LNO3+ | S | $\{ \ \mathtt{file} \ \}$ | save in a file | LN03.PLT |
| | A | | auxiliary port output | |
| | Р | { queue } | print on a queue | RDGL |
| ROLAND | S | $\{ \ \mathtt{file} \ \}$ | save in a file | RDGL.PLT |
| | A | | auxiliary port output | |

Table 2.43: HARDCOPY command print and save codes for non-bitmap devices

printers. Do *NOT* use compressed format output on other than these devices.

Examples

Suppose that the Hewlett-Packard pen plotter output has been enabled, with the command DEVICE HPPLOTTER. To print the graphics directly on queue QUENAME, enter:

```
HARDCOPY P QUENAME
```

Suppose that the HP LaserJet bitmap has been enabled with the DEVICE HPLASERJET command. To save the graphics in a file, FILE.PLT, for inclusion in a TeXor Lagrange TeXdocument in justified format, enter:

```
HARDCOPY TJ FILE.PLT
```

Suppose a PostScript printer has been enabled with the DEVICE POSTSCRIPT command. To save the graphics in a PostScript file, FILE.PSC, to be printed later, enter:

HARDCOPY S FILE.PSC

HELP

```
Syntax HELP { string ... }

HELP\LIBRARY libname { string ... }

Qualifiers \PAGE, \LIBRARY

Defaults \NOPAGE, \NOLIBRARY
```

For VMS users:

The HELP command invokes the on-line help facility. To get information about a specific topic, include it as the string with the HELP command. To leave help quickly, type control-z.

Paging the output

By default, output to the screen is not displayed by pages, but output to the screen continues until the information display ends. If the \PAGE qualifier is used, output to the screen is paged, that is, the output stops after each screen full of information is displayed.

User defined library

The HELP\LIBRARY command allows the user to specify a help library other than the default PHYSICA help library. The full filename specification is required, that is:

```
disk:[directory]libname.hlb
```

The \LIBRARY qualifier can be used with the \PAGE qualifier.

For UNIX users:

The HELP command invokes an on-line help facility that mimics the built-in VMS help facility. To browse the help information, enter just the HELP command. To get information about a specific topic, include it as the string with the HELP command, but then no subtopics will be displayed. To leave help quickly, type control-d.

INPUT

```
Syntax INPUT x1 { x2 ... x8 }
INPUT\MATRIX m nr nc

Qualifier \MATRIX

Default create vectors
```

The INPUT command is used to interactively enter data, from the terminal keyboard, into vector(s) or into a matrix. The default is to create vectors.

If the INPUT command is used in a script file, input will still be expected from the terminal keyboard.

Vectors

```
Syntax INPUT x1 { x2 ... x8 }
```

Enter one set of numbers per line, that is, enter $x1[i] \dots x8[i]$ all on one line. The maximum number of vectors that can be input with one command is 8.

VMS: Input is terminated by typing control-z.

UNIX: Input is terminated by typing control-d.

Making corrections

If you want to change a previous entry, say the n^{th} entry, enter In before the new entries. All entries must be present. For example, suppose you are creating three vectors, X, Y, and Z, and when you are entering the fourth set of numbers, you realize you have made a mistake in the second entry. The following example shows how you could correct the mistake.

```
PHYSICA: INPUT X Y Z
Enter 3 numbers, (control-Z ends)
(1) >> 10 .1 1000
(2) >> 20 2 2000
(3) >> 30 .3 3000
(4) >> 12 20 .2 2000
(4) >> 40 .4 4000
(5) >> 50 .5 5000 <control-z>
PHYSICA:
```

and then you will have the following vectors:

```
X = [ 10; 20; 30; 40; 50 ]
Y = [ 0.1; 0.2; 0.3; 0.4; 0.5 ]
Z = [ 1000; 2000; 3000; 4000; 5000 ]
```

Matrix

```
Syntax INPUT\MATRIX m nr nc
```

The INPUT\MATRIX command is used to interactively enter data, from the terminal keyboard, into a matrix. A new matrix will be created.

The number of rows, nr, and the number of columns, nc, must be entered. Enter one row of the matrix, that is, nc numbers, per line. Only nr lines will be requested.

Making corrections

If you want to change a previous row, say the \mathbf{n}^{th} row, enter In before the new entries. All entries must be present. For example, suppose you are creating a matrix, M, with 5 columns and 3 rows. When you are entering the third set of numbers, you realize you have made a mistake in the second set. The following example shows how you could correct the mistake.

```
PHYSICA: INPUT\MATRIX M 3 5
Enter 5 numbers

(row 1) >> 1 2 3 4 5

(row 2) >> 6 7 8 9 10

(row 3) >> 12 6 7 -8 9 10

(row 3) >> 11 12 13 14 15
PHYSICA:
```

and you will have the following matrix:

$$\mathbf{M} = \left(\begin{array}{ccccc} 1 & 2 & 3 & 4 & 5 \\ 6 & 7 & -8 & 9 & 10 \\ 11 & 12 & 13 & 14 & 15 \end{array}\right)$$

INQUIRE

```
Syntax INQUIRE 'prompt string' v1 { v2 ... }

Defaults if vI does not exist, it is assumed to be a scalar

Examples INQUIRE 'Enter a value >>' A

INQUIRE 'Enter YES or NO >>' TXT
```

The INQUIRE command is intended for use in script files. The prompt string is written to the monitor screen and you are expected to enter the correct number and type of values, corresponding to the variable names following the prompt string. You can inquire for scalars, vectors, matrices, or string variables.

If *just* a carriage return is typed in response to an INQUIRE prompt, the variable(s) that are being requested will keep their current value(s). This allows you to have default values for inquired variables.

If variable vI does not exist, it is assumed to be a scalar. If vI is a scalar, the user is expected to enter a literal constant or a scalar. For example,

```
S=3 ! default value is 3
INQUIRE 'Enter scalar >> ' S
```

If vI is a vector, the user is expected to enter a set of values or a vector. For example,

```
V=[1:10] ! default values
INQUIRE 'Enter vector >> ' V
```

If vI is a matrix, the user is expected to enter a matrix. For example,

```
MATRIX M 5 5
INQUIRE 'Enter matrix >> ' M
```

If vI is a string variable, the user is expected to enter a literal string or a string variable. For example,

```
A='yes' ! default is yes
```

```
INQUIRE 'Enter Yes or No >> ' A
```

Examples

Suppose you want a script that asks the user a question requiring a yes or no response, with the script branching depending on the response. Following is an example of this procedure. Note that the response is converted to uppercase, since it must be in uppercase for the checks to find a YES or NO.

```
DUM='yes'
                                    ! make a dummy string variable
ASK:
                                    ! a label
INQUIRE 'Enter YES or NO >>' DUM ! get a response from the keyboard
IF EQS(UCASE(DUM), 'YES') THEN
                                            ! check if YES
  . . .
  . . .
  GOTO NEXT
                                    ! it was YES, so continue
ENDIF
IF EQS(UCASE(DUM), 'NO') THEN
                                            ! check if NO
  . . .
  . . .
  GOTO NEXT
                                    ! it was NO, so continue
DISPLAY 'Invalid response: expecting YES or NO'
                                    ! go back and ask again
GOTO ASK:
NEXT:
                                    ! continue with the script
 . . .
 . . .
```

In the following example, the first command defines X to be a vector of length 1. The second command defines S to be a scalar. The third command creates a scalar N and assigns it the value 5.

```
VECTOR X 1 
 SCALAR S 
 N = 5 
 INQUIRE 'Enter a vector with '//RCHAR(N)//' entries and a scalar >>' X S
```

The INQUIRE command will write out the prompt and wait for you to type in a vector set and a single value. For example:

```
Enter a vector with 5 entries and a scalar >> [0;1;3.5;-4;100] 10.3
```

JOURNAL

Syntax JOURNAL filename Qualifiers \APPEND, \MACRO

Defaults \NOAPPEND, \NOMACRO, initial journal file = PHYSICA.JOURNAL

A journal file is a record of all user input to PHYSICA, as well as all non-graphics output, such as error messages, from PHYSICA. The initial state is to have the journaling of interactive input and output on, journaling of script file input and output off, and the initial journal file name is PHYSICA. JOURNAL.

Journaling can be disabled by entering the DISABLE JOURNAL command. This closes the file. Enter ENABLE JOURNAL to open the last journal file that was open, and to append subsequent journal entries to this file. To also enable the writing of script file input and output to the journal file, use the ENABLE JOURNAL\MACRO command.

To just disable journaling of script file input and output, use the DISABLE JOURNAL\MACRO command.

To open a new journal file, use the JOURNAL command. To open with append, use the \APPEND qualifier.

To find out whether a journal file is currently open, and if so, to display the name of the current journal file, enter the STATUS command.

Environment variables in file names

For UNIX users, it is now possible to use an environment variable in a file name, if the environment variable is preceded by a \$. For example,

```
setenv FILE myjournal
physica
journal $FILE
```

The environment variable can be just the first part of the filename, for example,

```
setenv FILE my
physica
journal $FILEjournal
```

KEYWORD

Syntax KEYWORD

The KEYWORD command enters an interactive mode, where you type a keyword and the online help locations of that keyword are displayed. The help locations are separated by blanks, while vertical bars, |, separate the levels within each location. For example, typing the keyword shell will display the help locations DISABLE|SHELL ENABLE|SHELL. You could then find information on shell by typing HELP DISABLE SHELL or HELP ENABLE SHELL.

The wildcard is *. An initial wildcard and/or a final wildcard are allowed. For example, *inc* displays inch inches nlxinc nlyinc nsxinc nsyinc which are valid keywords; while inc* displays inch inches.

Typing a TAB, or control-I, is similar to using a final wildcard, in that all matching keywords are displayed, unless there is a unique keyword, and then the keyword is completed for you.

LABFL

```
Syntax LABEL { 'textstring' }
Qualifiers \XAXIS, \YAXIS

Defaults no x-axis label, no y-axis label

Examples LABEL\XAXIS TXTVAR[5][1:10]
LABEL\XAXIS
LABEL\YAXIS 'This'//CHAR(39)//'s a label with a single quote'
LABEL\YAXIS '<alpha,beta,^>10'
```

The LABEL command sets the automatic x- or y-axis text label. Use the \XAXIS qualifier to set the x-axis label and use the \YAXIS qualifier to set the y-axis label.

The default font can be changed with the SET FONT command. To change the size of the x-axis text label use the SET command, with either XLABSZ or %XLABSZ. To change the size of the y-axis text label use the SET command, with either YLABSZ or %YLABSZ.

The x-axis automatic text label is drawn, centred, below the x-axis. The x-axis label is drawn only if and when the x-axis is drawn. The y-axis automatic text label will be drawn, centred, to the left of the y-axis. The y-axis label is drawn only if and when the y-axis is drawn. The commands that draw axes are: GRAPH, CONTOUR, DENSITY, and REPLOT.

Turning off the labels

Entering the LABEL\XAXIS command without a parameter indicates that no automatic x-axis text label is to be drawn.

Entering the LABEL\YAXIS command without a parameter indicates that no automatic y-axis text label is to be drawn.

Example

The following script demonstrates automatic axis labeling, and produced Figure 2.17 on page 137.

```
LABEL\YAXIS '<alpha,beta,gamma> X-AXIS TEST'
LABEL\YAXIS 'Y-AXIS TEST <sigma,^>2<_> > 5'
GRAPH\AXESONLY
```

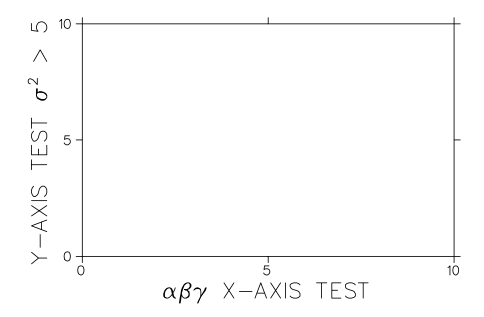


Figure 2.17: Automatic axis labels using the LABEL command

LEGEND

```
LEGEND ON | OFF

LEGEND FRAME ON | OFF | { xlo ylo xhi yhi }

LEGEND TRANSPARENCY ON | OFF

LEGEND AUTOHEIGHT ON | OFF

LEGEND NSYMBOLS n

LEGEND TITLE { 'title string' }

LEGEND STATUS

Defaults

legend off, frame on, transparency on, autoheight on,

\PERCENT, (xlo,ylo) = (55,22.5), (xhi,yhi) = (78,45),

number of plotting symbols = 1, no title
```

If LEGEND is ON, a legend entry is drawn into a legend frame box when the GRAPH command is entered. A legend entry consists of a short line segment with plotting symbols, and a string.

The string portion of a legend entry

If LEGEND is ON, the string portion of a legend entry is expected as the first parameter of the GRAPH command. For example:

```
GRAPH 'legend entry' X Y
```

If LEGEND is OFF, a string entered as a first parameter with the GRAPH command is ignored.

The string may contain formatting commands. See the PLOTTEXT command, page 165, for information on text formatting.

If LEGEND is ON then the data variables are not necessary, that is, you can enter: GRAPH\NOAXES 'Legend entry' and an entry is made to the legend but no curves are plotted.

Legend entry text height

Syntax LEGEND AUTOHEIGHT ON

LEGEND AUTOHEIGHT OFF

Default ON

By default, the height of the string portion of the legend entry will be determined so that the string fits "lengthwise" in the legend frame box. If you want to specify the height of the string portion of the legend entry, enter the LEGEND AUTOHEIGHT OFF command. The default is AUTOHEIGHT ON. When AUTOHEIGHT is OFF, the text height will be the current value of TXTHIT, as specified with the SET command. You can also include the text height as a text formatting command, <hr/>
'Hnn.n'> or <hr/>
'Hnn.n'>, within the string.

Do not include a text height formatting command when AUTOHEIGHT is ON. This will cause infinite looping, as the program attempts to adjust the text height to fit within the frame.

The line segment portion of a legend entry

The line segment portion of a legend entry is drawn with the same line type and plotting symbol as the corresponding data curve.

Plotting symbols

Syntax LEGEND NYSMBOLS n

Default number of symbols = 1

The number of plotting symbols drawn on the line segment can be specified by using the

NSYMBOLS keyword. The default is n=1, to plot one symbol. If one symbol is drawn, it will be in the middle of the line segment. If more one symbol is drawn, they will be equally spaced along the line segment, with one symbol at each end. If no symbols are drawn, the line segment will still be drawn. If no symbols are plotted on the data curve, no symbols will appear in the legend entry. Only the first n plotting symbols used for the data curve will be drawn in the legend entry.

Plotting symbols for the data curve are chosen with the SET PCHAR command.

The frame box

```
Syntax LEGEND FRAME { units } { xlo ylo xhi yhi }
LEGEND FRAME ON
LEGEND FRAME OFF

Defaults ON, \PERCENT, (xlo,ylo) = (55,22.5), (xhi,yhi) = (78,45)
```

Use the FRAME keyword to control the legend frame box. Legend entries are drawn starting from the top of the frame box. No clipping is done at the edges of the frame box, so entries may appear outside the frame box.

If LEGEND FRAME OFF is entered, the outline of the frame box will not be drawn. The default is ON, that is, to draw the legend frame box outline. If neither OFF nor ON is entered, it is assumed that you are entering the corner coordinates of the frame box.

Coordinates

The coordinates of the lower left corner of the legend frame box are (xlo,ylo), and the coordinates of the upper right hand corner are (xhi,yhi). The default lower left corner is (55,22.5), and the default upper right corner is (78,45), expressed as percentages of the current window.

If any of xlo, ylo, xhi, and yhi are not entered, the graphics cursor will be used to choose the missing coordinates.

Units

The frame box coordinates may be expressed in three types of units, which are chosen by command qualifier. The default is \PERCENT . See Table 2.44 for a listing of the qualifiers and their interpretations.

For example, if the \PERCENT qualifier is used, then a location of (50, 50) represents the centre of the current window. If the \WORLD qualifier is used, the coordinates are in units of the

| qualifier | interpretation of the coordinates |
|-----------|--|
| \PERCENT | percentages of the current window, as chosen with the WINDOW command. |
| \GRAPH | graph units, that is, the units defined by the minimum and maximum values for the last graph drawn. If no graph has been drawn yet, the defaults are $-1 \le x \le 1$ and $-1 \le y \le 1$ |
| \WORLD | centimeters or inches, as chosen with the SET UNITS command |

Table 2.44: Types of units recognized by the LEGEND command

world coordinate system, the plotting units. The default world coordinate system units are centimeters. See the DEVICE command for tables showing the dependence of plotting units on the graphics hardcopy output device.

Transparency

Syntax LEGEND TRANSPARENCY ON

LEGEND TRANSPARENCY OFF

Default ON

When the legend entries are drawn originally, the heights of the strings will likely all be different if AUTOHEIGHT is ON, and whatever is behind the frame box will be visible in the frame box. That is, the legend will be transparent. If the REPLOT command is entered, all the strings will be redrawn at the same height, using the minimum height of all the legend entries, and, if TRANSPARENCY is OFF, the background of the frame box will be erased. The default is TRANSPARENCY ON.

Note: This background erasing only applies to the monitor screen and to bitmap graphics hardcopy output, such as HP LaserJet output.

Remember, the legend frame box background will only be erased when a REPLOT command is entered.

Moving or resizing the legend frame box

The frame box can be moved and/or resized. First enter the LEGEND FRAME command to specify a new frame box, and then enter the REPLOT command.

The legend title

```
Syntax LEGEND TITLE { 'title string' }
Default no title
```

The default is to have no legend title. If there is a legend title, it is drawn just above the top of the frame box. The height of the legend title will be the current value of TXTHIT, as specified with the SET command. Text formatting commands may be included in the title string.

The legend title can be turned off by entering the LEGEND TITLE command without a title string.

Status

```
Syntax LEGEND STATUS
```

Use the STATUS keyword to display, on the monitor screen, the current status of the legend attributes.

Example

The following script produces figure 2.18 on page 143.

```
! first define the elements of a array string variable
T[1]='sin(<FITALIC.3>x<FTSAN>)<FITALIC.3>e<fITALIC.3,^>x<FTSAN>/5'
T[2]='\langle FITALIC.3\rangle \times \langle FTSAN\rangle/2-5'
T[3]='(<FITALIC.3>x<FTSAN>/3.5)<^>2<_>+3<FITALIC.3>x<FTSAN>/3.5'
T[4]='cos(<FITALIC.3>x<FTSAN>)<FITALIC.3>e<FTSAN,^>-<FITALIC.3>x<FTSAN>/9'
X = [0:4*PI:.5]
                                 ! generate some data
LEGEND ON
                                 ! turn legend on
LEGEND FRAME 20 50 50 80
                                 ! define the legend frame
LEGEND FRAME ON
                                 ! draw the frame box outline
LEGEND TITLE '<FROMAN.SERIF>Test legend'
SET LINTYP 1
                                 ! draw the first data curve
SET PCHAR 1
                                 ! plotting symbol
LEGEND NSYMBOLS 1
                                 ! one plotting symbol in legend entry
GRAPH T[1] \times SIN(X)*EXP(X/5)
                                 ! T[1] is the legend string entry
SET LINTYP 10
                                 ! draw the second data curve
SET PCHAR 2
                                 ! plotting symbol
LEGEND NSYMBOLS 2
                                 ! 2 symbols in legend entry
CURVE T[2] X X/2-5
                                 ! T[2] is the legend string entry
                                 ! draw the third data curve
SET LINTYP 5
SET PCHAR 3
                                 ! plotting symbol
LEGEND NSYMBOLS 3
                                 ! 3 symbols in legend entry
CURVE T[3] X (X/3.5)^2+3*X/3.5 ! T[3] is the legend string entry
SET LINTYP 7
                                 ! draw the fourth data curve
SET PCHAR 13
                                 ! plotting symbol
LEGEND NSYMBOLS 4
                                 ! 4 symbols in legend entry
CURVE T[4] X COS(X)*EXP(-X/9)
                                 ! T[4] is the legend string entry
REPLOT
                                 ! replot data on common scale
```

LINE

| Syntax | LINE |
|------------|--|
| | <pre>LINE\XYOUT x y { pen_code { line_type { colour { thickness }}}}</pre> |
| | <pre>LINE x y { pen_code { line_type { colour { thickness }}}}</pre> |
| Qualifiers | \GRAPH, \PERCENT, \WORLD, \XYOUT |
| Defaults | \PERCENT, interactive drawing |

This command draws line segments. If no parameters are entered with the LINE command, the graphics crosshair is used to draw interactively. If the \XYOUT qualifier is used, the graphics crosshair is used to draw the line segments, and the parameters are assumed to be output vectors in which to store the line segment coordinates and other attributes. If the \XYOUT qualifier is not used, and vectors are entered as parameters, then the input vectors

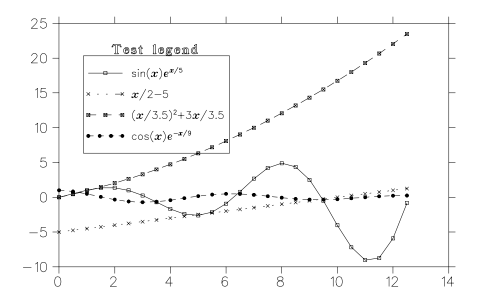


Figure 2.18: An example illustrating the graph LEGEND

are assumed to contain coordinates and other attributes for the line segments, and the graphics crosshair is not used.

Plotting units

If the \PERCENT qualifier is used, then the values of the coordinates will be interpreted as being in percentages of the current window, for example, (50,50) represents the centre of the current window. This is the default if no units qualifier is used.

If the \GRAPH qualifier is used, then the values of the coordinates will be interpreted as being in graph units, that is, the units defined by the minimum and maximum values for a graph. These units are taken from the last graph drawn, or, if no graph has been drawn yet, they are the default values: -1 <= x <= 1 and -1 <= y <= 1.

If the \WORLD qualifier is used, then the values of the coordinates will be interpreted as being in centimeters or inches, depending on the units type chosen with the SET UNITS command. The default world units are centimeters.

X Windows

When running under X Windows, mouse button two toggles the continuous display of the graphics cursor location. The PHYSICA keyword CUNITS is the units type for these numbers. If CUNITS = WORLD, the numbers depend on the current units type, either CM or IN, as chosen with SET UNITS. If CUNITS = GRAPH, the numbers displayed depend on the current graph axis

scales. If CUNITS = PERCENT, the numbers depend on the current window.

Non-interactive drawing

```
Syntax LINE x y { pen_code { line_type { colour { thickness }}}} 
Defaults pen_code = [3; 2; 2; ...], line_type = [1; 1; 1; ...], colour = [1; 1; 1; ...], thickness = [1; 1; 1; ...]
```

Vectors x and y should contain the x and y coordinates for the line segments' end points. Optional vector pen_code should contain the pen codes, either 2 (connect) or 3 (disconnect) for each point. Optional vector line_type should contain the line type codes (between 1 and 10) for the line segments. The line types are described in the SET LINE command section. Optional vector colour should contain the colour codes (between 1 and 8). Optional vector thickness should contain the the line thickness for bitmap and PostScript hardcopy output.

| key | action |
|-----|---|
| Q | quit the LINE command |
| M | display the command menu |
| / | clear the alpha-numeric terminal screen This has no affect on the graphics |
| Ū | start a line segment A mark will be drawn at the current location, but is not entered into the plot file, or into an open EDGR file, it is just there for reference |
| D | end a line segment A line segment is drawn from the last location that was chosen with the $\tt U$ or $\tt D$ key to the current location. This line segment will be entered into the plot file, and into an open EDGR file |
| T | try a line segment A line segment is drawn from the last location that was chosen with the U or D key to the current location. This line segment will <i>not</i> be entered into the plot file, or into an open EDGR file. If the line segment is acceptable, do not move the crosshair, and type the D key. If the line segment is not acceptable, simply move to another location and try again; this line segment will then be erased. |
| L | change the line type The current line type code is displayed and a new code can be entered. No carriage return is necessary after the new code is entered. The line type should be between 1 and 10, to choose number 10, enter an A. The line types are described in the SET LINE command section. The default line type is 1. |
| С | change the colour The current code is displayed and a new code can be entered. No carriage return is necessary after the new code is entered. 1=white, 2=red, 3=green, 4=blue, 5=yellow, 6=cyan, 7=magenta, 8=white |
| N | change the units The current units code is displayed and a new code can be entered. No carriage return is necessary after the new code is entered. 1 - percentages of the world coordinate system 2 - world coordinate system units (centimeters or inches) 3 - graph units 4 - percentages of the current window |
| Х | toggle the display of the x and y coordinates at the bottom of the terminal screen. This is for user reference only. By default, the x and y coordinates are not displayed. |

Table 2.45: The LINE command interactive menu

LIST

```
Syntax LIST x1 { x2 ... x5 } { n1 { n2 { n3 }}}
LIST\MATRIX matrix
LIST\MATRIX\FORMAT matrix (format)

Qualifiers \PAGE, \MATRIX, \FORMAT, \COUNTER

Defaults vector(s) expected, n1 = 1, n3 = 1, not paged, format = 1PD13.4, \COUNTER

Examples LIST X Y Z
LIST X Y Z 1 10 2
LIST\PAGE X[10:#] Y Z
LIST\PAGE X[10:#] Y Z
LIST\MATRIX\FORMAT M[2:11,5:20:2] (6(F10.3,2X))
```

By default, the LIST command displays, on the monitor screen, a listing of the specified vectors, xI[n1:n2:n3], with a counter displayed on the left under the heading #. n1 defaults to 1 and n3 defaults to 1 if not entered. If n2 is not entered, all of the elements of each vector are listed. If the counter is not desired, use the $\-COUNTER$ qualifier.

Paging the output

The \PAGE qualifier only applies to listings of vectors.

If the \PAGE qualifier is used, the listing will be paged, that is, after every twenty lines have been displayed, you will be asked if you want to continue or quit. Type the $\mathbb Q$ key to quit. Type any other key to continue the listing. No carriage return is necessary.

Listing a matrix

```
Syntax LIST\MATRIX matrix
LIST\MATRIX\FORMAT matrix (format)

Qualifiers \MATRIX, \FORMAT

Defaults format = 1PD13.4
```

The LIST\MATRIX command displays a matrix on the monitor screen. The format defaults to 1PD13.4. Use the \FORMAT qualifier to indicate that a user specified format has been entered. The format must be enclosed in parentheses, (and). Any standard FORTRAN format is valid, but only REAL variables can be displayed. Do not use INTEGER, LOGICAL, or CHARACTER formats.

LOAD

Syntax LOAD filename{,libraries}

Examples LOAD MYFILE

LOAD MYFILE, LIB1/LIB, LIB2/LIB

The LOAD command is only available under VAX/VMS.

The LOAD command dynamically loads and links an object module of a user written subroutine or a user written REAL*8 function. The filename is the compiled object module.

If a function is loaded, it can only be used in expressions, via the name USERN. If a subroutine is loaded, it can only be used with the CALL command, with no specified name.

If your subroutine or function requires other routines, either object modules files or object module libraries, just include the other file names and/or library names in the LOAD command. For example:

LOAD somename, OTHER1.OBJ, DISK: [DIR] LIBRARY/LIB

Restrictions

- The complete linked module has a size limit of 256,000 bytes (500 blocks).
- Only one object module can be loaded at any time. A subsequent LOAD command will simply replace the old module with the new.

Arguments

A user written function can have up to 20 REAL*8 arguments, all of which must be scalars.

A user written subroutine can have up to 15 arguments, which can be a mixture of contstants, scalars, vectors, matrices, quote strings, and string variables. The numeric arguments must be REAL*8.

The user *must* call the loaded subroutine or function with the correct number and type of arguments, else an addressing exception will result, and the program may be corrupted.

Subroutines

The name of a subroutine to be loaded dynamically via the LOAD command is irrelevant and can be anything the user desires. A user written subroutine *must* have the following form:

```
SUBROUTINE subname(IATYPE,ICODE,IUPDATE,IER,arg1,arg2,...)
INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
```

Other than the required arguments, IATYPE, ICODE, IUPDATE, and IER, there may be from 1 to 15 arguments in the subroutine argument list. The user is responsible for insuring that the correct number and type of arguments are used when actually employed with the CALL command. The parameters used in the CALL command, argI, which are passed as arguments to the subroutine, may be constants, scalars, vectors, matrices, quote strings, or string variables. The number of arguments and the type of argument *must* agree with the actual subroutine.

All of the numeric aguments, except for the required integer arguments IATYPE, ICODE, IUPDATE, and IER, must be REAL*8. A string argument is passed as a LOGICAL*1 array.

Note: The integer arguments IATYPE, ICODE, IUPDATE, and IER should *not* be mentioned as parameters with the CALL command.

See the file: PHYSICA\$DIR:PHYSICA_USER_FUNCTIONS.FOR for some subroutine examples.

IATYPE

IATYPE is an INTEGER*4 array, length 15, that indicates the type of each of the subroutine arguments argI. See Table 2.4 on page 15.

ICODE

ICODE is an INTEGER*4 array, dimensioned 3 by 15, that indicates the dimension of each of the subroutine arguments argI. Never extend variables beyond their original size as passed to the subroutine. If a variable is shortened inside the subroutine, the subroutine must update the new dimensions in the ICODE array, so that PHYSICA can reduce the variable dimensions appropriately. See Table 2.5 on page 16.

The ICODE array will be filled by PHYSICA with the current dimensions of the arguments, so the user written subroutine can check and, if necessary, update the dimensions of any of the subroutine arguments.

Never extend vectors, matrices, or string variables beyond their original sizes as passed to the user written subroutine. If a variable's size is shortened inside the subroutine, then the subroutine must update the ICODE array so that these variable dimensions can be reduced internally by PHYSICA upon return from the subroutine.

IUPDATE

IUPDATE is an INTEGER*4 array, length 15, that the user routine sets to indicate to PHYSICA whether one of the argI arguments has been modified inside that subroutine.

The default value for IUPDATE(i) is 0. Set IUPDATE(i) to 1 to indicate that the i^{th} argument, argI, has been modified. Never extend variables beyond their original size as passed to the subroutine. If a variable is shortened inside the subroutine, the subroutine must update the new dimensions in the ICODE array, so that PHYSICA can reduce the variable dimensions appropriately.

IER

IER is an INTEGER*4 variable that defaults to the value 0. Your routine can set IER to indicate to PHYSICA that an error has occured in the routine. Arithmetic errors, such as division by zero, over/underflow, will be asynchronously trapped. If other error tests are to be done inside the subroutine, the user flags the error by setting IER = -1 before the RETURN. If the CALL command was executed from within a script, this error flag causes PHYSICA to abort that script and control is passed back to the keyboard.

Numeric arguments

All the numeric arguments of your subroutine, except for the integer arguments IATYPE, ICODE, IUPDATE, and IER, must be REAL*8. A string argument is passed as a LOGICAL*1 array. Dimension numeric array arguments with length 1, for example:

```
REAL*8 X(1), Y(1), Z(1)
```

String arguments

All the string arguments of your subroutine must be LOGICAL*1, and should be dimensioned 1, for example:

```
LOGICAL*1 LFILE(1)
```

You can convert this to a string, say, CHARACTER*80 CFILE, using the following method:

```
LENF = ICODE(1,i)
DO I = 1, LENF
   CFILE(I:I) = CHAR(LFILE(I))
END DO
```

where LFILE is the i^{th} argument.

Accessing matrix data

If a matrix is passed as an argument to a user written subroutine, the elements of the matrix can only be accessed using a calculated index. To access element m[i,j] of the matrix m, use m[i+(j-1)*nrows] for $i=1, \ldots, nrows$ and $j=1, \ldots, ncols$.

Subroutine example

Consider the following subroutine, in file LOAD1.FOR, for calculating the cumulative product of the elements of a vector. Note the use of the mandatory integer arrays for checking input parameter types and sizes, for indicating which variables have changed, and for flagging errors.

```
SUBROUTINE LOAD1(IATYPE, ICODE, IUPDATE, IER, X, Y)
   INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
  REAL*8
             X(1), Y(1)
This subroutine requires two vector arguments, with the length of Y
greater than or equal to the length of X, and calculates the cumulative
product of X and stores the results in Y.
First check for input errors
   IF( IATYPE(3) .NE. -99 )THEN
     WRITE(*,*)' ERROR: too many arguments for loaded subroutine'
     IER = -1
     RETURN
  END IF
   IF( IATYPE(2) .EQ. -99 )THEN
     WRITE(*,*)' ERROR: not enough arguments for loaded subroutine'
     IER = -1
     RETURN
  END IF
   IF( IATYPE(1) .EQ. -99 )THEN
     WRITE(*,*)' ERROR: not enough arguments for loaded subroutine'
     IER = -1
     RETURN
  END IF
   IF( IATYPE(2) .NE. 1 )THEN
     WRITE(*,*)' ERROR: second argument is not a vector'
     IER = -1
     RETURN
  END IF
   IF( IATYPE(1) .NE. 1 )THEN
     WRITE(*,*)' ERROR: first argument is not a vector'
     IER = -1
     RETURN
  END IF
   IF( ICODE(1,2) .LT. ICODE(1,1) )THEN
     WRITE(*,*)' ERROR: second vector length < first vector length'</pre>
     IER = -1
     RETURN
   END IF
   IUPDATE(2) = 1
                     ! indicates the second argument is changed
  Y(1) = X(1)
  DO I = 2, ICODE(1,1)
     Y(I) = Y(I-1)*X(I)
  END DO
  RETURN
   END
```

To compile this source code, from within PHYSICA, creating an object module, load it, and then to make use of it, enter:

```
$FORTRAN LOAD1
LOAD LOAD1
X=[1:20]
VECTOR XPROD 20
CALL X XPROD
SCALAR\DUMMY J
LIST X XPROD RPROD(X[J],J,1:20) ! RPROD function is the same as LOAD1
```

Functions

A user written *function* should have the following form:

```
REAL*8 FUNCTION funcname(ARG1,ARG2,...,ARGM)
REAL*8 ARG2,ARG2,...,ARGM
```

The name of the function, funcname, is irrelevant, since, when the function is used in an expression, the name that is used *must* be USERN.

Note: There is no ICODE array for a function.

There may be from 1 to 20 arguments. All arguments *must* be REAL*8 scalars.

All arguments of the function are scalars. When the function is used in an expression, you can use vectors or matrices as arguments since the expression evaluator will process the function one element at a time.

Functions are *not* used with the CALL command. A user written function can be used wherever a function can be used in an expression.

Arithmetic errors, such as division by zero, overflow, or underflow, will be asynchronously trapped.

Note: There are also eight user written functions which are loaded at run time via a shareable image. These eight functions must be called USER1, USER2, ..., USER8.

Function example

Suppose you want to use a function, ANYNAME, which has three arguments, x_1 , x_2 , and x_3 , and returns the sum of x_1 and x_2 when x_3 is within $\frac{1}{2}$ of $\sqrt{x_1^2 + x_2^2}$, else it returns 0.

```
REAL*8 FUNCTION ANYNAME(X1,X2,X3)

IMPLICIT REAL*8 (A-H,O-Z)

ANYNAME = 0.0D0

IF( ABS( SQRT(X1**2+X2**2)-X3 ) .LT. 0.5D0 )ANYNAME = X1+X2

RETURN

END
```

To compile this source code, from within PHYSICA, creating an object module, load it, and then to make use of it, enter:

```
$FORTRAN ANYNAME
LOAD ANYNAME
=USERN(A,B,C)
```

A simple way to test the above function:

```
X=RAN([1:100])
Y=RAN([1:100])
Z=RAN([1:100])
U1=USERN(X,Y,Z)
U2=(X+Y)*(ABS(SQRT(X^2+Y^2)-Z)<.5)</pre>
```

and compare U1 with U2.

MAP

```
Syntax MAP\FIOWA global
MAP\FIOWABIG global
MAP\YBOS
MAP\HBOOK name
```

The MAP command is used to map onto shared memory or a global section. The FIOWA, FIOWABIG, and NOVA options are only available under VAX/VMS. The YBOS option is only available under DEC AXP OSF/1. The HBOOK option is available for global sections under VAX/VMS and for shared memory under all UNIX systems supported in the CERN libraries.

HBOOK

```
Syntax MAP\HBOOK name
```

A "snapshot" is taken of the data in the named global section or shared memory. To access the most current data, re-enter the MAP command. The variables created are the same as with the RESTORE\HBOOK command. Use the GLOBALS command under VAX/VMS to see the global section names to which you have access. Refer to the RESTORE\HBOOK command for information on what variables are created and their definitions.

FIOWA

Syntax MAP\FIOWA globalname MAP\FIOWABIG globalname

The MAP\FIOWA and MAP\FIOWABIG commands only work under VAX/VMS.

The MAP command is used to map onto a global section. A "snapshot" is taken of the data in the global section. To access the most current data, re-enter the MAP command. The variables created are the same as with the RESTORE\FIOWA command. Use the GLOBALS command to see the global section names to which you have access. Refer to the RESTORE command for information on what variables are created and their definitions.

Use the \FIOWABIG qualifier to access data sets made with the "big" version of FIOWA.

YBOS

Syntax MAP\YBOS

The MAP\YBOS command only works under Digital Unix.

The MAP command is used to map onto shared memory. A "snapshot" is taken of the data in the shared memory. To access the most current data, re-enter the MAP\YBOS command. The variables created are the same as with the RESTORE\YBOS and with the RESTORE\YBOS\DOTPLOT commands. Refer to the RESTORE command for information on what variables are created and their definitions.

MATRIX

```
Syntax MATRIX m1 { m2 ... } nrow ncol

Examples MATRIX M 10 20

MATRIX M1 M2 M3 10 20
```

The MATRIX command creates new matrices or changes the dimensions of existing matrices. Each mI will be a matrix with nrow rows and ncol columns.

If mI exists and is a matrix, then it will be either trimmed down to the specified dimensions or zero filled to expand it.

If mI exists but is not a matrix, it will be destroyed first and then re-created as a zero filled matrix with nrow rows and ncol columns. If mI does not exist, it will be created as a zero filled matrix with nrow rows and ncol columns.

MONITOR

Syntax MONITOR { keyword }

The MONITOR command is used to select a graphics output monitoring device. Table 2.46 lists the currently recognized graphics display device types.

Selecting a monitor type necessitates clearing the graphics.

The MONITOR command should be entered before opening an EDGR file.

| device | keyword |
|-------------------|---------|
| Digital VT640 | VT640 |
| Digital VT241 | VT241 |
| Citoh CIT-467 | CIT467 |
| Tektronix 4010/12 | TK4010 |
| Tektronix 4107 | TK4107 |
| Plessey PT-100G | PT100G |
| Seiko GR-1105 | GR1105 |
| X Window system | Х |
| Generic terminal | GENERIC |

Table 2.46: Monitor device types and corresponding keywords

Disabling/enabling graphics monitor output

Use the MONITOR OFF command to disable graphics output to the monitor. This can be useful when executing command files which the user is confident of and does not need to monitor. The hardcopy output is not affected, and is much quicker to obtain. To make the previous monitor type again available, enter the MONITOR ON command. When ON is used, the monitor type that was last in use will again be available for monitoring graphics.

The generic terminal driver

The generic terminal driver is a driver that is controlled by a data file that can be created and modified by the user. The file contains lists of escape sequences for performing the

various terminal functions. Creating such a file for a given terminal may allow it to be used with PHYSICA even though it is not explicitly supported.

This generic terminal data file name is passed to the program by a logical name, on VMS systems, or by an environment variable, on UNIX systems. This name must be assigned before you run the PHYSICA program.

VMS: \$ DEFINE TRIUMF\$GENTERM disk:[directory]filename

UNIX: % setenv TRIUMF_GENTERM <directory>filename

NEWS

Syntax NEWS

Use the NEWS command to display the latest information about changes in the program and new features pertaining to the PHYSICA program.

ORIENTATION

Syntax ORIENTATION keyword

Default initial orientation = LANDSCAPE

The ORIENTATION command sets the graphics orientation. This requires that the graphics be cleared, which is done automatically.

Note: The ORIENTATION command should be entered before opening an EDGR file with the EDGR OPEN command.

There are two basic graphics orientations available.

| keyword | result |
|-----------|---|
| LANDSCAPE | the world coordinate system large dimension is horizontal |
| PORTRAIT | the world coordinate system large dimension is vertical |

The initial orientation is landscape.

Landscape orientation means, for example, that text, with a text angle of zero, will be drawn across the wide dimension of the plotter paper, or that a graph will appear with the x-axis drawn across the wide dimension of the plotter paper.

Portrait orientation means, for example, that text, with a text angle of zero, will be drawn across the narrow dimension of the plotter paper, or that a graph will appear with the x-axis

drawn across the narrow dimension of the plotter paper.

The world coordinate system

The world coordinate system plotting units depend four things:

- the graphics orientation;
- the graphics hardcopy device type;
- the graphics hardcopy device size;
- the type of graphics units.

Refer to the DEVICE command, page 47, for tables showing the world coordinate system plotting units for the various devices.

The type of graphics units are chosen with the SET UNITS command.

PEAK

| Syntax | PEAK xout yout |
|------------|--|
| | PEAK\PNUM xout yout num |
| Qualifiers | \PNUM |
| Defaults | num = number of last drawn data curve, \NOPNUM |

The PEAK command brings up the graphics cursor, and interacts with the user through a keystroke menu, to find the minima or maxima of the current data curve drawn on the screen. The coordinates of the peaks so found are stored in the specified variables <code>xout</code> and <code>yout</code>. By default, the data curve referenced is the last one drawn. When the graphics cursor is brought up, the program is waiting for input of a one or two character keycode.

Choosing the data curve

```
Syntax PEAK\PNUM xout yout num
```

Any one of the data curves currently on the screen may be specified explicitly by using the \PNUM qualifier and including the scalar num. This scalar specifies which data curve, in the order they were drawn, is to be selected.

X Windows

When running under X Windows, mouse button two toggles the continuous display of the graphics cursor location. The PHYSICA keyword CUNITS is the units type for these numbers. If CUNITS = WORLD, the numbers depend on the current units type, either CM or IN, as chosen

with SET UNITS. If CUNITS = GRAPH, the numbers displayed depend on the current graph axis scales. If CUNITS = PERCENT, the numbers depend on the current window.

Code keys

Remember that this command assumes that a graph is present on the screen and none of the plot characteristics have been changed after the graph was done. After entering the PEAK command, the graphics cursor will appear. Typing a special code key results in a specific action. See Table 2.47 on page 158.

| key | action |
|-------|---|
| < | moves the graphics cursor to the nearest peak (or dip) to the left |
| > | moves the graphics cursor to the nearest peak (or dip) to the right |
| Pn | defines criterion for peak finding to be n points monotonically increasing, followed by n points monotonically decreasing. (default n= 1) <i>Note</i> : works only with > when enabled, n= 0 disables this function |
| Fn | fits least-squares parabola to 2*n+1 nearby points (n= $1,\ldots,9$) and moves the graphics cursor to the fitted peak location, e.g., F3 would peak fit the surrounding 7 pts |
| Cn | use the \mathtt{n}^{th} curve on the screen for subsequent codes |
| + | search for peaks (maxima) |
| _ | search for dips (minima). This is the default. |
| A | amplitude required for peak (or dip). The user is prompted to enter a limiting amplitude by returning two cursor y -coordinates (abort with Z) |
| Y | set the y tolerance parameter. Move the cursor to the desired y value and enter any character (abort with Z) |
| N | disable the y tolerance flag |
| Z | aborts entry of cursor coordinates for A,Y codes |
| D | display the last peak position in graph coordinates |
| R | record and save the last peak position |
| S | save the last peak position and mark with a symbol |
| / | alphanumeric clear |
| Q | quit |
| All o | ther characters ignored |

Table 2.47: Key codes for the PEAK command

PICK

```
Syntax PICK { xout yout { y1 { n1 } y2 { n2 } ... }}
PICK\NPTS num { xout yout { y1 { n1 } y2 { n2 } ... }}
Qualifiers \NPTS, \POLYGON, \MATRIX, \COUNTS, \MIN, \MAX, \DISPLAY

Defaults nI = I, \NONPTS, \NOPOLYGON, \NOMATRIX, \NOCOUNTS, \NOMIN, \NOMAX, \DISPLAY
```

The default action for the PICK command is to pick points off a graph which is currently displayed on the monitor screen, and to optionally save the values in output vectors, xout and yout. It is assumed that a graph is present on the screen and none of the plot characteristics have been changed after the plot was done. The graphics cursor is used, and various actions depend on which key is typed from the keyboard. If xout and yout are not entered, then no points will be saved.

X Windows

When running under X Windows, mouse button two toggles the continuous display of the graphics cursor location. The PHYSICA keyword CUNITS is the units type for these numbers. If CUNITS = WORLD, the numbers depend on the current units type, either CM or IN, as chosen with SET UNITS. If CUNITS = GRAPH, the numbers displayed depend on the current graph axis scales. If CUNITS = PERCENT, the numbers depend on the current window.

Specifying the number of points

```
Syntax PICK\NPTS num { xout yout { y1 { n1 } y2 { n2 } ... }}
```

When the \NPTS qualifier is used, a scalar, num, is expected which will be the maximum number of points that can be recorded with that command. When you have recorded num points, the command automatically stops.

Code keys

Remember that this command assumes that a graph is present on the screen and none of the plot characteristics have been changed after the graph was done. After entering the PICK command, the graphics cursor will appear. Typing a special code key results in a specific action. See Table 2.48.

If the \-DISPLAY qualifier is used, it turns off the message display for the default action, which is picking points off a graph. This was added to clean up the terminal window for those using the PICK command who do not desire to see the (x,y) locations or the menu. These can be turned on inside the PICK command by typing the 0 key.

| key | action |
|-----|--|
| | |
| D | Digitize the current crosshair position and display the coordinates corresponding to that location |
| R | same as D, but also record the coordinates, in graph units, of that location in the optional vectors xout and yout, record the interpolated value of curve nI in vector yI. The number of recorded points is displayed |
| М | same as R, but place a marker at the recorded location |
| C | same as ${\tt M}$, but also connect that point to the last recorded point with a line segment |
| 0 | turn on message display |
| Q | quit picking points |

Table 2.48: Key codes for the PICK command

Automatic digitizing of previously graphed data

It is possible to automatically digitize points off of previously drawn data curves by making use of the optional parameters yI (vector name) and nI (number). Assuming that the total number of curves that have been plotted on the currently displayed graph is at least nI, then the coordinate corresponding to the horizontal (independent) location of the crosshair will be interpolated on the nI^{th} curve and saved in the vector yI. Linear interpolation is used, and extrapolation is allowed. If the number nI is not present following the vector yI, then nI defaults to I.

Examples

Suppose you have entered the following commands to draw three data curves:

```
GRAPH\NOAXES X2 Y2
GRAPH\NOAXES X3 Y3
```

To pick points off the three data sets, you could enter: PICK XP YP YP1 YP2 YP3 Move the crosshair horizontally across the screen and type the R key at each location where you desire a value. The vector YP1 will the the interpolated values from curve (X1,Y1), vector YP2 will contain those from (X2,Y2), and vector YP3 will contain those from (X3,Y3). The horizontal (independent) coordinates will be contained in the vector XP.

Suppose you have drawn the same three data sets as above, but you only want to pick points off of the third curve drawn, that is, (X3,Y3). Enter: PICK XP YP YP3 3 Move the crosshairs horizontally across the screen and type the R key at the locations where you desire values off the third curve. The vector YP3 will contain the interpolated values from curve (X3,Y3), while the horizontal (independent) coordinates will be contained in vector XP.

Choosing the vertices of a polygon

```
Syntax PICK\POLYGON xp yp
```

The \POLYGON qualifier indicates that you are picking vertex points for a polygon. The only difference between \POLYGON and the default, \NOPOLYGON, is that a final point is added when you quit picking points. This final point will be the same as the first point picked, thus closing the polygon.

Matrices

```
Syntax PICK\MATRIX nrow ncol mxout myout
PICK\MATRIX\MIN { xin yin } m mxout myout
PICK\MATRIX\MAX { xin yin } m mxout myout

Defaults xin = [1;2;3;...], yin = [1;2;3;...]
```

The PICK\MATRIX command allows you to pick points off the screen using the graphics cursor. The x coordinates of the points are placed in matrix \mathtt{mxout} , while the y coordinates are placed in matrix \mathtt{myout} . The user must specify the size of these output matrices in \mathtt{nrow} , the number of rows, and \mathtt{ncol} , the number of columns. Both \mathtt{mxout} and \mathtt{myout} will be created with the same dimensions.

This feature was included to be used in conjunction with the BIN2D\MATRIX command, page 9.

Finding regional minima for a matrix

The PICK\MATRIX\MIN command allows you to choose circular regions on the screen, by choosing centres and radii. The minimum value of the matrix, m, contained within this circle will be found. The x coordinate of these minima will be output into matrix mxout, the y coordinates will be output into matrix myout.

Finding regional maxima for a matrix

The PICK\MATRIX\MAX command allows the user to choose circular regions on the screen, by choosing centres and radii. The maximum value of the matrix, m, contained within this

circle will be found. The x coordinate of these maxima will be output into matrix mxout, the y coordinates will be output into matrix myout.

Determining regional counts for data sets

```
Syntax PICK\COUNTS { xin } ydata counts PICK\COUNTS\MATRIX { xin yin } m counts Defaults xin = [1;2;3;...], yin = [1;2;3;...]
```

The PICK\COUNTS command allows you to pick pairs of points off the screen. The number of counts of the vector ydata inside that region will be displayed on the screen and stored in the output vector, counts. The input vector, xin, should contain the x coordinates for the data ydata. xin defaults to [1;2;3;...] if not entered.

Matrices

The PICK\COUNTS\MATRIX command allows you to pick rectangular regions off the screen. The number of counts of the matrix inside each such region will be displayed on the screen and stored in the output vector, counts. The input vectors, xin and yin, should contain the x and y coordinates for the matrix, where xin corresponds to the columns and yin corresponds to the rows, that is, m[i,j] has coordinates (xin[j],yin[i]). xin and yin both default to [1;2;3;...] if not entered.

PIEGRAPH

```
Syntax PIEGRAPH w e f c o r { cx cy { a }}

Qualifiers \GRAPH, \PERCENT, \WORLD

Defaults \PERCENT, a = 0
```

The PIEGRAPH command draws piecharts. The first five parameters are vectors which define each wedge of the pie. See Table 2.49.

Coordinates

The parameters r, cx, cy, and a should be literal constants or scalars. The radius of the pie chart is r. The parameters cx and cy represent the x and y location of the centre of the pie chart. The starting angle, in degrees, for the first wedge can be optionally specified with the parameter a, which defaults to zero.

If either cx or cy is not entered, the graphics cursor will be used to choose that coordinate. If both cx and cy are present, the graphics cursor will not be used.

```
 \begin{array}{|c|c|c|c|c|} \hline w & wedge \ values, \ in \ whatever \ units \ are \ applicable \\ the \ sum \ of \ all \ the \ wedge \ values \ is \ 100\% \ of \ the \ pie \ chart \\ \hline e \ & explode \ values, \ as \ a \ percent \ of \ the \ radius \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ radius \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ radius \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ radius \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ radius \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ no \ filling \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ no \ filling \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ no \ filling \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ no \ filling \\ f \ & explode \ values, \ as \ a \ percent \ of \ the \ no \ filling \\ f \ & explode \ outline \ of \ the \ pattern \\ f \ & explode \ outline \\ f \ &
```

Table 2.49: The pie wedge defining characteristics

Units

The pie chart coordinates may be expressed in three types of units, which are chosen by command qualifier. The default is \PERCENT. See Table 2.50 for a listing of the qualifiers and their interpretations.

| qualifier | interpretation of the coordinates | |
|-----------|--|--|
| \PERCENT | percentages of the current window, as chosen with the WINDOW command. | |
| \GRAPH | graph units, that is, the units defined by the minimum and maximum values for the last graph drawn. If no graph has been drawn yet, the defaults are $-1 \le x \le 1$ and $-1 \le y \le 1$ | |
| \WORLD | centimeters or inches, as chosen with the SET UNITS command | |

Table 2.50: Types of units recognized by the PIEGRAPH command

For example, if the \PERCENT qualifier is used, then a location of (50,50) represents the centre of the current window. If the \WORLD qualifier is used, the coordinates are in units of the world coordinate system, the plotting units. The default world coordinate system units are centimeters. See the DEVICE command for tables showing the dependence of plotting units on the graphics hardcopy output device.

Figure 2.19 illustrates how each individual pie wedge is drawn.

Pie wedge filling

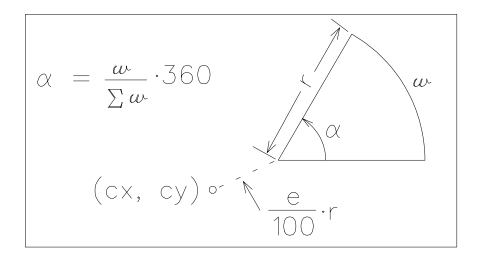


Figure 2.19: Pie chart wedge definition

Each wedge of the pie can be filled with either a grey scale dot fill pattern or a hatch pattern.

Hatch patterns

A hatch pattern is composed of an angle and one to ten spacings. These spacings are simply cycled through as the region is being filled, that is, a line is drawn inside the region at the specified angle, then a parallel line is drawn at the first spacing, then another parallel line is drawn at the second spacing, and so on for the number of spacings in that pattern. This process is repeated until the region is filled. The hatch patterns can be redefined with the SET HATCH command and displayed with the DISPLAY FILL command. There are ten hatch patterns available.

Dot fill patterns

A dot pattern is of the form: uv, where the digit u is the increment number of dots to light up horizontally, $1 \le u \le 9$, and the digit v is the increment number of dots to light up vertically, $1 \le v \le 9$. For example, a dot pattern of 34 means to light up every third dot horizontally and every fourth dot vertically. If uv is negative, then the dots are erased instead of turned on.

PostScript output

For PostScript output, set the PostRes keyword to the appropriate resolution for your hard-copy device, using the Set command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark inkjet printer has a resolution of 360 dpi. A "good" picture can be obtained with Postres = 180 and Linthk = 2.

Example

The following script produces Figure 2.20.

```
PIE_VALUES=[12.1;18.2;24.2;15.2;30.3]

EXPLODE=[10;10;0;25;0]

FILL=[8;33;44;7;22]

WCOLOUR=[1;2;3;4;5]

OCOLOUR=[5;4;3;2;1]

PIEGRAPH PIE_VALUES EXPLODE FILL WCOLOUR OCOLOUR 40 50 50
```

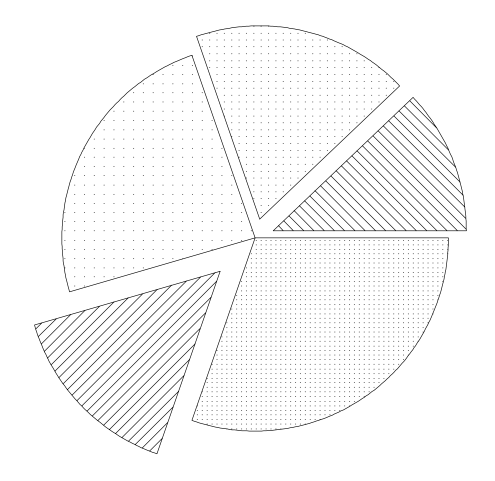


Figure 2.20: An example of a pie chart

PLOTTEXT

Syntax PLOTTEXT filename

The PLOTTEXT command is used to draw lines of formatted text that are read from a file. Text formatting commands can be interspersed in the lines of text. The text formatting

commands are summarized in Table 2.51.

Every text formatting command *must* be bracketed by the command delimiters, except for comment lines, which do not require the trailing command delimiter.

There is also a set of special reserved character names, see Figure 2.6 on page 61. These special names must be enclosed by the command delimiters, for example, <alpha><beta>. See the DISPLAY command on page 58 for information on how to display these special reserved names.

Environment variables in file names

For UNIX users, it is now possible to use an environment variable in a file name, if the environment variable is preceded by a \$. For example,

```
setenv FILE mytext
physica
plottext $FILE
```

The environment variable can be just the first part of the filename, for example,

```
setenv FILE my
physica
plottext $FILEtext
```

Comments

Command form: <! comment...

To insert comment lines in a formatted text file, that is, lines that will not be drawn on the plot, include <! in positions one and two of a line. The rest of the line will be considered to be a comment and will be ignored. This is useful for documenting formatted text files. For example, this input:

produces this output:

This is an example of a formatted text file with comments

Command delimiters

```
<!
                            comment line
               <Qx>,<Qxy> set command delimiters
                        <-> line continuation
                        < >
                             insert blank line
        <Hxx.x>, <Hxx.x%>
                             text height
        <Sxx.x>, <Sxx.x%>
                             line spacing
        <Mxx.x>, <Mxx.x%>
                             left margin
         <B>, <Bn>, <Bn:m>
                             grey scale or hatch fill
                      <Cn>
                             colour
              <Ffontname>
                             text font
<JC>, <JCxx.x>, <JCxx.x%>
                             centre
<JL>, <JLxx.x>, <JLxx.x%>
                             left justify
<JR>, <JRxx.x>, <JRxx.x%>
                             right justify
        <Zxx.x>, <Zxx.x%>
                             insert horizontal space
                        <_>
                             sub-script mode
                             super-script mode
                      <em>
                             italics mode
                        <X>
                             hexadecimal mode
                      <\b>
                             macron (bar) under previous character
                      <\d>
                             dot under previous character
                      <\^>
                             circumflex over previous character
                      <\ \>
                             acute over previous character
                      <\ '>
                             grave over previous character
                      <">
                             umlaut over previous character
                      <\^>
                             tilde over previous character
                      <\=>
                             macron (bar) over previous character
                      <\.>
                             dot over previous character
                      <\u>
                             breve over previous character
                      <\v>
                             v over previous character
                      <\H>
                             dieresis (double quote) over previous char-
                             acter
```

Table 2.51: PLOTTEXT command text formatting commands

Command form: <Qx>, <Qxy> Default delimiters: < and >

To set the leading command delimiter to x and the trailing command delimiter to y, include $\langle Qxy \rangle$ in the formatted text. If a single character, x, is to be the leading *and* the trailing command delimiter, simply include $\langle Qx \rangle$ in the formatted text. This is useful when the current command delimiters are required as characters to be drawn.

Within a file, the command delimiters will remain as set until changed with another <Qx> or <Qxy> command, but the command delimiters always default to < and > when the PLOTTEXT command is entered.

For example, this input:

```
<FTSAN><H.6><S1.2>Change the delimiters from <Q\> < and > to \Q<>\ \ and back to <Q\> < >\Q<>\
```

produces this output:

```
Change the delimiters from < and > to \setminus and back to < >
```

Continuation Lines

Command form: <->

To continue a line of formatted text onto the next line of a file, include <-> at the end of the line. The next line of text will be drawn at the end of the line with the continuation. Blanks at the end of the line with the continuation command, but before the <-> will be included, as well as blanks at the beginning of the next line.

This is useful when you have so many text formatting commands in a line that the line becomes very long. The maximum input line length is 255 characters, so it is necessary to continue the line onto two or more input lines.

```
<HO.5><S1>TH<FROMAN.FUTURA>IS <FROMAN.FASHON>is <->
<FROMAN.SERIF>a<FROMAN.SWISSL>n example <->
of a file
containing <->
continued lines
```

produces this output:

THIS is an example of a file containing continued lines

Inserting a blank line

Command form: < >

To insert a blank line in the plotted text use < >. Blank lines that are encountered in a formatted text file are simply ignored.

For example, this input:

<H.6><S1.2>The following blank line is ignored.

There will be a blank line after this.<>
This line is preceded by a blank line.

produces this output:

The following blank line is ignored. There will be a blank line after this.

This line is preceded by a blank line.

Character height

Command form: <Hxx.x>, <Hxx.x%>

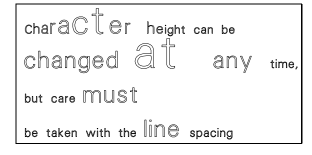
Default height: the current value of TXTHIT

To set the character height to xx.x units, include $\langle Hxx.x \rangle$ in the formatted text. This applies to subsequent text. The units, either centimeters or inches, are defined by the SET UNITS command. To set the height as a percentage of the height of the current window, that is, YUWIND - YLWIND, use $\langle Hxx.x\% \rangle$.

Within a file, the text height will remain as set until changed with another <hxx.x> or <hxx.x%> command, but the text height always defaults to the current value of TXTHIT when the PLOTTEXT command is entered.

```
<FROMAN.SWISSL><H0.5><S2>C<H.6>h<H.7>a<H.9>r<->
<H1.1>a<H1.4>c<H1.7>t<H1.4><->
<H1.1>e<H.9>r <H.7>h<H.6>e<H.5>ight can be
<H1>changed <H2>at <H1>any <H.5>time,
but care <H1>must<H.5>
<S2>be taken with the <H1>line<H.5> spacing
```

produces this output:



Line spacing

Command form: <Sxx.x>, <Sxx.x%> Default line spacing: 1.5× TXTHIT

To set the line spacing to xx.x units, include $\langle Sxx.x \rangle$ in the formatted text. The units, either centimeters or inches, are defined by the SET UNITS command. To set the line spacing as a percentage of the height of the current window, that is, YUWIND - YLWIND, use $\langle Sxx.x \rangle$.

The line spacing is the distance from the bottom of the previous text line to the bottom of the current text line. This spacing is the automatic vertical spacing to be used between text lines, but line spacing takes place *immediately*, so that each character in a text line may be drawn at any vertical distance, always measured from the bottom of the previous text line.

Within a file, the line spacing will remain as set until changed with another $\langle Sxx.x \rangle$ or $\langle Sxx.x \rangle$ command, but the line spacing always defaults to $1.5 \times$ the current value of TXTHIT when the PLOTTEXT command is entered.

```
<H0.5><S.8>Line spacing <S1.2>can be <S.8>changed
at any time in a line<S1.4>
and can be <S1.6>positive <S-.4>or negative<S1.4>
it may be set to zero
<S0><JL10>to continue a line<S1.2>
But be careful of setting to zero<S0>
at the end of a line
```

Line spacing changed at any time in a lineor negative

and can be positive

it may be set to zero to continue a line at the end of a line

But be careful of setting to zero

produces this output:

Left margin

Command form: <Mxx.x>, <Mxx.x%>

Default left margin: $0.01 \times (XUWIND - XLWIND)$

To set the left margin to xx.x units, include <Mxx.x> in the formatted text. The units, either centimeters or inches, are defined by the SET UNITS command. To set the left margin as a percentage of the width of the current window, that is, XUWIND - XLWIND, use <Mxx.x%>.

The left margin applies to the subsequent text lines. It does *not* apply to the current line.

Within a file, the left margin will remain as set until changed with another <Mxx.x> or <Mxx.x%> command, but the left margin always defaults to $0.01\times$ the width of the current window.

```
<H0.5><S1.2>An example using the left margin:
<M2.5>Margin changes take effect
on the next line, except it does
<JL3>affect justification immediately
<M3.5><JL2>affect justification immediately
as you can see
```

produces this output:

An example using the left margin:

Margin changes take effect

on the next line, except it does

affect justification immediately

affect justification immediately

as you can see

Bolding

Command form: <Bn>, <Bn:m>,

Default: no bolding

Bolding means that the text characters will be filled with a dot pattern or a hatch pattern. Bolding should only be used with the fonts: ROMAN.SERIF, ROMAN.FUTURA, ROMAN.FASHON, ROMAN.LOGO1, ROMAN.SWISSL, ROMAN.SWISSM, ROMAN.SWISSB, or TRIUMF.OUTLINE.

When <Bn> is encountered, subsequent characters will be filled. Within a file, bolding will remain as set until different fill pattern(s) are selected with another <Bn> or <Bn:m> command. Bolding is turned off by entering the command.

When $\langle \mathtt{Bn} : \mathtt{m} \rangle$ is encountered, the two *hatch* patterns $|\mathtt{n}|$ and $|\mathtt{m}|$ are used to fill the characters. This can be used to create a cross-hatching pattern. This feature *cannot* be used with two dot patterns.

```
<H.8><S1.5><B1><FROMAN.SERIF>Hatch fill bolding
font = <FROMAN.LOGO1>ROMAN.LOGO1
Bold pattern # 1
<B7:8>Bold patterns # 7 # 8<B>
<H.8><S2.5><B11>Dot pattern 11
<S1.5><B22>Dot pattern 22
<B33>Dot pattern 33<B>
```

Hatch fill bolding
font = ROMAN.LOGOI

Bold pattern # I

Bold patterns * 7 * *

Dot pattern II

Dot pattern ZZ

Pot pattern B3

produces this output:

Dot fill patterns

A dot pattern is of the form: uv, where the digit u is the increment number of dots to light up horizontally, $1 \le u \le 9$, and the digit v is the increment number of dots to light up vertically, $1 \le v \le 9$. For example, a dot pattern of 34 means to light up every third dot horizontally and every fourth dot vertically. If uv is negative, then the dots are erased instead of turned on. Note that 200 is interpreted the same as 211, that is, every dot is lit.

PostScript output

For PostScript output, set the POSTRES keyword to the appropriate resolution for your hard-copy device, using the SET command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark inkjet printer has a resolution of 360 dpi. A "good" picture can be obtained with POSTRES = 180 and LINTHK = 2.

Hatch patterns

A hatch pattern is composed of an angle and one to ten spacings. These spacings are simply cycled through as the region is being filled, that is, a line is drawn inside the region at the specified angle, then a parallel line is drawn at the first spacing, then another parallel line is drawn at the second spacing, and so on for the number of spacings in that pattern. This process is repeated until the region is filled. The hatch patterns can be redefined with the SET HATCH command and displayed with the DISPLAY FILL command. There are ten hatch patterns available.

Colour

Command form: <Cn>

Default colour: the current colour as set by the COLOUR command

To set the colour, include <Cn> in the text. This applies to subsequent text.

Within a file, the colour will remain as set until changed with another <Cn> command, but the colour always defaults to the current colour as set by the COLOUR command. See Table 2.6, page 20, for the colour associated with each colour code.

For example:

```
<FSCRIPT.2><S1.2><H.6><C2>Colour <C3>cannot <C4>be <C5>displayed <C6>on <C7>photocopies, <C1>but <C2>try <C3>this <C4>formatted <C5>text <C6>on a <C7>colour device
```

Font

Command form: <Ffontname>

Default font: the current font as selected with the SET FONT command

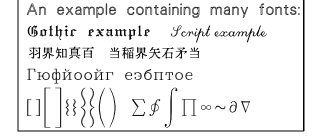
To select a font, include <ffontname> in the formatted text. This will apply to subsequent text. Within a file, the font will remain as set until changed with another <ffontname> command, but the font always defaults to the current font as set by the SET FONT command.

The manual TRIUMF GRAPHICS FONTS contains font tables. See Table 2.60 on page 243 for a list of the font names.

For example, this input:

```
<HO.5><S1><FROMAN.SWISSL>An example containing many fonts:
<FGOTHIC.ENGLISH>Gothic example <FSCRIPT.2>Script example
<FKANJI4>Kanji example
<FCYRILLIC.2>Cyrillic example
<S1.5><FMATH>0123456789 SJKMOPQR
```

produces this output:



Centre justification

Command form: <JC>, <JCxx.x>, <JCxx.x%>

To centre text at the location xx.x units from the left edge of the current window, that is, XLWIND, include <JCxx.x> before the text. The units, either centimeters or inches, are defined by the SET UNITS command. To centre the text at a location defined as a percentage of the width of the current window, that is, XUWIND - XLWIND, use <JCxx.x%>.

If no number is present, that is, <JC>, the text will be centred at the location midway in the window between the right edge, XUWIND, and the left edge.

The centred text will include all text after the <JC>, <JCxx.x> or <JCxx.x%> command up to the next justification command (justify left, right, or centre) or up to the end of the line, whichever comes first.

It is possible to centre about any position on the current line, in any order, and to mix left and right justifications with centering on the same line.

For example, this input:

```
<H0.5><S1.2>Example using centre justification:
<JC>This is centred on the page
<JC5>Column 1<JC12>Column 2
<JC5>-0.002<JC12>198.32
<S1.4><JC5><PM>0.75<JC12><Integrl><beta>d<sigma>
```

produces this output:

Left justification

Command form: <JL>, <JLxx.x>, <JLxx.x%>

To left justify text at the location xx.x units from the left edge of the current window, that is, XLWIND, include $\langle JLxx.x \rangle$ before the text. The units, either centimeters or inches, are defined by the SET UNITS command. To left justify the text at a location defined as a percentage of the width of the current window, that is, XUWIND - XLWIND, use $\langle JLxx.x\% \rangle$.

If no number is present, that is, <JL>, the text will be left justified at the left edge of the window.

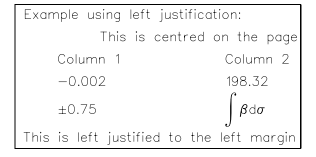
The left justified text will include all text after the <JL>, <JLxx.x> or <JLxx.x%> command and up to the next justification command (justify left, right, or centre) or up to the end of the line, whichever comes first.

It is possible to left justify about any position on the current line, in any order, and to mix left and right justifications with centering on the same line.

For example, this input:

```
<H0.5><S1.2>Example using left justification:
<JC>This is centred on the page
<JL11>Column 2<JL2>Column 1
<JL2>-0.002<JL11>198.32
<S1.5><JL2><PM>0.75<JL11><INTEGRL><beta>d<sigma>
This is left justified to the left margin
```

produces this output:



Right justification

Command form: <JR>, <JRxx.x>, <JRxx.x%>

To right justify text at the location xx.x units from the left edge of the current window, that is, XLWIND, include $\langle JRxx.x \rangle$ before the text. The units, either centimeters or inches, are defined by the SET UNITS command. To right justify the text at a location defined as a percentage of the width of the current window, that is, XUWIND - XLWIND, use $\langle JRxx.x\% \rangle$.

If no number is present, that is, <JR>, the text will be right justified at the right edge of the window, that is, XUWIND.

The right justified text will include all text after the <JR>, <JRxx.x> or <JRxx.x%> command and up to the next justification command (justify left, right, or centre) or up to the end of

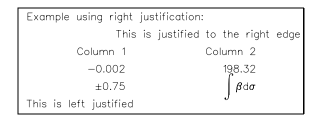
the line, whichever comes first.

It is possible to left justify about any position on the current line, in any order, and to mix left and right justifications with centering on the same line.

For example, this input:

```
<H0.5><S1.2>Example using right justification:
<JR>This is justified to the right edge
<JR16>Column 2<JR7>Column 1
<JR7>-0.002<JR16>198.32
<JR7><PM>0.75<JR16><INTEGRL><beta>d<sigma>
This is left justified
```

produces this output:



Horizontal spacing

Command form: <Zxx.x>, <Zxx.x%>

To insert a horizontal space of xx.x units, include $\langle Zxx.x \rangle$ in the formatted text. The units, either centimeters or inches, are defined by the SET UNITS command. To set the horizontal space as a percentage of the width of the current window, that is, XUWIND - XLWIND, use $\langle Zxx.x\% \rangle$.

The horizontal space, which may be positive or negative, is measured from the current location.

Note: The right, centre and left justification tabs, are measured from the left edge of the window.

```
<H0.5><S.8>Example of horizontal spacing:
<H.8><S2>Include a 2cm sp<Z2>ace
or move back-<Z-5><S4>wards and down
<S2>Remember to reset
the line spacing
```

produces this output:

Example of horizontal spacing:

Include a 2cm sp ace

or move back—

wards and down

Remember to reset

the line spacing

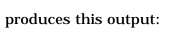
Sub-script mode

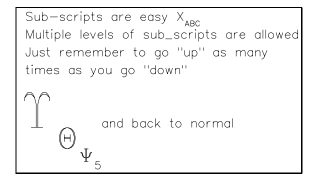
Command form: <_>

To enter sub-script mode, include <_> in the formatted text. Subsequent text will have 60% the current height and will be vertically spaced down a distance equal to 60% of the current height. This allows for multiple levels of sub-scripts, but for every level of sub-scripting, there must be a corresponding level of super-scripting to bring the text back "up".

Within a file, each level of sub-scripting remain in effect until <^>, super-script mode, is encountered.

```
<H0.5><S1>Sub-scripts are easy X<_>ABC<^>
Multiple levels of sub_scripts are allowed
Just remember to go "up" as many
times as you go "down"
<S3><H2><UPSILON><_><THETA><_><PSI><_>5<^><^><>><-><
<H.5>and back to normal
```





Super-script mode

Command form: <^>

To enter super-script mode, include <^> in the formatted text. Subsequent text will have 60% the current height and will be vertically spaced up a distance equal to 60% of the current height. This allows for multiple levels of super-scripts, but for every level of super-scripting, there must be a corresponding level of sub-scripting to bring the text back "down".

Within a file, each level of super-scripting remain in effect until <_>, sub-script mode, is encountered.

For example, this input:

```
<H0.5><S1>Super-scripts are easy X<^>ABC<_>
Multiple levels of super_scripts are allowed
Just remember to go "down" as many
times as you go "up"
<S3><H2><UPSILON><^><THETA><^><PSI><^>5<_><_><_><</p>
<H.5>and back to normal
```

produces this output:

Slanted mode

Command form:

To control slanted, or emphasis, mode, include in the formatted text. acts like a toggle switch in that the first time is encountered slanted mode will be turned on, that is, subsequent text will be slanted, and the next time it is encountered, slanted mode will be turned off. Slanted mode may be used for any character in any font, even with bolding on.

For example, this input:

```
<em><HO.6><S1.2>Slanted in the TSAN font
<FSTANDARD>Slanted in the STANDARD font
<FROMAN.SWISSL>Slanted in the ROMAN.SWISSL font
<FSCRIPT.2>Slanted in a SCRIPT font
<FROMAN.SERIF>Slanted in font ROMAN.SERIF
The final <Q\> <em>\Q<>\ turns <em>slanted mode off
```

produces this output:

Slanted in the TSAN font

Slanted in the STANDARD font

Slanted in the ROMAN.SWISSL font

Slanted in a SERIFI font

Slanted in font ROMAN.SBRIF

The final turns slanted mode off

Note: The graphics editor EDGR does not recognize slanted mode. So, if you open an EDGR file and include slanted text, when you edit the graphics with EDGR, the text will not be slanted.

Hexadecimal mode

Command form: <X>

To control hexadecimal text input mode, include <X> in the formatted text. <X> acts like a toggle switch in that the first time <X> is encountered, hexadecimal mode is turned on, and the next time it is encountered, hexadecimal mode is turned off. Hexadecimal mode means that the text is be assumed to be pairs of hexadecimal digits that represent non-keyboard characters.

The hexadecimal codes for characters depend on which font is being used. Refer to the font tables in the TRIUMF GRAPHICS FONTS manual for these codes. Also, see the DISPLAY FONT command for information on how to display any font table.

```
<H0.5><S.8>An example of hexadecimal input
in the default font <Rightarrow> TSAN
Greek letters:
<H1><S1.7><X>CACBCCCDCECFDADBDCDDDEDF<X>
<H.5><S1.2>and other symbols:
<H1><S1.7><X>4AAFB96954555657<X>
```

produces this output:

An example of hexadecimal input in the default font \Rightarrow TSAN Greek letters: $\alpha\beta\gamma\delta\epsilon\zeta\eta\delta\iota\kappa\lambda\mu$ and other symbols: $\hbar\phi \lesssim \beta\subset\cup\supset\cap$

Accents

| Command | |
|---------|---------------------------------------|
| <\b> | macron (bar) under previous character |
| <\d> | dot under previous character |
| <\^> | circumflex over previous character |
| <\',> | acute over previous character |
| ' | grave over previous character |
| <\"> | umlaut over previous character |
| <\~> | tilde over previous character |
| <\=> | macron (bar) over previous character |
| <\.> | dot over previous character |
| <\u> | breve over previous character |
| <\v> | v over previous character |
| <\H> | dieresis (double quote) over previous |
| | character |

Table 2.52: Formatted text accent special characters

To place one of the special accents on a character, insert the appropriate command immediately after that character. The accent will be centred over or under that character. See Table 2.52 for a listing of the accent commands. See Figure 2.21 for examples of the accents on the letter "o" in the TRIUMF.2 font.

Not all fonts allow for accents. For example, the cyrillic font or the hiragana font. The accent

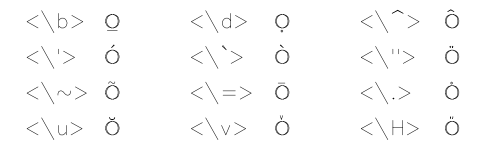


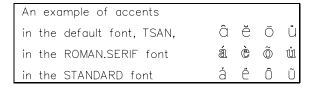
Figure 2.21: Example accents on the letter "o"

will be positioned fairly well over the lower case letters "a", "e", "o", and "u", but not perfectly positioned. You can use EDGR to relocate the accent if its position is not to your liking.

For example, this input:

```
<H0.3><S0.8>An example of accents
in the default font, TSAN, <->
<JL40%><H0.5>a<\^> e<\u> o<\=> u<\"><H0.3>
<FTSAN>in the ROMAN.SERIF font <->
<FROMAN.SERIF><JL40%><H0.5>a<\'> e<\'> o<\^> u<\.><H0.3>
<FTSAN>in the STANDARD font <->
<FSTANDARD><JL40%><H0.5>a<\v> e<\H> o<\^> u<\.><H0.3>
```

produces this output:



POLYGON

Syntax POLYGON xpoly ypoly xdata ydata key Qualifiers \INSIDE \INSIDE

The POLYGON command creates a vector, key, which will have the same length as the input vectors, xdata and ydata. By default, key[i] = 1 if the point (xdata[i],ydata[i]) is inside the polygon defined by input vectors xpoly and ypoly, otherwise key[i] = 0. If the \noint (noint is used, key[i] = 0 if the point (xdata[i],ydata[i]) is inside the polygon, otherwise key[i] = 1.

Example

You can use this command is conjunction with the PICK\POLYGON command to choose a polygon and the DESTROY command to eliminate data points within the chosen polygon. The following script produces Figure 2.22.

```
GEN\RANDOM X -5 5 200
                       ! generate some "data"
GEN\RANDOM Y 10 20 200 !
SET PCHAR -16
                       ! choose unjoined point plotting symbol
WINDOW 5
                       ! display the data graphically
GRAPH X Y
PICK\POLYGON XP YP
                       ! interactively choose a polygon around the
                          data you want to eliminate
POLYGON XP YP X Y K
                       ! create key vector, K
DESTROY X Y IFF (K=1)
                       ! eliminate unwanted data
WINDOW 6
GRAPH X Y
                       ! display data without unwanted points
SET PCHAR O
                       ! choose no plotting symbol, joined
GRAPH\NOAXES XP YP
                       ! overlay the polygon you chose above
```

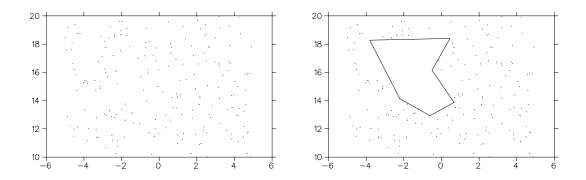


Figure 2.22: An example demonstrating the POLYGON command

QUIT

Syntax QUIT
Qualifiers \CLEAR
Defaults \NOCLEAR

The QUIT command is the cleanest way to stop the program. This does a complete FORTRAN STOP. You cannot re-enter PHYSICA after quitting, without re-running the program.

If the \CLEAR qualifier is appended to the QUIT command, the graphics is cleared before the program is stopped.

READ

```
READ file{\line_range} x1{\c1} { x2{\c2} ... }
Syntax
          READ\FORMAT file{\line_range} (frmt) x1 { x2 ... }
          READ\UNFORMATTED file{\line_range} (frmt) x1 { x2 ... }
          READ\SCALAR file\{\n\} s1\{\c1\} { s2\{\c2\} ... }
          READ\SCALARS\FORMAT file{\n} (frmt) s1 { s2 ... }
          READ\SCALARS\UNFORMATTED file{\n} (frmt) s1 { s2 ... }
          READ\MATRIX file{\n} m nrows { ncols }
          READ\MATRIX\FORMAT file{\n} (frmt) m nrows { ncols }
          READ\MATRIX\UNFORMATTED file\n (frmt) m nrows { ncols }
          READ\TEXT file{\line_range} txtvar
          READ\TEXT\FORMAT file{\line_range} (frmt) txtvar
          READ\TEXT\UNFORMATTED file{\line_range} (frmt) txtvar
Qualifiers \VECTORS, \SCALARS, \MATRIX, \TEXT, \ASCII, \UNFORMATTED, \FORMAT,
          \CONTINUE, \CLOSE, \APPEND, \OVERLAY, \EXTEND, \ERRSTOP, \ERRFILL,
          \ERRSKIP, \FLIPPED, \MESSAGES
          \VECTORS, \ASCII, \-FORMAT, \-CONTINUE, \-CLOSE, \-APPEND,
Defaults
          \-OVERLAY, \EXTEND, \ERRSTOP, \FLIPPED, \MESSAGES
Examples READ FILE.DAT X Y Z
          READ\APPEND\FORMAT FILE.DAT\3 (2X,3F10.2) X Y Z
          READ\FORMAT FILE.DAT\[2:100:2] (6X,F10.3,2X,F10.3) X\3 Y\1 Z\7
          READ FILE.DAT 4X
          READ\SCALAR FILE.DAT A B C
          READ\SCALAR\FORMAT FILE.DAT\3 (2X,3F10.2) A B C
          READ\SCALAR FILE.DAT\2 A\3 B\1 C\7
          READ\MATRIX FILE.DAT M 10 20
          READ\MATRIX\FORMAT FILE.DAT\3 (7(F10.3,2X)) M 10 20
          READ\TEXT FILE.DAT T
          READ\TEXT FILE.DAT\3 T
          READ\TEXT\FORMAT FILE.DAT\3 (2X,A10) T[2]
          READ\TEXT FILE.DAT\[2;3;15] T
```

READ is a general purpose command for reading vectors, scalars, a matrix, or string variables from a file. The maximum record length that can be read is 32768 bytes.

The variable type that will be read is determined by a command qualifier. The default, requiring no special qualifier, is to read data into vectors. Reading data into other variable types is chosen by using the appropriate qualifier. Refer to Table 2.53. The parameters that

are expected depend on which of these qualifiers is used.

| variable type | qualifier | |
|--------------------------------|-----------|-----------|
| multiple vectors | \VECTORS | (default) |
| multiple scalars | \SCALARS | |
| one matrix | \MATRIX | |
| one string or one string array | \TEXT | |

Table 2.53: Variables that can be read and their required qualifiers

By default, informational messages are displayed on the terminal monitor. If the \NOMESSAGES, or \-MESSAGES, qualifier is used, these informational messages will not be displayed.

Environment variables in file names

For UNIX users, it is now possible to use an environment variable in a file name, if the environment variable is preceded by a \$. For example,

```
setenv FILE dum.dat
physica
read $FILE x y z
```

The environment variable can be just the first part of the filename, for example,

```
setenv FILE dum
physica
read $FILE.dat x y z
```

Opening and closing files

By default, the file is closed after the read, so that a subsequent READ of the same file will start reading at the first record of the file. If the \CONTINUE qualifier is used, the file will not be closed after processing that READ command, and a subsequent READ of the same file will begin reading at the next record. If the \CLOSE qualifier is used, the file will be closed first and then reopened before reading, so that READ\CLOSE always begins reading with the first record in the file.

Examples

The command READ\SCALARS\CLOSE\CONTINUE DUM.DAT N first closes the file DUM.DAT, reads the first record, and leaves the file open.

If you entered: READ\SCALARS DUM.DAT N after the previous command, the second record would be read.

```
The commands READ\CLOSE\CONTINUE DUM.DAT\[1:10] X READ\CONTINUE DUM.DAT\[1:10] Y
```

would read the first ten records into vector X and the next ten records into vector Y.

Reading data into vectors

By default, vectors are read from columns of numbers in an ASCII file. The file is read by records, using free format. The cI^{th} column is placed into vector xI. cI defaults to I. Every record is read, from record 1 to the end of file. If xI exists, it will be destroyed, and a new xI vector created.

ASCII files

```
Syntax READ file{\line_range} x1{\c1} { x2{\c2} ... }
READ\FORMAT file{\line_range} (frmt) x1 { x2 ... }

Qualifiers \FORMAT, \CONTINUE, \CLOSE, \APPEND, \OVERLAY, \EXTEND, \ERRSTOP, \ERRSTILL, \ERRSKIP, \MESSAGES

Defaults \-FORMAT, \-CONTINUE, \-CLOSE, \-APPEND, \-OVERLAY, \EXTEND, \ERRSTOP, \MESSAGES
```

By default, or if the \ASCII qualifier is used, the file is assumed to be an ASCII file and is read by records, starting with the first record.

A scalar appended to the file name, file\n, specifies the starting record. The first n-1 records will be skipped.

A vector appended to the file name, file\x, specifies from which records to read data. The first x[1] - 1 records will be skipped. The data will be read from records x[1], x[2], ..., x[#]. Records x[i] + 1 to x[i+1] - 1 will be skipped. The vector x must be monotonically increasing.

By default, the I_{th} column is placed into vector xI. The column number can be specified by appending a scalar, cI, to the vector name as a qualifier. In this case, the cI_{th} column can be placed into the xI vector. For example, after the command:

```
READ DUM.DAT W\2 X\4 Y Z\1
```

W would contain column 2, X would contain column 4, Y would default to column 3, and Z would contain column 1.

By default, free format is used for reading the data. Number fields can be separated by blanks or by commas. The \FORMAT qualifier must be used to indicate that a format is

present. The format must be enclosed in parentheses, (and). If a format is used, column numbers *cannot* be specified. Standard FORTRAN formats are valid, but only REAL variables can be read, so do not use INTEGER, LOGICAL or CHARACTER formats. All values are converted to REAL*8 for internal storage.

To read different numbers of elements into vectors with a single READ command, use the \noextend, or \-extend, qualifier. The output length of a vector will be number of values that are read, to a maximum of that vector's original length. For example, suppose that vector X has length 10 and vector Y has length 20, and suppose you enter:

```
READ\-EXTEND file\[1:20] X Y
```

If 20 records are read, vector X will be made with a length of 10 and vector Y will be made with a length of 20. If only 15 records are read, vector X will have length 10 but vector Y will only have length 15.

By default, a new vector is created to hold the newly read data. If the \OVERLAY qualifier is used, an existing vector will have the newly read data overlayed on the original data. The resultant vector length may be longer, but never shorter. Use the \APPEND qualifier to append the newly read data onto the end of existing vectors.

```
\-EXTEND is incompatible with \OVERLAY. \-EXTEND is incompatible with \APPEND.
```

Field counts are specified by an integer preceding the vector name. Field counts must be literal integers, that is, they *cannot* be scalar variables. For example:

```
READ FILE DAT 3X 2Y Z
```

will read 6 numbers from each record, placing the first 3 numbers into X[i], X[i+1], X[i+2], the next 2 numbers into Y[i], Y[i+1], and the last number into Z[i]. This command is equivalent to:

```
READ FILE.DAT X X X Y Y Z
```

Records beginning with an exclamation mark, !, are considered to be comments and are ignored.

By default, \ERRSTOP, an invalid field stops the read, but the data that has been read up to the error is saved. If the \ERRSKIP qualifier is used, an invalid field causes the entire record to be skipped. If the \ERRFILL qualifier is used, an invalid field causes the entire record to be filled with the value of ERRFILL if a format was entered, or only the invalid field will be set

to ERRFILL if no format was entered. By default, ERRFILL=0, but it's value can be changed with the SET command and it's value obtained with the GET command.

Example 1

| command | result |
|-----------------------|--|
| READ FILE.DAT 3X 2Y Z | X = [1;2;3;7;8;9;13;14;15] Y = [4;5;10;11;16;17] Z = [6;12;18] |
| | Y = [4;5;10;11;16;17] |
| | Z = [6;12;18] |
| READ FILE.DAT 3X | X = [1; 2; 3; 7; 8; 9; 13; 14; 15] |
| READ DUM.DAT 6X | X = [1;2;3;4;5;6;7;8;9;10;11;12;13;14;15;16;17;18] |

Example 2

| command | result | |
|--|--|--|
| READ DUM.DAT X Y | X = [1;2;3;4;5;6] | |
| | Y = [23.7; -31.4; 9.09; 10.001; -2.0; 30.2] | |
| READ DUM.DAT\3 X Y | X = [3;4;5;6] Y = [9.09;10.001;-2.0;30.2] | |
| | Y = [9.09;10.001;-2.0;30.2] | |
| READ DUM.DAT\[2:4] X\3 Y | X = [.0002; .03; 4.0] | |
| | X = [.0002; .03; 4.0] Y = [-31.4; 9.09; 10.001] | |
| READ\FORMAT DUM.DAT\[1:5:2] (2X,F8.4,E9.4) X Y | X = [23.7; 9.09; -2.0] | |
| | Y = [.000001;.03;50.0] | |

Example 3

Sample data file: This script reads data into vectors and allows the user to choose when to stop reading.

```
10 20 30
                           LNUM = 1
40 50 60
                           START:
70 80 90
                           READ DUM.DAT\LNUM X Y Z
                           IF "EXIST('X') THEN GOTO DONE ! end of file
this is a test
100 200 300
                           LIST X Y Z
                           ANS='Y'
400 500 600
700 800 900
                           INQUIRE 'read again ? (Y|n)' ANS
                           IF NES(UCASE(ANS), 'Y') THEN GOTO DONE
test line two
-10 -20 -30
                           LNUM = LEN(X) + LNUM + 1
-40 -50 -60
                           GOTO START
-70 -80 -90
                           DONE:
```

Unformatted binary files

```
Syntax READ\UNFORMATTED file{\line_range} (frmt) x1 { x2 ... }

Qualifiers \CONTINUE, \CLOSE, \APPEND, \OVERLAY, \EXTEND, \MESSAGES

Defaults \-CONTINUE, \-CLOSE, \-APPEND, \-OVERLAY, \EXTEND, \MESSAGES
```

If the \UNFORMATTED qualifier is used, the file is assumed to be an unformatted binary file. The two methods of reading data from unformatted binary files, by record or stream, are indicated by the prescription paramter, (frmt), which is required when the \UNFORMATTED qualifier is used. The frmt prescription must be enclosed in parenthesis, (and). For example:

```
\label{eq:condition} \begin{split} &\text{real*4 x(100), y(100)}\\ &\text{do i = 1, 100}\\ &\text{write(unit)x(i),y(i)} &\text{! written by record} &\text{( 100 records )}\\ &\text{end do}\\ &\text{write(unit)x,y} &\text{! written as a stream ( 1 record )}\\ &\text{write(unit)(x(i),y(i),i=1,100)} &\text{! written as a stream ( 1 record )} \end{split}
```

The qualifiers \ERRSTOP, \ERRSKIP, and \ERRFILL cannot be used with the \UNFORMATTED qualifier.

Reading by record

The frmt codes: R4 R8 I1 I2 I4 L1 L4 Xn

specify that the data is to be read by record, and indicate the data type and the number of bytes for each value in each record. The Xn code specifies skipping over n bytes. All values are converted to REAL*8 for internal storage.

For example: READ\UNFORMATTED FILE.DAT (3R4,X8,I2) X Y Z A indicates 22 bytes per record: REAL*4, REAL*4, REAL*4, skip 8 bytes, INTEGER*2.

A vector appended to the file name, file\x, specifies from which records to read data. The first x[1] - 1 records will be skipped. The data will be read from records $x[1], x[2], \ldots, x[\#]$. Records x[i] + 1 to x[i+1] - 1 will be skipped. The vector x must be monotonically increasing.

To read different numbers of elements into vectors with a single READ command, use the \noextend, or \-extend, qualifier. The output length of a vector will be number of values that are read, to a maximum of that vector's original length. For example, suppose that vector X has length 10 and vector Y has length 20, and suppose you enter:

```
READ\UNFORMATTED\-EXTEND file\[1:20] (2R4) X Y
```

If 20 records are read, vector X will be made with a length of 10 and vector Y will be made with a length of 20. If only 15 records are read, vector X will have length 10 but vector Y will only have length 15.

By default, a new vector is created to hold the newly read data. If the \OVERLAY qualifier is used, an existing vector will have the newly read data overlayed on the original data. The resultant vector length may be longer, but never shorter. Use the \APPEND qualifier to append the newly read data onto the end of existing vectors.

```
\-EXTEND is incompatible with \OVERLAY. \-EXTEND is incompatible with \APPEND.
```

Field counts must be literal integers, that is, they *cannot* be scalar variables. Field counts are specified by an integer preceding the vector name.

For example: READ\UNFORMATTED FILE.DAT (6R8) 3X 2Y Z will read 6 numbers from each record, placing the first 3 numbers into X[i], X[i+1], X[i+2], the next 2 numbers into Y[i], Y[i+1], and the last number into Z[i]. This command is equivalent to: READ\UNFORMATTED FILE.DAT (6R8) X X X Y Y Z

Stream reading

The frmt codes: 1B 2B 4B 8B

specify that the data is to be read as a stream, and indicate the data type only. All values are converted to REAL*8 for internal storage. The number of values to read is indicated by creating the vectors before issuing the READ command. The number of values to read into a vector will be exactly the current length of that vector.

For example: VECTOR X 100

VECTOR Y 200

READ\UNFORMATTED FILE.DAT (4B) X Y

indicate that 100 REAL*4 values are to be read into X and 200 REAL*4 values into Y.

A scalar appended to the file name, file\n, specifies the starting record. The first n-1 records will be skipped.

Restrictions: vector line ranges *cannot* be used

held counts cannot be used \-EXTEND is always in effect \OVERLAY cannot be used \APPEND cannot be used

Example 1

Suppose you have written some data using the code fragment:

You could read this data using the commands:

```
READ\UNFORMATTED\SCALARS DUM.DAT (I2) N ! get length of vectors

VECTOR X Y Z N ! create vectors with length N

READ\UNFORMATTED DUM.DAT\2 (8B) X Y Z ! stream read vectors
```

Example 2

Suppose you have written some data using the code fragment:

You could read this data using the commands:

```
READ\UNFORMATTED\SCALARS DUM.DAT (I2) N ! get length of vectors READ\UNFORMATTED\MATRIX DUM.DAT\2 (8B) M 3 N ! stream read as a matrix X = M[1,*] ! extract first row Y = M[2,*] ! extract second row Z = M[3,*] ! extract third row
```

Example 3

Suppose you have written some data using the code fragment:

```
REAL*8 X(10), Y(10), Z(10)
...
OPEN(UNIT=20,FILE='DUM.DAT',FORM='UNFORMATTED')
DO I = 1, 10
  WRITE(20)X(I),Y(I),Z(I) ! write the arrays by record
END DO
...
```

You could read this data using the command:

```
READ\UNFORMATTED DUM.DAT (3R8) X Y Z ! read by record
```

Example 4

Suppose you have written some data using the code fragment:

```
REAL*8 X(10), Y(10), Z(10)

INTEGER*2 NUM
...

OPEN(UNIT=20,FILE='DUM.DAT',FORM='UNFORMATTED')

NUM = 10

WRITE(20)NUM
! first record contains length of arrays

WRITE(20)X
! stream write the X array

WRITE(20)Y
! stream write the Y array

WRITE(20)Z
! stream write the Z array
...
```

You could read this data using the commands:

```
READ\UNFORMATTED\SCALARS\CONTINUE DUM.DAT (I2) N ! don't close file

VECTOR X Y Z N ! create vectors of length N

READ\UNFORMATTED\CONTINUE DUM.DAT (8B) X ! stream read

READ\UNFORMATTED\CONTINUE DUM.DAT (8B) Y ! stream read

READ\UNFORMATTED\CONTINUE DUM.DAT (8B) Z ! stream read
```

Example 5

To read the unformatted binary file DUM.DAT, bytes 5-8 into X, bytes 9-12 into Y, bytes 13-16 into Z, enter the command:

```
READ\UNFORMATTED DUM.DAT (X4,3R4) X Y Z
```

To read the same file, but putting all the data into the vector Y, that is, bytes 5-8 into Y[i], bytes 9-12 into Y[i+1], bytes 13-16 into Y[i+2], where i=[1;4;7;...], enter the command:

```
READ\UNFORMATTED DUM.DAT (X4,3R4) 3Y
```

To start reading at the end of the second record of the file and to read THETA (INTEGER*4) from bytes 1-4, and PHI (REAL*8) from bytes 5-12, enter the command

```
READ\UNFORMATTED DUM.DAT\2 (I4,R8) THETA PHI
```

Reading data into scalars

The READ\SCALARS command reads scalar numbers from one record of a file. By default, the first record is read from an ASCII file, and, if no errors are encountered on the read, the \mathfrak{I}_{th} number is placed into scalar \mathfrak{sI} . New scalar variables are created. By default, no scalars will

be made if an invalid field is encountered on the read. A scalar appended to the file name, file\n, specifies the starting record. The first n-1 records will be skipped.

ASCII files

```
Syntax READ\SCALARS file{\n} s1{\c1} { s2{\c2} ... }
READ\SCALARS\FORMAT file{\n} (frmt) s1 { s2 ... }

Qualifiers \FORMAT, \CONTINUE, \CLOSE, \MESSAGES

Defaults \-FORMAT, \-CONTINUE, \CLOSE, \MESSAGES
```

By default, or if the \ASCII qualifier is used, the file is assumed to be an ASCII file and the first record is read. Specify which record to read by appending a scalar to the file name as a qualifier, file\n. By default, no scalars will be created if an invalid field is encountered while reading.

By default, the I_{th} number field is placed into scalar sI. The field number can be specified by appending a scalar, cI, to the scalar name as a qualifier. In this case, the cI_{th} field can be placed into the sI scalar.

For example, after the command: READ\SCALARS DUM.DAT W\2 X\4 Y Z\1 W would contain field 2, X would contain field 4, Y would default to field 3, and Z would contain field 1.

The \FORMAT qualifier must be used to indicate that a format is present. The format must be enclosed in parentheses, (and). If a format is used, field numbers *cannot* be specified.

Standard Fortran formats are valid, but only REAL variables can be read, so do not use INTEGER, LOGICAL or CHARACTER formats. All values are converted to REAL*8 for internal storage.

If the \ERRFILL qualifier is used, an invalid field causes the invalid field to be set to ERRFILL. By default, ERRFILL = 0, but it's value can be changed with the SET command, and obtained with the GET command. \ERRFILL cannot be used with a format.

Example

```
1 23.7 0.1000E-5
2 -31.4 0.2000E-3
3 9.09 0.3000E-1
```

| command | result | |
|--|-------------|-------------|
| READ\SCALARS DUM.DAT A B | A = 1 | B = 23.7 |
| READ\SCALARS DUM.DAT A\3 B | A = .000001 | B = 23.7 |
| READ\SCALARS DUM.DAT\3 A B | A = 3 | B = 9.09 |
| READ\SCALARS DUM.DAT\2 A\3 B | A = .0002 | B = -31.4 |
| READ\SCALARS\FORMAT DUM.DAT (2X,F8.4,E9.4) A B | A = 23.7 | B = .000001 |

Unformatted binary files

```
\begin{tabular}{ll} Syntax & READ\SCALARS\UNFORMATTED file{$\n$} & frmt) s1 $\{ s2 \dots \}$ \\ Qualifiers & $\CONTINUE, \CLOSE, \MESSAGES \\ \hline \end{tabular}
```

If the \UNFORMATTED qualifier is used, the file is assumed to be an unformatted binary file. By default, the first record is read. Specify which record to read by appending a scalar to the file name as a qualifier, file\n. No scalars will be created if an invalid field is encountered while reading. The (frmt) paramter is a prescription that specifies how the record is to be read and is required when the \UNFORMATTED qualifier is used. The prescription must be enclosed in parenthesis, (and).

The frmt codes: R4 R8 I1 I2 I4 L1 L4 Xn indicate the data type and the number of bytes for each value in the record. The Xn code is used to skip over n bytes. All values are converted to REAL*8 for internal storage.

Examples

```
READ\SCALARS\UNFORMATTED DUM.DAT (3R4,X8,I2) A B C D
```

reads 22 bytes from the first record: REAL*4, REAL*4, REAL*4, skip 8 bytes, INTEGER*2 and creates 4 scalars from these numbers.

```
READ\SCALARS\UNFORMATTED DUM.DAT (X10,R8) A
```

creates scalar A from the REAL*8 number in bytes 11 - 18 of the first record.

Reading data into a matrix

By default, a two dimensional array is read by records, starting with the first record, from an ASCII file, in free format, where nrows is the first dimension, the number of rows, and ncols is the second dimension, the number of columns. The first dimension *must* be entered

exactly. If the second dimension is not known, do not enter a value for ncols, and the read will continue until the end of file. The actual second dimension will be displayed when the read operation is done. A new matrix variable will be made. No matrix will be made if an error is encountered on the read.

A scalar appended to the file name, file\n, specifies the starting record. The first n-1 records will be skipped.

ASCII files

By default, or if the \ASCII qualifier is used, the file is assumed to be an ASCII file and is read by records. A scalar appended to the file name as a qualifier, file\n, specifies the starting record. The file will be read from the n_{th} record to the end of file.

The \FORMAT qualifier must be used to indicate that a format is present. The format must be enclosed in parentheses, (and).

Standard Fortran formats are valid, but only REAL variables can be read, so do not use INTEGER, LOGICAL or CHARACTER formats. All values are converted to REAL*8 for internal storage.

Flipped

Suppose you have a matrix M which has 3 rows and 4 columns. When you enter WRITE\MATRIX file M the rows of the matrix are written into records of the file. There will be 3 records, each containing 4 numbers. But, if you then entered READ\MATRIX file M 3 4 the input matrix would be scrambled, because READ puts the first record into the first column, the second record into the second column, and so on. Thus, the matrix is transposed, or flipped. To read it in properly, you would have to enter READ\MATRIX file M 4 3 and then take the transpose, M = <-M. So, there is a qualifier, \-FLIPPED, which has the syntax:

```
READ\MATRIX\-FLIPPED file matrix ncolumns nrows
```

Note: In the default, without the \-FLIPPED qualifier, the syntax remains: READ\MATRIX file matrix nrows in

Example 1

the code fragment creates the file REAL*4 M(7,5) 11. 21. 31. DO J = 1, 541. 51. 61. 71. 12. 22. DO I = 1, 7M(I,J) = 10*I+J32. 42. 52. 62. 72. 13. END DO 23. 33. 43. END DO 53. 63. 73. WRITE(1,10)M 14. 24. 34. 10 FORMAT(3F4.0) 44. 54. 64. 74. 15. 25. 35. 45. 55. 65.75. command result 11 12 13 14 15READ\MATRIX\FORMAT file (3F4.0) M 7 5 21 22 23 24 25 41 42 43 44 45 51 52 53 54 55 Example 2 REAL*4 M(7,5) DO J = 1, 5DO I = 1, 7

the code fragment

END DO
END DO
WRITE(1,10)M
10 FORMAT(14F4.0)

M(I,J) = 10*I+J

creates the file

11. 21. 31. 41. 51. 61. 71. 12. 22. 32. 42. 52. 62. 72.

13. 23. 33. 43. 53. 63. 73. 14. 24. 34. 44. 54. 64. 74.

15. 25. 35. 45. 55. 65. 75.

| command | result |
|--|--|
| READ\MATRIX\FORMAT file (14F4.0) M 7 5 | $M = \begin{pmatrix} 11 & 12 & 13 & 14 & 15 \\ 21 & 22 & 23 & 24 & 25 \\ 31 & 32 & 33 & 34 & 35 \\ 41 & 42 & 43 & 44 & 45 \\ 51 & 52 & 53 & 54 & 55 \\ 61 & 62 & 63 & 64 & 65 \\ 71 & 72 & 73 & 74 & 75 \end{pmatrix}$ |
| Example 3 | |
| the code fragment | creates the file |
| REAL*4 M(7,5) DO J = 1, 5 DO I = 1, 7 M(I,J) = 10*I+J END DO END DO DO J = 1, 5 WRITE(1,10)(M(I,J),I=1,7) END DO 10 FORMAT(4F4.0) | 11. 21. 31. 41. 51. 61. 71. 12. 22. 32. 42. 52. 62. 72. 13. 23. 33. 43. 53. 63. 73. 14. 24. 34. 44. 54. 64. 74. 15. 25. 35. 45. 55. 65. 75. |
| command | result |
| READ\MATRIX file M 7 5 | $M = \begin{pmatrix} 11 & 12 & 13 & 14 & 15 \\ 21 & 22 & 23 & 24 & 25 \\ 31 & 32 & 33 & 34 & 35 \\ 41 & 42 & 43 & 44 & 45 \\ 51 & 52 & 53 & 54 & 55 \\ 61 & 62 & 63 & 64 & 65 \\ 71 & 72 & 73 & 74 & 75 \end{pmatrix}$ |

Example 4

the code fragment

creates the file

| | REAL*8 M(7,5) | 11. | 12. | 13. | 14. | 15. |
|----|-----------------------------------|-----|-----|-----|-----|-----|
| | DO J = 1, 5 | 21. | 22. | 23. | 24. | 25. |
| | DO I = 1, 7 | 31. | 32. | 33. | 34. | 35. |
| | M(I,J) = 10*I+J | 41. | 42. | 43. | 44. | 45. |
| | END DO | 51. | 52. | 53. | 54. | 55. |
| | END DO | 61. | 62. | 63. | 64. | 65. |
| | WRITE(1,10)((M(I,J),J=1,5),I=1,7) | 71. | 72. | 73. | 74. | 75. |
| 10 | FORMAT(5F5.0) | | | | | |

command

result

```
11 21 31 41 51 61 71
READ\MATRIX file M 5
                                                                      12 \ \ 22 \ \ 32 \ \ 42 \ \ 52 \ \ 62 \ \ 72
                                                                      13 23 33 43 53 63 73
14 24 34 44 54 64 74
```

Example 5

the code fragment

creates the file

```
21.
  REAL*8 M(5,8)
                                                  31. 41. 51.
                                        11.
  DO J = 1, 8
                                        12.
                                             22.
                                                  32.
                                                       42.
                                                           52.
    DO I = 1, 5
                                        13.
                                             23.
                                                  33. 43. 53.
      M(I,J) = 10*I+J
                                        14.
                                             24.
                                                  34.
                                                       44. 54.
    END DO
                                        15.
                                             25.
                                                  35.
                                                       45.
  END DO
                                        16.
                                             26.
                                                  36. 46. 56.
                                        17.
  WRITE(1,10)((M(I,J),I=1,5),J=1,8)
                                             27.
                                                  37. 47. 57.
10 FORMAT(5F5.0)
                                        18.
                                             28.
                                                  38. 48. 58.
```

| command | | result |
|---|------------------|---|
| READ\MATRIX file M 5 | | $\mathbf{M} = \begin{pmatrix} 11 & 12 & 13 & 14 & 15 & 16 & 17 & 18 \\ 21 & 22 & 23 & 24 & 25 & 26 & 27 & 28 \\ 31 & 32 & 33 & 34 & 35 & 36 & 37 & 38 \\ 41 & 42 & 43 & 44 & 45 & 46 & 47 & 48 \\ 51 & 52 & 53 & 54 & 55 & 56 & 57 & 58 \end{pmatrix}$ |
| Example 6 | | |
| the code fragment | | creates the file |
| REAL*8 M(5,8) DO J = 1, 8 DO I = 1, 5 M(I,J) = 10*I+J END DO END DO WRITE(1,10)((M(I,J),J=1,8),1) 10 FORMAT(4F5.0) | (=1,5) result | 11. 12. 13. 14. 15. 16. 17. 18. 21. 22. 23. 24. 25. 26. 27. 28. 31. 32. 33. 34. 35. 36. 37. 38. 41. 42. 43. 44. 45. 46. 47. 48. 51. 52. 53. 54. 55. 56. 57. 58. |
| READ\MATRIX file M 4 | M = | $\begin{pmatrix} 11 & 15 & 21 & 25 & 31 & 35 & 41 & 45 & 51 & 55 \\ 12 & 16 & 22 & 26 & 32 & 36 & 42 & 46 & 52 & 56 \\ 13 & 17 & 23 & 27 & 33 & 37 & 43 & 47 & 53 & 57 \\ 14 & 18 & 24 & 28 & 34 & 38 & 44 & 48 & 54 & 58 \end{pmatrix}$ |
| command | result | |
| READ\MATRIX file M 8 | М = | 11 21 31 41 51 12 22 32 42 52 13 23 33 43 53 14 24 34 44 54 15 25 35 45 55 16 26 36 46 56 17 27 37 47 57 18 28 38 48 58 |

Example 7

Suppose you have the file DUM.DAT

Header line 1

Header line 2

1.0 2.0 3.0 4.0 5.0 6.0 7.0

1.1 2.1 3.1 4.1 5.1 6.1 7.1

1.2 2.2 3.2 4.2 5.2 6.2 7.2

1.3 2.3 3.3 4.3 5.3 6.3 7.3

1.4 2.4 3.4 4.4 5.4 6.4 7.4

command

result

```
READ\MATRIX DUM.DAT\3 M 7  M = \begin{pmatrix} 1.0 & 1.1 & 1.2 & 1.3 & 1.4 \\ 2.0 & 2.1 & 2.2 & 2.3 & 2.4 \\ 3.0 & 3.1 & 3.2 & 3.3 & 3.4 \\ 4.0 & 4.1 & 4.2 & 4.3 & 4.4 \\ 5.0 & 5.1 & 5.2 & 5.3 & 5.4 \\ 6.0 & 6.1 & 6.2 & 6.3 & 6.4 \\ 7.0 & 7.1 & 7.2 & 7.3 & 7.4 \end{pmatrix}
```

Unformatted binary files

```
Syntax READ\MATRIX\UNFORMATTED file{\n} (frmt) m nrows { ncols }
Qualifiers \CONTINUE, \CLOSE, \MESSAGES
Defaults \-CONTINUE, \-CLOSE, \MESSAGES
```

If the \UNFORMATTED qualifier is used, the file is assumed to be an unformatted binary file. The two methods of reading data from unformatted binary files, by record or stream, are indicated by the prescription, (frmt), paramter, which is required whith the \UNFORMATTED qualifier. A scalar appended to the file name as a qualifier, file\n, specifies the starting record. The frmt prescription must be enclosed in parenthesis, (and).

```
For example: real*8 x(100,10)

do j = 1, 10

    write(unit)(x(i,j),i=1,100) ! write 10 records

end do

    write(unit)x ! stream write

    write(unit)(x(i,j),i=1,100),j=1,10) ! stream write
```

Reading by record

```
Syntax \qquad \texttt{READ} \backslash \texttt{MATRIX} \backslash \texttt{UNFORMATTED file} \{ \backslash \texttt{n} \} \ \ (\texttt{frmt}) \ \ \texttt{m nrows} \ \ \{ \ \ \texttt{ncols} \ \ \}
```

The frmt codes: R4 R8 I1 I2 I4 L1 L4 Xn specify that the data is to be read by record, and indicate the data type and the number of bytes for each value in each record. The Xn code specifies skipping over n bytes. For example, (20R4) indicate 80 bytes per record: 20 REAL*4 values. All values are converted to REAL*8 for internal storage. If the number of columns, ncols, is not entered, records will be read until an end of file is reached.

Stream reading

```
Syntax READ\MATRIX\UNFORMATTED file{\n} (frmt) m nrows ncols
```

The frmt codes: 1B 2B 4B 8B

specify that the data is to be read as a stream, and indicate the data type only. The number of values to read is indicated by the number of rows, nrows, and the number of columns, ncols, *both* of which are required. All values are converted to REAL*8 for internal storage.

For example: READ\MATRIX\UNFORMATTED DUM.DAT (4B) M 10 100 indicate that 1000 REAL*4 values are to be stream read into matrix M with 10 rows and 100 columns.

Example 1

Consider the code fragment:

```
REAL*8 X(7,5)

INTEGER*2 NR, NC

DATA NR /7/, NC /5/

DO J = 1, 5

DO I = 1, 7

X(I,J) = 10*I+J

END DO

END DO

OPEN(UNIT=20,FILE='DUM.DAT',FORM='UNFORMATTED')

WRITE(20)NR,NC ! first record contains dimensions

WRITE(20)X ! stream write the array
```

| commands | result | | | | | | |
|---|--------|----------------|----------|----------|----------|-----------------|----------|
| READ\UNFORM\SCALARS DUM.DAT (212) NR NC | | 11 | 12 | 13 | 14 | 15 | <u> </u> |
| READ\UNFORM\MATRIX DUM.DAT\2 (8B) M NR NC | | 21 31 | 22 32 | 23 33 | 24 34 | $\frac{25}{35}$ | |
| | M = | 41 | 42 | 43 | 44 | 45 | |
| | | 51 | 52 | 53 | 54 | 55 | |
| | | 61 | 62 | 63 | 64 | 65 | |
| | | \setminus 71 | 72 | 73 | 74 | 75 / | / |

Example 2

Consider the code fragment:

```
REAL*8 X(7,5)

INTEGER*2 NR

DATA NR /7/

DO J = 1, 5

DO I = 1, 7

X(I,J) = 10*I+J

END DO

END DO

OPEN(UNIT=20,FILE='DUM.DAT',FORM='UNFORMATTED')

WRITE(20)NR ! first dimension, number of rows

DO J = 1, 5

WRITE(20)(X(I,J),I=1,7) ! write by records ( 5 records )

END DO
```

commands result

Reading data into a string variable

By default, the READ\TEXT command reads the first record from an ASCII file as a string, making a string variable. The maximum line length that can be read is 255 characters. If a single line is read, a string variable will be made. If multiple lines are read, an array string

variable will be made.

A scalar appended to the file name, file\n, specifies the record to read as data. The first n-1 records will be skipped. A string variable will be made.

A vector appended to the file name, file\x, specifies from which records to read data. The first x[1]-1 records will be skipped. The data will be read from records $x[1],x[2],\ldots,x[\#]$. Records x[i]+1 to x[i+1]-1 will be skipped. The vector x must be monotonically increasing. Multiple lines will be read only if multiple line numbers are indicated. a string array variable will be made.

ASCII files

```
Syntax READ\TEXT file{\line_range} txtvar
READ\TEXT\FORMAT file{\line_range} (frmt) txtvar

Qualifiers \FORMAT, \CONTINUE, \CLOSE, \MESSAGES

Defaults \-FORMAT, \-CONTINUE, \-CLOSE, \MESSAGES
```

By default, when no format is entered, the entire line, up to 255 characters is read. The \FORMAT qualifier must be used to indicate that a format is present. The format must be enclosed in parentheses, (and).

```
The frmt codes: nX means to skip over n characters An means to read n characters
```

For example: (2X,A10) means to skip the first two characters and read the next ten.

Examples

| command | result |
|-------------------------|---|
| READ\TEXT DUM.DAT\4 TXT | reads the fourth record into a scalar string variable |
| | reads 5 strings from records 4 to 8 |
| | into a string array variable |

Unformatted binary files

```
Syntax READ\TEXT\UNFORMATTED file{\line_range} (frmt) txtvar

Qualifiers \CONTINUE, \CLOSE, \MESSAGES

Defaults \-CONTINUE, \-CLOSE, \MESSAGES
```

If the \UNFORMATTED qualifier is used, the file is assumed to be an unformatted binary file. By default, the first record is read. The (frmt) paramter is a prescription that specifies

how the record is to be read and is required when the \UNFORMATTED qualifier is used. The prescription must be enclosed in parenthesis, (and). The only allowable prescription is (An), which means to read n characters. For example: (A10) means to read the first ten characters from the record.

REBIN

Syntax REBIN v vout n
REBIN m mout nr nc

The REBIN command rebins the data in either:

the vector v by the compression factor n; or the matrix m by the row compression factor nc and the column compression factor nc.

Rebinning vectors

Syntax REBIN v vout n

Suppose that the length of vector v is N then:

$$\mathtt{vout}[i] = \sum_{k=1}^{\mathtt{n}} \mathtt{v}[(i-1)\mathtt{n} + k] \quad \mathbf{for} \quad i = [1:\frac{N}{\mathtt{n}}],$$

that is, the length of vout will be $\frac{N}{n}$, and

$$\mathtt{vout}[1] = \sum_{i=1}^{\mathtt{n}} \mathtt{v}[i]$$

$$\mathtt{vout}[2] = \sum_{i=\mathtt{n}+1}^{2\mathtt{n}} \mathtt{v}[i]$$

. . .

$$\mathtt{vout}[rac{N}{\mathtt{n}}] = \sum_{i=(rac{N}{\mathtt{n}}-1)\mathtt{n}+1}^{rac{N}{\mathtt{n}}\mathtt{n}}\mathtt{v}[i]$$

If $(\frac{N}{n}) \times n$ is not equal to N, then the last element of vout will be incomplete. For example, if N=10 and n=3 then vout will have 3 elements:

vout[1] = v[1] + v[2] + v[3]
vout[2] = v[4] + v[5] + v[6]
vout[3] = v[7] + v[8] + v[9]

and v[10] will not be included in vout.

Examples

Suppose that vector X has 20 elements. After the command: REBIN X XOUT 2

```
XOUT[1] = X[ 1] + X[ 2]
XOUT[2] = X[ 3] + X[ 4]
XOUT[3] = X[ 5] + X[ 6]
...
XOUT[10] = X[19] + X[20]
```

After the command: REBIN [1:1000] X 3

$$X[1] = 1+2+3$$

 $X[2] = 4+5+6$
 $X[3] = 7+8+9$
...
 $X[333] = 997+998+999$

A warning will be given that the length of [1:1000] is not evenly divisible by 3 and so the last bin is incomplete.

Rebinning matrices

Syntax REBIN m mout nr nc

Suppose that matrix m has N rows and M columns, then:

$$\mathtt{mout}[i,j] = \sum_{l=1}^{\mathtt{nc}} \sum_{k=1}^{\mathtt{nr}} \mathtt{m}[(i-1)\mathtt{nr} + k, (j-1)\mathtt{nc} + l] \quad \mathbf{for} \quad i = [1:\frac{N}{\mathtt{nr}}], j = [1:\frac{M}{\mathtt{nc}}],$$

that is, the matrix mout will have $\frac{N}{nr}$ rows and $\frac{M}{nc}$ columns.

$$\mathtt{mout}[1,1] = \sum_{j=1}^{\mathtt{nc}} \ \sum_{i=1}^{\mathtt{nr}} \mathtt{m}[i,j]$$

$$\mathtt{mout}[1,2] = \sum_{j=\mathtt{nc}+1}^{2\mathtt{nc}} \sum_{i=1}^{\mathtt{nr}} \mathtt{m}[i,j]$$

$$\mathtt{mout}[2,1] = \sum_{j=1}^{\mathtt{nc}} \sum_{i=\mathtt{nr}+1}^{2\mathtt{nr}} \mathtt{m}[i,j]$$

$$\mathtt{mout}[2,2] = \sum_{j=\mathtt{nc}+1}^{2\mathtt{nc}} \sum_{i=\mathtt{nr}+1}^{2\mathtt{nr}} \mathtt{m}[i,j]$$

. . .

If $(\frac{N}{\mathtt{nr}} \times \mathtt{nr}$ is not equal to N, then the last row of mout will be incomplete. If $(\frac{M}{\mathtt{nc}} \times \mathtt{nc}$ is not equal to M, then the last column of mout will be incomplete.

Example

Suppose that M is a matrix:

$$\mathtt{M} = \begin{pmatrix} 1 & 2 & 3 & 4 & 5 & 6 \\ 2 & 4 & 6 & 8 & 10 & 12 \\ 3 & 6 & 9 & 12 & 15 & 18 \\ 4 & 8 & 12 & 16 & 20 & 24 \\ 5 & 10 & 15 & 20 & 25 & 30 \\ 6 & 12 & 18 & 24 & 30 & 36 \\ 7 & 14 & 21 & 28 & 35 & 42 \\ 8 & 16 & 24 & 32 & 40 & 48 \end{pmatrix}$$
 Then after the command: REBIN M MOUT 2 3

$$\mathtt{MOUT} = \begin{pmatrix} 18 & 45 \\ 42 & 105 \\ 66 & 165 \\ 90 & 225 \end{pmatrix}$$

Note that:

$$\begin{aligned} \text{MOUT}[1,1] &= 1+2+3+2+4+6 \\ \text{MOUT}[1,2] &= 4+5+6+8+10+12 \\ \\ \text{MOUT}[2,1] &= 3+6+9+4+8+12 \\ \\ \text{MOUT}[2,2] &= 12+15+18+16+20+24, \quad \text{etc.} \end{aligned}$$

REFRESH

Syntax REFRESH

The REFRESH command is relevant only when using an X Window monitor. The X Window graphics window, and the zoom window, if it exists, will be redrawn. This has no affect on a graphics hardcopy.

RENAME

| Syntax | RENAME oldname newname |
|--------|------------------------|
| | RENAME old* new* |
| | RENAME *old *new |

The RENAME command renames variables. The simplest case is to rename one variable. For example:

```
RENAME XOLD XX ! renames the variable XOLD to XX
```

If the wildcard * is the last character in the old name, all variables beginning with the preceding characters will be renamed. For example:

```
RENAME X1* XX* ! renames X1 to XX, X1X to XXX, and X1Y to XXY
```

If the wildcard is the first character in the old name, all variables that end with the succeeding characters will be renamed. For example:

```
RENAME *X *Y2 ! renames X to Y2, AX to AY2, XX to XY2
```

REPLOT

```
Syntax REPLOT { nw }
REPLOT\ALLWINDOWS

Qualifiers \AXES, \ALLWINDOWS, \TEXT

Defaults \AXES, nw = current window number, \-TEXT
```

By default, information pertaining to curves produced by the GRAPH command, as well as an automatically plotted curve from the ELLIPSE command, are saved in replot buffers. When a set of axes is drawn and then data is overlayed on those axes, all of the data may not appear within the axis boundaries. You can use the REPLOT command to have all of the data appear on a single autoscaled graph. The window number, nw, defaults to the current window number. When the \NOAXES qualifier is used, the data will be replotted but the graph axes will not be drawn.

What is saved

For each data set, the window number in which it was drawn is saved, along with the plotting symbols, including their types, colours, connectivity, angles, and sizes are saved for replotting. Also saved are: error bar types, line thicknesses, line types, colours, and histogram types.

Strings

Strings that were drawn with the TEXT command will also be replotted. If the \GRAPH qualifier is used with the TEXT command, the location of the replotted text will remain fixed with respect to the graph's coordinate system. Otherwise, the location of the replotted text will remain fixed with respect to the window.

The \backslash -TEXT qualifier means to replot the data curves but not any strings that had also been drawn using the TEXT command. The default is \backslash TEXT, which means to also replot any such strings.

Enable/Disable

After the DISABLE REPLOT command is entered, subsequent data curves will not be saved in the replot buffers. Use the ENABLE REPLOT command to re-enable saving. If replotting is enabled, it can be disabled for individual commands by use of the \normale normale qualifier, for example, GRAPH\normale normale and commands by use of the \normale normale normale and commands by use of the \normale normale normale and commands by use of the \normale normale normale

Windows

If the window number nw is entered, the current window becomes window nw. Only the data drawn into window nw will be replotted.

If the REPLOT command is entered without a window number, and without the \ALLWINDOWS qualifier, the graphics will be automatically cleared. If using a window number or \ALLWINDOWS , it is up to the user to ensure that the appropriate windows are cleared before entering the REPLOT command.

Clearing the graphics

Use the CLEAR\NOREPLOT command to clear all graphics without emptying the replot buffers.

Use the ERASEWINDOW, page 76, command to erase the graphics from a window. This only applies to the terminal monitor, to PostScript graphics, and to bitmap graphics.

Use the CLEAR\REPLOTONLY command to clear the replot buffers without clearing the graphics.

Redraw all windows

Use the REPLOT\ALLWINDOWS command to replot all windows that have some data stored in the replot buffers. It is up to the user to ensure that the appropriate windows are cleared before entering the REPLOT\ALLWINDOWS command.

Examples

The following sequence of commands draws a graph of 20*X versus 10*X and overlays a graph of $(X^3)/10$ versus X^2 on the same set of axes. To see the complete drawing, the REPLOT command is used.

Suppose that you have plotted a graph of vectors Y1 vs. X1 in window number 5; a graph of vectors Y2 vs. X2 in window number 7, and overlain a plot of Y3 vs. X3 also in window 7. To replot both windows, use the REPLOT\ALLWINDOWS command.

```
WINDOW 5 ! choose window

GRAPH X1 Y1 ! plot data and axes

WINDOW 7 ! choose window

GRAPH X2 Y2 ! plot data and axes

GRAPH\NOAXES X3 Y3 ! overlay a curve

CLEAR\NOREPLOT ! clear graphics but not replot buffers

REPLOT\ALL ! replot window 5 and window 7
```

RESIZE

```
Syntax RESIZE
```

The RESIZE command brings up the graphics cursor, allowing PHYSICA to know when the graphics window is resized by dragging a corner or side of the graphics window. Quit RESIZE by pressing the Q key while the cursor is in the graphics window.

Using the RESIZE command before changing the shape of the graphics window is not necessary under VMS or Digital Unix. Resizing should be done with the RESIZE command under Linux, Silicon Graphics IRIX, Sun SOLARIS, and IBM AIX or the PHYSICA window will not match the actual graphics window.

Use the REFRESH command to simply redraw the contents of the graphics window, which is sometimes needed after resizing or uncovering the graphics window.

RESTORE

```
Syntax RESTORE filename

Qualifiers \PHYSICA, \FIOWA, \XFIOWA, \MUD, \MSR, \IMSR, \CHAOS, \YBOS, \HBOOK

Defaults \PHYSICA
```

The RESTORE command is used to restore specially formatted data sets:

previously saved PHYSICA sessions; FIOWA type data sets; μ SR type data sets; μ SR MUD type data sets; μ SR type data sets; CHAOS type data sets; YBOS type data sets, HBOOK type data sets.

Environment variables in file names

For UNIX users, it is now possible to use an environment variable in a file name, if the environment variable is preceded by a \$. For example,

```
setenv FILE mysession
physica
restore $FILE
```

The environment variable can be just the first part of the filename, for example,

```
setenv FILE my
physica
restore $FILEsession
```

PHYSICA sessions

```
Syntax RESTORE file
Qualifier \TTBUFFERS

Default \TTBUFFERS
```

By default, the RESTORE command, or RESTORE\PHYSICA, restores a previously saved PHYSICA run. Normally it would be used immediately after entering PHYSICA to continue a previous session that was saved with the SAVE command, page 224. An attempt is made to restore

the plots that were active at the time the SAVE command was issued, similar to entering the REPLOT\ALLWINDOWS command. Since some graphics is not REPLOTable, such as contour plots and density plots, this does not always work as expected. The hardcopy device as chosen with the DEVICE command is restored, as well as the orientation.

Note: This command should be used with caution, since the current run will be irrevocably lost in the process of restoring a previous run.

The run time loaded subroutine or function information is not restored, but must be regenerated explicitly in the restored session with the LOAD command, page 147.

The input line recall buffers, that is, the dynamic buffer, the static buffer, and the keypad buffer, are restored also, by default. If you do not want to restore these buffers when resuming a PHYSICA session, use the \-TTBUFFERS qualifier.

FIOWA data sets

Syntax RESTORE\FIOWA filename

The RESTORE\FIOWA command restores FIOWA type data files containing histograms and scatterplots. No special qualifiers are need to restore data sets made with the "big" version of FIOWA. When you restore a file (or map to shared memory), the following PHYSICA variables are created automatically:

| | variable name | type | description |
|---|---------------|--------|---|
| ٠ | DATA | vector | all the histogram and scatterplot data |
| | NHIST | scalar | number of histograms |
| | NSCAT | scalar | number of scatterplots |
| | HTITLE | string | string containing all the histogram titles |
| | HXLABEL | string | string containing all the histogram labels |
| | STITLE | string | string containing all the scatterplot titles |
| | SXLABEL | string | string containing all the scatterplot xlabels |
| | SYLABEL | string | string containing all the scatterplot ylabels |
| | | | |

Histograms

The following histogram related vectors are created automatically:

| variable name | description |
|---------------|---------------------------|
| NBINS | number of bins |
| HSTART | starting index |
| HXLO | x minimum |
| HXINC | bin size |
| HYLO | y minimum |
| HYINC | y increment |
| HLO | underflow |
| HHI | overflow |
| STRTHT | starting index for title |
| STRTHX | starting index for xlabel |
| LENHT | length of title |
| LENHX | length of xlabel |
| | |

Scatterplots

The following scatterplot related variables are created automatically:

| variable name | type | description |
|---------------|--------|---------------------------|
| NSBINX | vector | number of bins in x |
| NSBINY | vector | number of bins in y |
| ISCAT | vector | index of first bin |
| SXLO | vector | x minimum |
| SXINC | vector | x increment |
| SYLO | vector | y minimum |
| SYINC | vector | y increment |
| OUTSID | matrix | underflows and overflows |
| STRTST | vector | starting index for title |
| STRTSX | vector | starting index for xlabel |
| STRTSY | vector | starting index for ylabel |
| LENST | vector | length of title |
| LENSX | vector | length of xlabel |
| LENSY | vector | length of ylabel |

The scatterplot overflows and underflows are stored in a matrix named OUTSID, which always has 8 rows, and is defined below for scatterplot number j.

| x-low | x-ok | x-high |
|-------------|----------------------|----------------------|
| y-high | y-high | y-high |
| OUTSID[4,j] | OUTSID[3,j] | OUTSID[2,j] |
| x-low | x-ok | x-high |
| y-ok | <i>y</i> - ok | <i>y</i> - ok |
| OUTSID[5,j] | | OUTSID[1,j] |
| x-low | x-ok | x-high |
| y-low | y-low | y-low |
| OUTSID[6,j] | OUTSID[7,j] | OUTSID[8,j] |

FIOWA examples

To draw histogram n:

```
label\xaxis htitle[strtht[n]:strtht[n]+lenht[n]-1]
scalar\dummy j
graph\hist loop(hxlo[n]+(j-0.5)*hxinc[n],j,1:nbins[n]) -
data[hstart[n]:hstart[n]+nbins[n]-1]
```

To draw scatterplot n, using the diffusion type of density plot:

```
label\xaxis stitle[strtst[n]:strtst[n]+lenst[n]-1]
scalar\dummy j
x = loop(sxlo[n]+(j-0.5)*sxinc[n],j,1:nsbinx[n])
y = loop(sylo[n]+(j-0.5)*syinc[n],j,1:nsbiny[n])
m = fold(data[iscat[n]:iscat[n]+nsbinx[n]*nsbiny[n]-1],nsbinx[n])
density\profiles\diffusion x y <-m ! <- is the transpose operator</pre>
```

XFIOWA data sets

```
Syntax RESTORE\XFIOWA filename
```

The RESTORE\XFIOWA command restores eXpanded FIOWA type data files containing histograms and scatterplots (as modified by Tom Huber from the original FIOWA). These data files are ASCII and so are transportable between platforms, unlike the original FIOWA files which are written as unformatted data.

No special qualifiers are need to restore data sets made with the "big" version of FIOWA. When you restore an XFIOWA file, you get the same PHYSICA variables as with a FIOWA file.

HBOOK data sets

Syntax RESTORE\HBOOK filename

RESTORE\HBOOK\DIR filename directory

Examples RESTORE\HBOOK FILE.DAT

RESTORE\HBOOK\DIR FILE.DAT '/SUB1/SUB2'

The RESTORE\HBOOK command restores HBOOK type data files containing histograms and 2d histograms (scatterplots). If the \DIR qualifier is used, the ZEBRA RZ directory within the file can be specified, using the absolute pathname in the ZEBRA syntax. When you restore a file, you get the following PHYSICA variables:

| variable name | type | description |
|---------------|--------|--|
| DATA | vector | all the histogram and scatterplot data |
| HERROR | vector | histogram and scatterplot errors |
| NHIST | scalar | number of histograms |
| NSCAT | scalar | number of scatterplots |
| HTITLE | string | string containing all the histogram titles |
| STITLE | string | string containing all the scatterplot titles |

The HERROR vector contains the error contents of all the histograms and scatterplots. Access this information in the same way as you access the /verb+DATA+ vector, for example, for histogram /verb+n+: error = herror[hstart[n]:hstart[n]+nbins[n]-1].

Listing

If you use the \LIST qualifer, a listing of the contents of the data file will be displayed on your monitor screen. If the \LIST qualifier is used, the ZEBRA RZ directory within the file can be specified, using the absolute pathname in the ZEBRA syntax. The listing is produced by the HLDIR routine from the CERN library.

Ntuples

```
Syntax RESTORE\HBOOK\RWN filename id RESTORE\HBOOK\RWN\DIR filename directory id
```

This command will restore the Row Wise Ntuple with identification number id from an HBOOK data file. If the \DIR qualifier is used, the ZEBRA RZ directory within the file can be entered, using the absolute pathname in the ZEBRA syntax. By default, a vector will be created for each variable in the Ntuple, where the names of these vectors will be the tag

names of the variables in the Ntuple.

For now, Column Wise Ntuples cannot be restored in PHYSICA.

If the \MATRIX qualifier is used, an array string variable, named \Mathbb{NTAGSn} , will be created containing the tag names for the variables in the Ntuple, and a matrix, named $\Mathbb{NTUPLEn}$, will be created containing the data, where n is the identification number id.

Histograms

The following histogram related vector variables will be created automatically:

| variable na | me description |
|-------------|-----------------------------|
| ID1 | histogram identifier |
| NBINS | number of bins |
| NOENT | number of entries |
| HMEAN | mean value |
| HSTD | standard deviation |
| HEQUIV | number of equivalent events |
| HSTART | starting index |
| HXLO | x minimum |
| HXINC | bin size |
| HLO | underflow |
| HHI | overflow |
| STRTHT | starting index for title |
| LENHT | length of title |
| | |

NOENT[j] is the number of original entries in histogram [j], including underflows and overflows.

 ${\tt HLO[j]}$ is the sum of the weights for events below the histogram lower limit. If you did not use weights when filling the histogram, then ${\tt HLO}$ is the number of events below the histogram lower limit.

HHI[j] is the sum of the weights for events above the histogram upper limit. If you did not use weights when filling the histogram, then HHI is the number of events above the histogram upper limit.

The number of equivalent events returned in HEQUIV is based on the channel contents only. If w_i represents the contents of event i, then

number of equivalent events
$$= \sum (w_i)^2 / \sum (w_i^2)$$

This is what it is supposed to be, *but* it seems to be just the total number of entries in the histogram, which is the same as NOENT. To calculate this value inside PHYSICA, you could use:

```
y = data[hstart[n]:hstart[n]+nbins[n]-1]
! hsum will be the sum of the weights inside the histogram limits
statistics\-message y hsum\sum
! hsum2 will be the sum of the squares of the weights
statistics y*y hsum2\sum
hequiv[n] = hsum*hsum/hsum2
```

The standard deviation returned in HSTD is based on the channel contents only. If x_i and w_i represent the value and contents of event i, and $W = \sum (w_i)$, then

expectation value
$$E(x) = \sum (w_i x_i)/W$$

mean value $= E(x)$
central moment of order n, $M(n) = E((x - E(x))^n)$
standard deviation $= \sqrt{(M(2))}$

The mean value, or expectation value, returned in HMEAN is based on the channel contents only. If x_i and w_i represent the value and contents of event i, and $W = \sum (w_i)$, then

mean value =
$$\sum (w_i x_i)/W$$

Scatterplots

The following 2d histogram related vector variables will be created automatically:

| variable name | type | description |
|---------------|--------|--------------------------|
| ID2 | vector | scatterplot identifier |
| SNOENT | vector | number of entries |
| NSBINX | vector | number of bins in x |
| NSBINY | vector | number of bins in y |
| ISCAT | vector | index of first bin |
| SXLO | vector | x minimum |
| SXINC | vector | x increment |
| SYLO | vector | y minimum |
| SYINC | vector | y increment |
| OUTSID | matrix | underflows and overflows |
| STRTST | vector | starting index for title |
| LENST | vector | length of title |

SNOENT[j] is the number of original entries in scatterplot [j], including underflows and overflows.

The overflows and underflows for scatterplot j are stored in a matrix named OUTSID, which always has 8 rows, and is defined below.

| x-low | x-ok | x-high |
|-------------|-------------|-------------|
| y-high | y-high | y-high |
| OUTSID[4,j] | OUTSID[3,j] | OUTSID[2,j] |
| x-low | x-ok | x-high |
| y-ok | y-ok | y-ok |
| OUTSID[5,j] | | OUTSID[1,j] |
| x-low | x-ok | x-high |
| y-low | y-low | y-low |
| OUTSID[6,j] | OUTSID[7,j] | OUTSID[8,j] |

HBOOK examples

To draw histogram n:

```
label\xaxis htitle[strtht[n]:strtht[n]+lenht[n]-1]
scalar\dummy j
graph\hist loop(hxlo[n]+(j-0.5)*hxinc[n],j,1:nbins[n]) -
data[hstart[n]:hstart[n]+nbins[n]-1]
```

To draw scatterplot n, using the diffusion type of density plot:

```
label\xaxis stitle[strtst[n]:strtst[n]+lenst[n]-1]
scalar\dummy j
x = loop(sxlo[n]+(j-0.5)*sxinc[n],j,1:nsbinx[n])
y = loop(sylo[n]+(j-0.5)*syinc[n],j,1:nsbiny[n])
m = fold(data[iscat[n]:iscat[n]+nsbinx[n]*nsbiny[n]-1],nsbinx[n])
density\profiles\diffusion x y <-m ! <- is the transpose operator</pre>
```

YBOS data sets

```
Syntax RESTORE\YBOS filename
Qualifier \DOTPLOT
Default \NODOTPLOT
```

The RESTORE\YBOS command restores YBOS type data files containing histograms or dotplots. Use the \DOTPLOT qualifier to specify that you want to restore a file containing dotplots. By default, the file is assumed to contain histogram data.

Histograms

When you restore (or map to shared memory) a YBOS histogram file, for histogram number j, you get the following PHYSICA variables:

| variable name | type | YBOS structure name or description |
|---------------|--------------|------------------------------------|
| HRUNNO | scalar | YBS.I_RUNNO |
| HTITLE[j] | array string | YBS.HISTO_ST(j).DOT_NAME |
| HTEST[j] | array string | YBS.HISTO_ST(j).TEST_NAME |
| HDATA[j] | array string | YBS.HISTO_ST(j).DATA_NAME |
| HIST | vector | YBS.HISTO |
| HSTART[j] | vector | starting index for histo j |
| HNBIN[j] | vector | YBS.HISTO_ST(j).N_BINS |
| HLOW[j] | vector | YBS.HISTO_ST(j).XLOW |
| HHI[j] | vector | YBS.HISTO_ST(j).XHI |
| HBINW[j] | vector | (HHI[j]-HLOW[j])/HNBIN[j] |
| HUNDER[j] | vector | YBS.HISTO_ST(j).XUNDER |
| HOVER[j] | vector | YBS.HISTO_ST(j).XOVER |
| HSUM[j] | vector | sum of histo j |
| HMEAN[j] | vector | mean value of histo j |
| HSIGMA[j] | vector | sigma of histo j |

Dotplots

Syntax RESTORE\YBOS\DOTPLOT file

When you restore (or map to shared memory) a YBOS dot plot file, for dot plot number j, you get the following PHYSICA variables:

| variable name | type | YBOS structure name or description |
|------------------------|--------------|--|
| DRUNNO | scalar | YBS.I_RUNNO |
| <pre>X_SYMBOL[j]</pre> | array string | YBS.DOT_EXTRA(j).X_SYMBOL |
| Y_SYMBOL[j] | array string | YBS.DOT_EXTRA(j).Y_SYMBOL |
| STEST_ALL[j] | array string | YBS.DOT_EXTRA(j).STEST_ALL |
| DOT_NAME[j] | array string | YBS.DOT_ST(j).DOT_NAME |
| SCTEST[j] | array string | YBS.DOT_EXTRA(j).SCTEST[1-4] |
| XDOT | vector | YBS.XDOT |
| YDOT | vector | YBS.YDOT |
| XLOW[j] | vector | YBS.DOT_ST(j).X_LOW |
| XHIGH[j] | vector | YBS.DOT_ST(j).X_HIGH |
| YLOW[j] | vector | YBS.DOT_ST(j).Y_LOW |
| YHIGH[j] | vector | YBS.DOT_ST(j).Y_HIGH |
| SCTEST_START[j,1:4] | matrix | starting character index for SCTEST[j] |
| SCTEST_END[j,1:4] | matrix | final character index for SCTEST[j] |
| COL_POINTER[j,1:4] | matrix | YBS.DOT_ST(j).COL_ST(1:4).COL_POINTER |
| COL_OFF[j,1:4] | matrix | YBS.DOT_ST(j).COL_ST(1:4).COL_OFF |

Furthermore: SCTEST[j][SCTEST_START[j,k]:SCTEST_END[j,k]] = YBS.DOT_EXTRA(j).SCTEST[k]

YBOS examples

To draw histogram n:

```
label\xaxis htitle[n]
scalar\dummy j
graph loop(hlow[n]+(j-1)*hbinw[n],j,1:hnbin[n]) -
hist[hstart[n]:hstart[n]+hnbin[n]-1]
```

To draw dotplot n:

```
do j = [1:4]
  if (col_off[n,j] > 0) then
    scale xlow[n] xhigh[n] 5 ylow[n] yhigh[n] 5
    graph\axesonly
    colour j
    graph\noaxes -
    xdot[col_pointer[n,j]:col_pointer[n,j]+col_off[n,j]-1] -
    ydot[col_pointer[n,j]:col_pointer[n,j]+col_off[n,j]-1]
    set
    %xloc 32
    %yloc 90-3*j
    %txthit 2.0
      cursor -2
    text rchar(col_off[n,j],'F6.0')//'//sctest[n]
  endif
enddo
```

μ SR MUD data sets

```
Syntax RESTORE\MUD file
```

The RESTORE\MUD command restores μ SR MUD type data files containing histograms. The actual histogram data is stored in a vector, HISTDATA, and histogram number j can be extracted using: HISTDATA[HISTSTART[j]:HISTEND[j]]. The global title is in a string variable called TITLE. The title for histogram number j is in the array string variable HISTITLE[j]. For more information on the MUD data format, visit the URL:

http://www.triumf.ca/dagroup/modas/mud_friendly.html

General variables

| scalars | strings |
|------------|--------------|
| RUNNUMBER | TITLE |
| RUNDESC | LAB |
| EXPTNUMBER | AREA |
| ELAPSEDSEC | METHOD |
| TIMEBEGIN | APPARATUS |
| TIMEEND | INSERT |
| | SAMPLE |
| | ORIENT |
| | DAS |
| | EXPERIMENTER |

Comments

| scalars | vectors | strings |
|-------------|-------------|---------------|
| COMMENTTYPE | COMMENTPREV | COMMENTAUTHOR |
| COMMENTNUM | COMMENTNEXT | COMMENTTITLE |
| | COMMENTTIME | COMMENTBODY |

Histograms

| scalars | vectors | strings |
|---------|-----------------|-----------|
| HISTNUM | NUMBINS | HISTTITLE |
| | HISTSTART | |
| | HISTEND | |
| | HISTDATA | |
| | HISTTYPE | |
| | HISTNUMBYTES | |
| | HISTBYTESPERBIN | |
| | HISTFSPERBIN | |
| | HISTTO_PS | |
| | HISTTO_BIN | |
| | HISTGOODBIN1 | |
| | HISTGOODBIN2 | |
| | HISTBDGD1 | |
| | HISTBKGD2 | |
| | NUMEVENTS | |

Scalers

| scalars | vectors | strings |
|------------|--------------|-------------|
| SCALERTYPE | SCALERCOUNTS | SCALERLABEL |

Independent variables

| scalars | vectors | strings |
|------------|----------------|-------------------|
| INDVARTYPE | INDVARLOW | INDVARNAME |
| | INDVARHIGH | INDVARDESCRIPTION |
| | INDVARMEAN | INDVARUNITS |
| | INDVARSTDDEV | |
| | INDVARSKEWNESS | |

μ SR data sets

```
Syntax RESTORE\MSR file
```

The RESTORE\MSR command restores μ SR type data files containing histograms. Histogram number j is stored in column j of the matrix IH. The global title is in a string variable called TITLE. The title for histogram number j is in the array string variable HTITLE[j]. There are also associated scalars:

```
IRUN is the run number

NHISTS is the number of histograms

NBINS is the number of bins

NS_BIN is the binning increment
```

The associated vectors:

| TOTEV[j] | is the total number of events in histogram j |
|----------|---|
| ITO[j] | is the start of background for histogram <code>j</code> |
| IT1[j] | is the start of data for histogram j |
| IT2[j] | is the end of data for histogram j |

To plot histogram J, enter:

```
GRAPH\HISTOGRAM [0:(NBINS-1)*NSBIN:NSBIN] IH[1:NBINS,J]
```

$I\mu$ SR data sets

```
Syntax RESTORE\IMSR file
```

The RESTORE\IMSR command restores $|\mu$ SR type data files. Histogram number j is stored in column j of the matrix IX. The run number is stored in a scalar called IRUNNO. The version number is stored in a scalar called IVERS. The global title is in a string variable called ITITLE. The subtitle, if it exists in the file, is in a string variable called ISUBTITLE. The title for histogram number j is in the array string variable XTITLE[j].

To plot histogram J, enter:

```
GRAPH\HISTOGRAM [1:VLEN(M)(1)] IX[*,J]
```

CHAOS data sets

The RESTORE\CHAOS command restores CHAOS type data files containing histograms. The run number is stored in a scalar variable named RUN_NUMBER. The number of events analyzed will be stored in a scalar variable named I_ANALYZED. The number of histograms and the number of channels are stored in scalar variables named NHIST and NCHAN. The vectors XLO, XHI, NBINS, and HSTART will be created. Each of these will have NHIST elements. The histogram data will be stored in a vector named HIST, with NCHAN elements.

The array string variable HNAMES will be created, with NHIST elements. HNAMES[i] is the name of histogram number i. The array string variable TITLES will be created, with NHIST elements. TITLES[i] is the title of histogram number i. The array string variable EVENT_CALIB will be created, with NHIST elements. EVENT_CALIB[i] is the event/calibration flag for histogram number i.

The data for histogram number I is located in HIST[HSTART[i]:NBINS[i]+HSTART[i]-1]. To make an x vector for plotting histogram number I, enter:

GENERATE X XLO[I],,XHI[I] NBINS[I]+HSTART[I]-1

RETURN

Syntax RETURN

The RETURN command is to be used in conjunction with script files.

If the RETURN command is encountered in a script file, control passes back to the calling script, if there is one, or to the keyboard, if that script was the top level script. You can also type RETURN from the keyboard after a TERMINAL command to abort that script.

SAVE

Syntax SAVE file

The SAVE command saves all the data associated with the current run, that is, all variable names, contents, and histories. This includes all scalars, vectors, matrices, and string variables. This information is written in a special binary format into the specified file. The run may be resumed later by means of the RESTORE command, page 211.

The plot information necessary for a replot is also saved. The window definitions are saved,

as well as the colour, the default filename extension for executable files, the stack file, autoscaling type, and any aliases that have been defined. The hardcopy device as chosen with the DEVICE command is saved, as well as the graphics ORIENTATION.

User defined routine information is not saved, but must be regenerated explicitly in the restored session.

SCALAR

```
Syntax SCALAR s1 { s2 ... }

Qualifiers \DUMMY, \VARY

Examples SCALAR A B C

SCALAR\DUMMY I J K

SCALAR\VARY P1 P2
```

The SCALAR command defines sI to be a scalar. If sI exists and is a scalar, there is no affect. If sI exists but is not a scalar, it will be destroyed first. If sI does not exist, it will be created and initialized to the value one (1).

Fit parameters

```
Syntax SCALAR\VARY s1 { s2 ... }
```

The SCALAR\VARY command defines sI to be a scalar variable as above, but it also allows sI to vary during a FIT. These fit parameters are treated as fixed value scalars, except by the FIT command. The FIT command can vary a parameter's value to minimize the least squares residual and the result is still a scalar.

To fix a parameter so it can no longer vary use the SCALAR command with no qualifier.

Dummy variables

```
Syntax SCALAR\DUMMY s1 { s2 ... }
```

The SCALAR\DUMMY command defines sI to be a scalar dummy variable for use in functions requiring dummy variables: LOOP, SUM, PROD, RSUM, and RPROD. Dummy variables cannot be used in other contexts, since they have no fixed values.

SCALES

```
Syntax SCALES { minx maxx nlxinc miny maxy nlyinc }
SCALES minx maxx miny maxy

Qualifiers \COMMENSURATE

Defaults non-commensurate axes

Examples SCALES
SCALES SCALES
SCALES 0 0 0 .1 .5 0
SCALES\COMM -10 5 3 .01 .05 4
```

The SCALES command turns off the autoscaling feature, as set with the SET AUTOSCALE command, and sets the graph scales for subsequent graphs as follows:

```
minx - minimum value to display on the x-axis

maxx - maximum value to display on the x-axis

nlxinc - number of large (numbered) tic marks on the x-axis

miny - minimum value to display on the y-axis

maxy - maximum value to display on the y-axis

nlyinc - number of large (numbered) tic marks on the y-axis
```

If no parameters are entered, the scales will be frozen at their current values, and the current values of minx, maxx, nlxinc, miny, maxy, and nlyinc will be displayed.

If four (4) parameters are entered, the number of labeled tic marks on the x and y-axes, that is, nlxinc and nlyinc, default to zero (0), which means the program will choose "nice" numbers of increments.

Commensurate axis scaling

If the \COMMENSURATE qualifier is used, the axes will be drawn with commensurate scaling, that is, the lengths of both axes will be adjusted to give a commensurate graph. The lengths of the axes are adjusted at the time the SCALES\COMMENSURATE command is given, which means that if you change windows, with the WINDOW command, after the SCALES\COMMENSURATE command, the axis lengths will be incorrect. Choose the window you want and then set the axis scales.

Labeled tic marks

The number of large, labeled, tic marks to be displayed on the x-axis is controlled by nlxinc. Its initial value is 5%.

| nlxinc | < 0 | the number of large tic marks on the x -axis will be $nlxinc +1$. If the x - |
|--------|-----|---|
| | | axis is logarithmic, that is, if ${\tt XLOG}>1$, then ${\tt nlxinc}$ may be changed to |
| | | avoid fractional powers. If the x -axis is linear, that is, if $XLOG < 1$, then |
| | | the x -axis minimum and maximum will not be labeled with numbers. |
| nlxinc | = 0 | the number of large tic marks on the x-axis will be automatically deter- |
| | | mined for the user. The virtual x -axis minimum and maximum will be |
| | | determined, so the x -axis may not begin or end at a large tic mark |
| nlxinc | > 0 | the number of large tic marks on the x -axis will be $nlxinc +1$. If $XLOG > 1$, |
| | | logarithmic x -axis, $nlxinc$ may be changed to avoid fractional powers. If |
| | | XLOG < 1, linear x -axis, $nlxinc$ will not be changed. |

The number of large, labeled, tic marks to be displayed on the y-axis is controlled by \mathtt{nlyinc} . Its initial value is 5%.

| nlyinc | < 0 | the number of large tic marks on the y -axis will be $\mathtt{nlyinc} + 1$. If the y -axis |
|--------|-----|---|
| | | is logarithmic, that is, if $YLOG > 1$, then $nlyinc may be changed to avoid$ |
| | | fractional powers. If the y -axis is linear, that is, if $YLOG < 1$, then the y -axis |
| | | minimum and maximum will not be labeled with numbers. |
| nlyinc | =0 | the number of large tic marks on the y-axis will be automatically deter- |
| | | mined for the user. The virtual y -axis minimum and maximum will be |
| | | determined, so the y -axis may not begin or end at a large tic mark |
| nlyinc | > 0 | the number of large tic marks on the y -axis will be $nlyinc +1$. If $YLOG > 1$, |
| | | logarithmic y -axis, nlyinc may be changed to avoid fractional powers. If |
| | | YLOG < 1 , linear y -axis, nlyinc will not be changed. |

Example

The following script produces Figure 2.23:

```
WINDOW 5
                                  ! choose a window
SCALES -.5 .5 2 -5 5 2
                                  ! set the graph scales
                                  ! plot only the axes
GRAPH\AXESONLY
GET
                                  ! get some GPLOT keyword values
 %XLAXIS XLX
 %XUAXIS XUX
                                  ! don't forget the blank line
SET
                                  ! set some GPLOT keyword values
 CURSOR -2
                                  ! centred text
                                  ! x reference point for text
 %XLOC (XLX+XUX)/2
 %YLOC 50
                                  ! y reference point for text
                                  ! don't forget the blank line
TEXT 'SCALES -.5 .5 2 -5 5 2'
                                  ! draw a string
WINDOW 6
SCALES 10 50 4 -100 -80 2
GRAPH\AXESONLY
TEXT 'SCALES 10 50 4 -100 -80 2' !
WINDOW 7
SET YLOG 10
                                  ! y-axis logarithmic, base 10
SCALES 10 50 4 2 6 4
GRAPH\AXESONLY
TEXT 'SCALES 10 50 4 2 6 4'
SET %YLOC 40
TEXT 'YLOG = 10'
SET %YLOC 50
WINDOW 8
SET YLOG -2
                                  ! base 2, numbered without powers
GRAPH\AXESONLY
TEXT 'SCALES 10 50 4 2 6 4'
SET %YLOC 40
                                  !
TEXT 'YLOG = -2'
```

SET

```
Syntax SET { keyword { value }}
SET { keyword = value }

Examples SET %XLAXIS 20
SET %XLAXIS=XLX
```

The SET command sets the values of the GPLOT plot characteristic keywords and the PHYSICA specific keywords. Use the GET command, page 107, to obtain the values of these keywords.

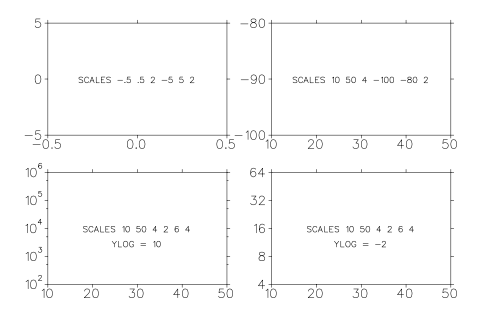


Figure 2.23: An example using the SCALES command

How the SET command works

If the SET command is entered with no parameters, more than one keyword value can be obtained without re-entering the SET command. Other keywords and values will be requested, until a blank line is entered, at which time the user is put back into command line entry mode.

If the SET command is entered with no parameters in a script file, a blank line is *necessary* to indicate that the SET command is finished.

If a keyword is entered with the SET command, then only that one keyword's value can be changed, with that command. If a value is not entered after the keyword, the current value is displayed. The current value is unchanged if no new value is entered.

Examples

To display the current value of XMIN, enter: GET XMIN

The following script gets the current value of XMIN and then changes it to XMIN-10, and sets the value of XMAX to XMIN+100

```
GET XMIN A ! makes a scalar A

SET

XMIN A-10

XMAX A+100 ! don't forget the blank line to finish
...
```

GPLOT keywords

See **Appendix A** for descriptions of all the GPLOT plotting characteristic keywords. The tables produced by the MENU contain most of the keywords that can be accessed with the SET and GET commands, along with their current values.

The GPLOT keywords: MASK, ALIAS, PMODE, PTYPE, and ERRBAR, should *not* be changed in PHYSICA, as these are internally adjusted and used by various commands.

The PHYSICA keywords

The keywords, ARROLEN, ARROTYP, and ARROWID, apply to arrows drawn with the FIGURE ARROW command, page 82. See Figure 2.24 for examples of the available arrow styles.

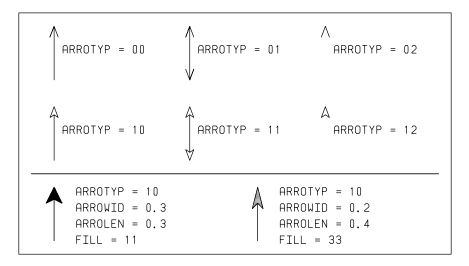


Figure 2.24: Arrow styles

ARROLEN

Default value: ARROLEN = 0.20

ARROLEN is the arrow head length expressed as a fraction of the length of the arrow's shaft.

ARROTYP

Default value: ARROTYP = 00

ARROTYP determines the type of arrow to draw. See Table 2.54.

| ARROTYP | | arrow style |
|---------|--------------|---|
| 00 | open head, | line segment with arrowhead at one end |
| 01 | open head, | line segment with arrowhead at each end |
| 02 | open head, | arrowhead at one end with no line segment |
| 10 | closed head, | line segment with arrowhead at one end |
| 11 | closed head, | line segment with arrowhead at each end |
| 12 | closed head, | arrowhead at one end with no line segment |

Table 2.54: The ARROTYP code and corresponding arrow styles

ARROWID

Default value: ARROWID = 0.15

ARROWID is the arrow head width expressed as a fraction of the length of the arrow's shaft.

AUTOSCALE

```
ParametersON | OFF | XAXIS | YAXIS | COMMENSURATE

Parameter \VIRTUAL

Qualifier

Examples SET AUTOSCALE ON

SET AUTOSCALE XAXIS\VIRTUAL

SET AUTOSCALE COMMENSURATE
```

The AUTOSCALE keyword controls autoscaling for graph axes. Autoscaling means to automatically choose the minimum and maximum values for the axes, as well as the number of large, numbered, tic marks for the axes. Autoscaling affects commands that draw axes, that is, the commands GRAPH, CONTOUR, DENSITY, REPLOT, and SLICES. The type of autoscaling that is done depends on the keyword that is used.

| keyword | result |
|--------------|--|
| ON | Autoscale the horizontal and the vertical axes |
| COMMENSURATE | Autoscale the horizontal and vertical axes and |
| | change the lengths of the axes so that they will be commensurate |
| XAXIS | Autoscale the horizontal axis only, |
| | the vertical axis will remain as currently set |
| YAXIS | Autoscale the vertical axis only, |
| | the horizontal axis will remain as currently set |
| OFF | turn off all autoscaling, |
| | the axes will appear as they are currently set |

Autoscaling remains in effect until either the command SET AUTOSCALE OFF is entered, or the SCALES command is entered. The default is ON.

When the \VIRTUAL qualifier is used, the virtual minima and maxima for the axes will be determined, so that the axes may not begin or end at a large tic mark. If the keyword ON is used, both x- and y-axes will have virtual minima and maxima. If the keyword YAXIS is used, only the x-axis will have virtual minimum and maximum. If the keyword YAXIS is used, only the y-axis will have virtual minimum and maximum.

CNTSEP

Default value: %CNTSEP = 50%

The separation between contour labels can be set with CNTSEP or %CNTSEP. If %CNTSEP is used, the separation is a percentage of the height of the window, that is, YUWIND-YLWIND. If CNTSEP is used, the separation is expressed in centimeters or inches, depending on the units type as set with the SET UNITS command.

LABSIZ

Default value: %LABSIZ = 1.5

LABSIZ, or %LABSIZ, is the size of the contour labels. %LABSIZ is the size as a percentage of the height of the window, that is, YUWIND-YLWIND, while LABSIZ is the size expressed in centimeters or inches, depending on the units type as set with the SET UNITS command.

LEGSIZ

Default value: LEGSIZ = 1.6

LEGSIZ, or %LEGSIZ, is the size of the contour plot and density plot legend entries. %LEGSIZ is the size as a percentage of the height of the window, that is, YUWIND-YLWIND, while LEGSIZ is the size expressed in centimeters or inches, depending on the units type as set with the SET UNITS command.

LEGFRMT

Default value: LEGFRMT = $^{\circ}1PE10.3'$

The numeric legend entries drawn by the DENSITY and CONTOUR commands are written using the LEGFRMT format.

ERRFILL

Default value: ERRFILL = 0

If the \ERRFILL qualifier is used with the READ command, and an invalid field is encountered on a record in the data file, then ERRFILL will be used.

format used: all fields on that record will be set to ERRFILL no format used: only the invalid field will be set to ERRFILL

FILL

Default value: FILL = 0

FILL is used in the FIGURE command, with the fillable figures: BOX, POLYGON, WEDGE, CIRCLE, ELLIPSE, and ARROWS with closed heads. It is also used for filling the boxes with the DENSITY\BOX command. See Table 2.55 for a description of the interpretations of the FILL keyword.

| | | FILL | = | 0 | no filling |
|----|--------|------|--------|----|--|
| 1 | \leq | FILL | \leq | 10 | fill with hatch pattern FILL |
| 11 | \leq | FILL | \leq | 99 | fill with dithering pattern |
| | | | | | if $ FILL = nm$, n is the increment number of dots hori- |
| | | | | | zontally and m is the increment number of dots to light |
| | | | | | up vertically |
| | | FILL | < | 0 | erase using FILL as above |

Table 2.55: Interpretations of the FILL keyword

For example, FILL = 34 means to light up every 3^{rd} dot horizontally and every 4^{th} dot verti-

cally. FILL = -34 means to erase every 3^{rd} dot horizontally and every 4^{th} dot vertically. FILL = 8 means to fill with hatch pattern 8. FILL = -8 means to erase using hatch pattern 8.

See the SET HATCH command for information on changing the hatch pattern definitions. See the DISPLAY HATCH command for information on how to display examples of the hatch patterns.

HATCH

```
Parameters: hnum { spaces angle }
```

The SET HATCH command is used for changing the hatch pattern definitions that are used for text bolding, for filling areas under curves or histograms, and for use by the TILE, PIEGRAPH, and FIGURE commands.

The SET HATCH command does *not* choose the hatch pattern to be used by other commands. It only alters the definition of a hatch pattern.

If just the keyword HATCH is entered, a table of the spacings and angles for all ten hatch patterns is displayed. You will be requested to enter a hatch pattern number, hnum, a spacing scalar or a vector of spacings, spaces, and an angle, angle. Typing the <RETURN> key, without entering anything, will leave the current values unchanged. If the hatch pattern number, a spacing scalar or vector and an angle are entered, then nothing will be displayed. For example, to set hatch pattern 3 to have spacings of [0.05;0.1;0.2] and an angle of 45°, enter:

```
SET HATCH 3 [0.05;0.1;0.2] 45
```

while to set hatch pattern 7 to have a single spacing of 0.5 and an angle of -45°, enter:

```
SET HATCH 7 .5 -45
```

The hatch pattern number, hnum, should be between one and ten. A spacing vector, spacevec, must have no more than ten elements. A hatch pattern is composed of an angle and from one to ten spacings. The default spacings and angles are listed in Table 2.56. The angles are in degrees and the spacing lengths, by default, are expressed in centimeters, but if the units are changed to inches, with the SET UNITS command, the lengths will be converted to inches. See Figure 2.7 on page 61 for examples of the hatch patterns. See the DISPLAY command, page 58, for information on how to display examples of the hatch patterns.

When an object is being filled, a line is drawn inside the object at the specified angle, then a parallel line is drawn at the first spacing, and so on for the number of spacings in that

pattern. This process is repeated until the object is filled.

| Pattern | Spac | cings | |
|---------|------|-------|-------|
| Number | 1 | 2 | angle |
| 1 | 0.01 | | 0 |
| 2 | 0.01 | | 90 |
| 3 | 0.05 | | 0 |
| 4 | 0.05 | | 90 |
| 5 | 0.10 | | 0 |
| 6 | 0.10 | | 90 |
| 7 | 0.20 | | 45 |
| 8 | 0.20 | | -45 |
| 9 | 0.20 | 0.10 | 45 |
| 10 | 0.20 | 0.10 | -45 |

Table 2.56: The hatch pattern defaults

LINE

Parameters: n { v }

The SET LINE command is used for changing the definition of the line types that are used by the commands: GRAPH, LINE, PICK, ELLIPSE, FIGURE, and ZEROLINES.

This command does *not* choose the line type to be used by other commands. It only alters the definition of a line type. To choose a line type, use the SET LINTYP command.

If just the keyword LINE is entered, a table of the spacings for all ten line types is displayed. You will be requested to enter a line type number, n, and a vector of spacings, v. Typing the <RETURN> key, without entering anything, will leave the current values unchanged. If the line type number and a spacing vector are entered, then nothing will be displayed. For example, to set line type 2 to [0.05;0;0], enter:

```
SET LINE 2 [0.05;0;0]
```

The line type number, n, should be between one and ten. The spacing vector, v, must have three elements. The line types come in four different styles:

- ordinary solid line
- double line of specified width
- dashed line with specified dash and space lengths
- dashed line with two different dash lengths

The different line types are achieved by specifying the three lengths v[1], v[2], and v[3] as shown in Table 2.57.

| v[1] | v[2] | v[3] | Result |
|------|------|------|---|
| =0 | | | ordinary solid line |
| > 0 | =0 | | double line with width v[1] |
| > 0 | > 0 | =0 | dashed line with dash length v[1], space length v[2] |
| > 0 | > 0 | > 0 | dashed line with first dash length $v[1]$, space length $v[2]$, and second dash length $v[3]$ |
| | | | v[2], and second dash length v[3] |

Table 2.57: Line type definitions

There are ten line types available. The defaults are listed in Table 2.58. The lengths are expressed in centimeters, the default, but if the units are changed to inches, with the SET UNITS command, the lengths will be converted to inches. See Figure 2.8 on page 62 for examples of the default line types. See the DISPLAY command, page 58, for information on how to display examples of the line types.

| line type | v[1] | v[2] | v[3] |
|-----------|------|------|------|
| 1 | 0.00 | 0.00 | 0.00 |
| 2 | 0.07 | 0.00 | 0.00 |
| 3 | 0.50 | 0.30 | 0.00 |
| 4 | 0.50 | 0.30 | 0.10 |
| 5 | 0.30 | 0.30 | 0.00 |
| 6 | 0.30 | 0.30 | 0.10 |
| 7 | 0.20 | 0.20 | 0.00 |
| 8 | 0.20 | 0.20 | 0.05 |
| 9 | 0.05 | 0.20 | 0.00 |
| 10 | 0.05 | 0.30 | 0.00 |

Table 2.58: The line type defaults

SHOWHISTORY

Default value: $\mathtt{SHOWHISTORY} = 5$

SHOWHISTORY controls how many lines of history to display for each numeric variable as a result of the SHOW command.

MAXHISTORY

SHOWHISTORY

 $egin{array}{lll} n & < & 0 & \longrightarrow & \mbox{all stored history lines will be displayed} \\ n & = & 0, 1 & \longrightarrow & \mbox{only the latest history line will be displayed} \\ n & > & 0 & \longrightarrow & \mbox{a maximum of n lines of history will be displayed for each variable} \\ \end{array}$

Table 2.59: The SHOWHISTORY keyword interpretation

Default value: MAXHISTORY = 5

MAXHISTORY is the maximum number of history lines to store for each numeric variable. MAXHISTORY was added because if a variable had its value changed within a large DO loop, a new history line was added each time the loop was processed, which could lead to virtual memory problems.

WRAP

Default value: WRAP = 0

If WRAP = 0, history lines and string variable contents lines are not wrapped when displayed with the SHOW command. If WRAP is non-zero, these lines are wrapped.

TENSION

Default value: TENSION = 1.0

TENSION controls the spline tension for the functions using cubic splines:

DERIV, INTEGRAL, INTERP, SMOOTH, SPLINTERP, and SPLSMOOTH.

SEED

Default value: SEED = 12345

SEED is the random number seed value. This seed is updated whenever the GENERATE\RANDOM command is entered, or the RAN is used.

POSTRES

Default value: POSTRES = 180

POSTRES controls the PostScript graphics output resolution, in dots per inch. This applies to dot filled text characters and dot types of DENSITY plots. The resolution can be changed at

any time, so different parts of a single drawing can be drawn with different resolutions.

SPEED

Default value: SPEED = 20

SPEED controls the pen plotter speed. This applies to Hewlett-Packard, Houston, and Roland RDGL || pen plotters. The speed can be changed at any time, so different parts of a single drawing can be drawn at different speeds.

WIDTH

Default value: WIDTH = 80

WIDTH controls the character width of the alphanumeric monitor screen. The value for WIDTH should be between 2 and 132.

XPREV

Default value: XPREV = 0

XPREV is the last world x-coordinate that was drawn by any graphics command. The value of this keyword is automatically updated.

YPREV

Default value: YPREV = 0

YPREV is the last world y-coordinate that was drawn by any graphics command. The value of this keyword is automatically updated.

NCURVES

Default value: NCURVES = 0

NCURVES is the total number of data curves that have been drawn, using the GRAPH command, since the last CLEAR command. The value of this keyword is automatically updated.

PCHAR

Parameters: symbol { size { colour { angle }}}

PCHAR controls the plotting symbols, or the appearance of the histogram bars, when the GRAPH command is entered. Each of the parameters may be either a literal constant, a scalar, or a vector. Entering a constant or a scalar is similar to entering a vector whose elements are all equal to the constant or the value of the scalar.

The interpretations of the parameters depend on the value of the GPLOT keyword HISTYP at the time that the GRAPH command is entered. HISTYP is also changed with the SET command. See Table 2.39 on page 118 for an explanation of HISTYP. If HISTYP = 0, the default value, subsequent graphs will be normal line graphs.

Plotting symbols

When plotting normal line graphs, the first parameter, symbol, determines the plotting symbol to draw at each data point, and whether or not to connect the symbols with line segments. Plotting symbol zero means do not draw a symbol, but just draw line segments joining the data points. This is the default.

The symbol that will be plotted at the point (x[i],y[i]) will be the ASCII decimal equivalent of the absolute value of symbol[i]. For example, 54 represents the ASCII character 6, 35 represents the ASCII character #, and 78 represents the ASCII character \$. The maximum absolute value of a plotting symbol is ninety-five (95), so the lower case alphabetic characters are not available. These symbols have their lower left corners at the data point.

There are eighteen (18) special plotting symbols which are centred at the data points using the code numbers from one (1) to eighteen (18). See Figure 2.9 on page 62. These symbols are *not* translated into their ASCII decimal equivalents. See the DISPLAY command, page 58, for information on displaying the plotting symbols on the monitor screen.

A plotting symbol value of zero means do not draw a symbol, but connect that point to the previous point. If the value of symbol[i+1] is greater than zero, the symbol drawn at point (x[i+1],y[i+1]) will be connected to the symbol drawn at point (x[i],y[i]). If the value of symbol[i+1] is less than zero, the symbol drawn at point (x[i+1],y[i+1]) will *not* be connected to the symbol drawn at point (x[i],y[i]).

Plotting symbol size

When plotting normal line graphs, the first optional parameter, size, controls the relative size of each individual plotting symbol. If size is entered, the size of the symbol at point (x[i],y[i]) will be $size[i] \times CHARSZ$. The GPLOT keyword CHARSZ can be changed with the SET command.

Plotting symbol colour

When plotting normal line graphs, the second optional vector, colour, is interpreted as the colour code for the plotting symbol. If colour is entered, the colour code of the plotting symbol at point (x[i],y[i]) will be colour[i]. Table 2.6 on page 20 shows the recognized colours and their associated colour codes.

Plotting symbol angle

When plotting normal line graphs, the third optional parameter, angle, is interpreted as the rotation angle, in degrees, of the plotting symbol. If angle is entered, the angle of the plotting symbol drawn at point (x[i],y[i]) will be angle[i]. The angle of all of the plotting symbols defaults to CHARA, which has a default value of 0° . The GPLOT keyword CHARA can be changed with the SET command.

Histograms

```
Parameters: fill { w { c }}
```

The GRAPH command used with the \HISTOGRAM qualifier plots histograms. Alternatively, SET HISTYP n where n>0, forces subsequent graphs to be histogram type graphs. See Table 2.39 on page 118 for an explanation of HISTYP.

```
\begin{array}{ll} {\tt HISTYP} = 0 & {\tt non\text{-}histogram} \\ \\ {\tt HISTYP} = 2 \ {\tt or} \ 4 & {\tt tails \ will \ be \ drawn} \\ & {\tt the \ user \ has \ control \ over \ individual \ histogram \ bar \ filling,} \\ & {\tt width, \ and \ colour} \\ \\ {\tt HISTYP} = 1 \ {\tt or} \ 3 & {\tt no \ tails \ are \ drawn} \\ & {\tt the \ user \ has \ control \ over \ individual \ histogram \ bar \ width,} \\ & {\tt and \ colour} \end{array}
```

The third optional parameter, angle, is ignored if a histogram is plotted.

Fill patterns

If HISTYP is 1 or 3, the first parameter, fill, is ignored. To fill the area under the histogram in this case, use the SET command to set the value of LINTYP to 100+ the hatch pattern number or to 200+ the dot pattern number. For example, 107 refers to hatch pattern number 7, while 244 refers to a dot fill pattern of every fourth dot both horizontally and vertically.

If HISTYP is 2 or 4, the first parameter, fill, determines the fill pattern for the individual bars of the histogram. The filling can be done with grey scales or with hatch patterns.

If $200 \le |fill[i]| \le 299$, the grey scale pattern fill[i] -200 will be used to fill the histogram bar at (x[i],y[i]). For example, if fill[i] = 234, then grey scale dot pattern 34 will be used.

```
fill[i] = 200 means every dot is lit (same as 211)
fill[i] > 0 means draw
fill[i] < 0 means erase
```

A grey scale dot pattern is of the form: uv, where the digit u is the increment number of dots to light up horizontally, $1 \le u \le 9$, and the digit v is the increment number of dots to light up vertically, $1 \le v \le 9$. For example, a grey scale dot pattern of 34 means to light up every third dot horizontally and every fourth dot vertically. If uv is negative, then the dots are erased instead of lighted. A grey scale pattern of zero, 00, is interpreted the same as pattern 11, that is, every dot is lit.

For PostScript output, set the POSTRES keyword to the appropriate resolution for your hard-copy device, using the SET command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark inkjet printer has a resolution of 360 dpi. A "good" picture can be obtained with POSTRES = 180 and LINTHK = 2.

If $100 \le |\text{symbol}[i]| \le 110$, the hatch pattern fill[i]-100 will be used to fill the histogram bar at (x[i],y[i]). For example, if fill[i] = 108, then hatch pattern 8 will be used.

| fill[i] | = | 100 | means do no fill the histogram bar |
|---------|---|-----|---|
| fill[i] | > | 0 | means the histogram bar outline will be drawn |
| fill[i] | < | 0 | means the histogram bar outline will not be drawn |

A hatch pattern is composed of an angle and one to ten spacings. These spacings are simply cycled through as each histogram bar is being filled, that is, a line is drawn inside the histogram bar at the specified angle, then a parallel line is drawn at the first spacing, then another parallel line is drawn at the second spacing, and so on for the number of spacings in that pattern. This process is repeated until the histogram bar is filled. There are ten hatch patterns available. See Figure 2.7 on page 61 for hatch pattern examples.

See the HATCH keyword for information on changing the hatch pattern definitions. See the DISPLAY command for information on how to display examples of the hatch patterns.

Histogram bar width

The first optional parameter, w, allows the user to control the widths of individual histogram bars. If w is entered, the width of the histogram bar at (x[i],y[i]) will be the normal width times w[i].

Histogram bar colour

The second optional parameter, c, allows the user to control the colours of individual histogram bars. If c is entered, the colour code for the histogram bar at (x[i],y[i]) will be c[i]. See Table 2.6, page 20, for the colour associated with each colour code.

UNITS

Parameter values: CM | IN Default value: UNITS = CM

Note: The value of the UNITS keyword is a string instead of a numeric value.

UNITS controls the plotting units type, either centimeters, CM, the default, or inches, IN.

CUNITS

Parameter values: GRAPH | PERCENT | WORLD

Default value: CUNITS = PERCENT

CUNITS controls the units type for the cursor readout when the graphics cursor is invoked by the PICK, PEAK, LINE, or FIGURE command when running under X Windows and mouse button two is pressed. If WORLD is chosen, the numbers displayed depend on the current units type, either centimeters or inches, as chosen with SET UNITS. If GRAPH is chosen, the numbers displayed depend on the current graph axis scales.

FONT

Default value: FONT = TSAN

Note: The value of the FONT keyword is a string instead of a numeric value.

FONT controls the graphics font. For a list of the font names, see Table 2.60. If just the FONT keyword is entered, a table of the font names is displayed, and you will be asked to enter a new font name. If you type the <RETURN> key, without entering anything, the font is left unchanged.

The DISPLAY FONT command will draw a font table for any font.

| STANDARD | ITALIC.2 ITALIC.2A | GOTHIC.ENGLISH GOTHIC.FRAKTUR | ROMAN.2 ROMAN.2A |
|---------------|-----------------------|-------------------------------|---------------------|
| SANSERIF.1 | ITALIC.3 | GOTHIC.ITALIAN | ROMAN.3 |
| SANSERIF.2 | | | |
| SANSERIF.CART | SCRIPT.1 | CYRILLIC.2 | OLDALPH |
| | SCRIPT.2 | | |
| HELVETICA.1 | | KATAKANA | KANJI1 |
| | GREEK.1 | HIRAGANA | KANJI2 |
| TRIUMF.1 | GREEK.2 | | KANJI3 |
| TRIUMF.2 | GREEK.2A | | KANJI4 |
| TSAN | GREEK.CART | | KANJI5 |
| | | | |
| ROMAN.FUTURA | ROMAN.SWISSL | SPECIAL | HEBREW |
| ROMAN.SERIF | ROMAN.SWISSM | MATH | |
| ROMAN.FASHON | ROMAN.SWISSB | TRIUMF.OUTLINE | |
| ROMAN.LOGO1 | | | |

Table 2.60: The font names

SHOW

```
Syntax SHOW v1 { v2 ... }

Qualifiers \text{VECTORS, \SCALARS, \MATRICES, \TEXT, \FIXED, \DUMMY, \VARY}

Examples SHOW X*

SHOW \text{VECTORS\SCALARS *ABC*}
```

The SHOW command displays, on the terminal screen, a list of:

- the current vectors, their lengths and histories
- the current scalars, their values and histories
- the current matrices, their dimensions and histories
- the current string variables, their lengths and values

If a variable name is entered, only the information about that variable is listed. Wildcards,

 \ast , are allowed in the name, and then all variables that match will be listed.

If the \VECTORS qualifier is used, only the vector variables will be listed.

If the \SCALARS qualifier is used, only the scalar variables will be listed.

If the \MATRICES qualifier is used, only the matrix variables will be listed.

If the \TEXT qualifier is used, only the string variables will be listed.

The qualifiers FIXED, DUMMY, and VARY are *only* valid when used with the \SCALARS qualifier.

```
\begin{tabular}{lll} $ \line & - FIXED & \longrightarrow & do not show scalars that are fixed (with respect to fitting) \\ $ \line & - DUMMY & \longrightarrow & do not show scalars that are dummy variables \\ $ \line & - VARY & \longrightarrow & do not show scalars allowed to vary in a fit \\ \end{tabular}
```

Combinations are allowed, for example, \-FIXED\-DUMMY means only show scalars that are allowed to vary in a fit.

Ordered Vectors

All vectors now have an order property. Vectors are either in ascending order, descending order, or un-ordered. The type is displayed in the SHOW command, where +0 means ascending order, -0 means descending order, and no symbol means un-ordered. For now, being ordered only has an affect on the vector union, /|, and the vector intersection, /&. These operations are much faster if the vector operands are ordered. The WHERE function produces an ascending order vector, as does the SORT\UP command. The SORT\DOWN command produces a descending order vector. This new vector property will be utilised more in the future to enhance speed and efficiency.

Examples

Suppose you have defined the following variables:

```
vectors:
                    Х
                         XX
                              AX
                                    XB
scalars:
                   SX
                              S_1
                        XSX
matrices:
                   M1
                        MX1
string variables:
                   T1
                         T2
                              ST
                                   TXT
SHOW *X
```

```
SHOW *X the information about X, XX, AX, SX and XSX will be displayed

SHOW X* the information about X, XX, XB and XSX will be displayed

SHOW V X* the information about X, XX and XB will be displayed

SHOW X the information about X only will be displayed

SHOW *X* the information about X only will be displayed

SHOW *X* the information about X, XX, AX, XB, SX, XSX, MX1, and TXT will be displayed

SHOW T *X* the information about TXT will be displayed
```

SLICES

```
Syntax SLICES n x y z { a }

Defaults a = 30°

Examples SLICES 5 X Y Z

SLICES 10 X Y Z 45
```

The SLICES command bins the x, y, and z vectors into n slices, and produces a graphical representation of this binned data. The x, y and z vectors are assumed to represent scattered data points (x[i],y[i]) with altitude z[i]. The z-axis is drawn at an angle, a, in degrees. a defaults to 30° .

This command is open for suggestions.

Example

The following script produced Figure 2.25.

```
X=COS([-100:100:.5]) ! generate some "data"
Y=SIN([10:20:.025]) !
Z=SIN(X)*COS(Y) !
SET PCHAR -16 ! set plotting character to a point
SLICES 5 X Y Z 35 !
```

SORT

```
Syntax SORT x { x1 x2 ... }

Qualifiers \UP, \DOWN

Defaults \UP

Examples SORT X Y

SORT\DOWN X Y Z

SORT\UP X Y Z
```

The SORT command is used for sorting vectors into ascending or descending order. By default, the vector \mathbf{x} is sorted into ascending order. To sort vector \mathbf{x} into ascending order, it is not necessary to use the \UP qualifier. Ascending order means that element 1 will be the smallest element. To sort vector \mathbf{x} into descending order, use the \DOWN qualifier. Descending order means that element 1 will be the largest. Vector \mathbf{x} will be altered.

Associated vectors

If other vectors, xI, are entered, they will not be sorted. They will be re-arranged in the same

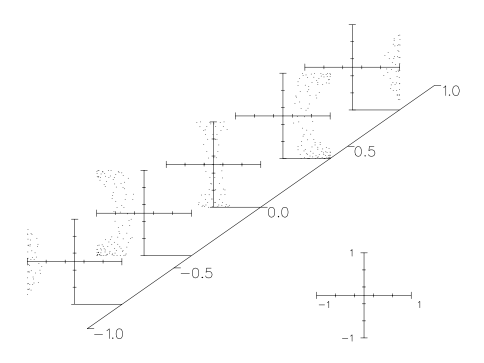


Figure 2.25: An example using the SLICES command

way that x is re-arranged, that is, if element x[i] becomes x[j] because of sorting then xI[i] will become xI[j]. This feature was included so that if data vectors are associated with each other they can be sorted without the associations being lost.

Examples

Suppose you have three data vectors, X, Y, and Z, which represent rectangular coordinates and an associated altitude:

```
X=[1;2;3;4;5]
Y=[10;8;6;4;2]
Z=[-0.3;-1.0;-0.5;2.0;-2.0]
```

If you want to sort Z into descending order, without breaking up the triplets, enter:

SORT\DOWN Z X Y

However, if you want to sort Z into descending order, and are not concerned about keeping the Y and Z associations, simply enter:

```
SORT\DOWN Z
```

Suppose you now want to re-sort Y data into ascending order, and you are concerned about keeping the X and Z associations. Enter:

```
SORT Y X Z
```

```
X=[5;4;3;2;1]
Y=[2;4;6;8;10]
```

after which:

Z=[-2.0;2.0;-0.5;-1.0;-0.3]

STACK

Syntax STACK filename

Qualifiers \APPEND, \EXECUTE

Defaults \NOAPPEND, \EXECUTE

Examples STACK FILE.STK

STACK\APPEND FILE.STK STACK\NOEXE FILE.STK

A stack file will contain all the subsequently entered commands. The STACK command is a way of interactively creating a command script file, and is meant to be used in conjunction with the EXECUTE command, page 76. Commands that are being stacked in a file can be executed later.

To turn off the stacking feature, use the DISABLE STACK command, after which commands will no longer be written to the previously named stack file. To re-enable the same stack file, use the command ENABLE STACK, after which subsequent commands will be appended to that file.

Environment variables in file names

For UNIX users, it is now possible to use an environment variable in a file name, if the environment variable is preceded by a \$. For example,

```
setenv FILE mysession
physica
stack $FILE
```

The environment variable can be just the first part of the filename, for example,

```
setenv FILE my
```

```
physica
stack $FILEsession
```

Appending to a stack file

The default, \NOAPPEND , is to open a new file. If the \APPEND qualifier is used, then subsequent commands will be appended to the stack file.

Executing commands while stacking

The default, $\$ is to execute commands normally. If the $\$ NOEXECUTE qualifier is used, then subsequently entered commands will be written to the file, but will *not* be executed at that time.

STATISTICS

```
Syntax STATISTICS x { s1\keyword { s2\keyword ... }}
STATISTICS\MOMENT w x n { sout }
STATISTICS\PEARSON x y rcof prob

Qualifiers \MESSAGES, \WEIGHTS, \MOMENT, \PEARSON

Defaults \MESSAGES, \NOWEIGHTS, \NOMOMENT, \NOPEARSON

Examples STATISTICS X
STATISTICS\NOMESS X XMED\MEDIAN XMEAN\MEAN
STATISTICS\WEIGHTS W X XVAR\VARIANCE XSUM\SUM
STATISTICS\MOMENT Y X 3 M3
```

The STATISTICS command calculates¹ various statistics for the input variable x, which can be a vector or a matrix. Specific statistics are chosen with qualifier keywords which are appended to the output parameters with the backslash, \setminus .

Table 2.61 shows the parameter qualifier keywords and corresponding output values for extrema. Table 2.62 shows the parameter qualifier keywords and corresponding output values for central measures. Table 2.63 shows the parameter qualifier keywords and corresponding output values for dispersion and skewness.

Informational messages

The default is to display all the calculated statistics on the terminal screen. If the \NOMESSAGES command qualifier is used, and if at least one output scalar is entered, then the values of

¹The definitions used here are taken from "Numerical Recipes – The Art of Scientific Computing" by W.H. Press, B.P. Flannery, S.A. Teukolsky, W.T. Vetterling, Cambridge University Press, 1986.

| keyword | output value |
|----------------------------|--|
| \MAX | maximum value of x |
| \setminus IMAX | index of the maximum if x is a vector |
| | row index of the maximum if x is a matrix |
| $\backslash \texttt{JMAX}$ | column index of the maximum if x is a matrix |
| \MIN | minimum value of x |
| \IMIN | index of the minimum if x is a vector |
| | row index of the minimum if x is a matrix |
| $\backslash \mathtt{JMIN}$ | column index of the minimum if x is a matrix |

Table 2.61: The STATISTICS command extrema keywords

| keyword | output value |
|---------|-----------------------------|
| \SUM | arithmetic sum (unweighted) |
| \MEAN | arithmetic mean |
| \GMEAN | geometric mean |
| \MEDIAN | median value |
| \RMS | root-mean-square |

Table 2.62: The STATISTICS command central measure keywords

| keyword | output value |
|-----------|--------------------|
| VARIANCE | variance |
| \SDEV | standard deviation |
| \ADEV | average deviation |
| \KURTOSIS | kurtosis |
| \SKEWNESS | skewness |

Table 2.63: The STATISTICS command dispersion and skewness keywords

all the statistics will not be displayed on the terminal screen.

Weights

```
Syntax \qquad {\tt STATISTICS} \\ {\tt WEIGHTS w x { s1\keyword { s2\keyword ... }}}
```

You *must* use the \WEIGHTS qualifier to indicate that a weight vector is present. Weights cannot be applied to matrix data.

A weighting factor, $w_i \ge 0$, could be the frequency, the probability, the mass, the reliability, or some other multiplier. The minimum of the lengths of w and x will be used. If the lengths are different, a warning message to that effect will be displayed on the terminal screen.

Definitions

Suppose that x is a vector with N elements.

If a weight vector, w, is entered, remember to use the \WEIGHTS command qualifier. The length of w is assumed to also be N. If no weight is entered, let w_i default to 1, for $i=1,\ldots,N$. Define the total weight: $W=\sum_{j=1}^N w_j$

Sum

The sum is defined by $\sum_{j=1}^{N} x_j$

Mean value

The mean value, \bar{x} , is defined by $\bar{\mathbf{x}} = \frac{1}{W} \sum_{j=1}^{N} w_j x_j$

Geometric mean

The geometric mean, \mathcal{G}_x , is defined, if each $x_i \geq 0$, by: $\mathcal{G}_x = e^{\frac{1}{W}\sum_{j=1}^N w_j \log(x_j)}$

Median

The median is the element of x which has equal numbers of values above it and below it. If N is even, the median is the average of the unique two central values.

Root-mean-square

The root-mean-square, \mathcal{RMS} , is defined by $\mathcal{RMS} = \sqrt{\frac{1}{W} \sum_{j=1}^{N} w_j x_j^2}$

Variance

The variance, μ , is defined by $\mu = \frac{N}{W(N-1)} \sum_{j=1}^{N} w_j (x_j - \bar{x})^2$

Standard deviation

The standard deviation, σ , is defined by $\sigma = \sqrt{\mu}$

Average deviation

The average deviation, or mean deviation, δ , is defined by $\delta = \sum_{j=1}^N w_j |x_j - \bar{x}|/W$

Skewness

The skewness, or third moment, skew, is a nondimensional quantity that characterizes the degree of asymmetry of a distribution around its mean. The skewness is a pure number that characterizes only the shape of the distribution, and is defined by

$$skew = \sum_{j=1}^{N} w_j \left[\frac{x_j - \bar{x}}{\sigma} \right]^3 / W$$

A positive value of skewness signifies a distribution with an asymmetric tail extending out towards more positive x; a negative value signifies a distribution whose tail extends out towards more negative x.

Kurtosis

The kurtosis, kurt, is a nondimensional quantity which measures the relative peakedness or flatness of a distribution, relative to a normal distribution. A distribution with positive kurtosis is termed leptokurtic; a distribution with negative kurtosis is termed platykurtic. An in-between distribution is termed mesokurtic. The kurtosis is defined by

$$kurt = (\sum_{j=1}^{N} w_j [\frac{x_j - \bar{x}}{\sigma}]^4) - 3$$

where the -3 term makes the value zero for a normal distribution.

Moments

 $\textbf{\textit{SYNTAX}} \qquad \texttt{STATISTICS} \backslash \texttt{MOMENT w x n } \{ \texttt{ s } \}$

If the \model moment command qualifier is used, the \mathbf{n}^{th} moment of vector \mathbf{x} , with weight \mathbf{w} , is calculated and optionally stored in output scalar \mathbf{s} . The moment number, \mathbf{n} , can be any integer >0.

s=S/W, where: $S=\sum_{i=1}^N w_i \times x_i^n$ and $W=\sum_{i=1}^N w_i$ and where N is the length of x and w.

If x and w have different lengths, a warning message to that effect will be displayed, and the minimum of the lengths of x and w will be used. If W = 0, then an error message is displayed and s is returned as 0.

Linear correlation coefficient

Pearson's r, or the linear correlation coefficient, is widely used as a measure of association between variables that are continuous. For pairs of quantities (x_i, y_i) , for $i = 1, \ldots, N$, the linear correlation coefficient r is given by the formula:

$$r = \frac{\sum_{i=1}^{N} (x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum_{i=1}^{N} (x_i - \bar{x})^2} \sqrt{\sum_{i=1}^{N} (y_i - \bar{y})^2}}$$

where \bar{x} is the mean of the x_i 's, and \bar{y} is the mean of the y_i 's.

The value of r lies between -1 and +1, inclusive. It takes on a value of +1 when the data points lie on a straight line with positive slope, x and y increase together. The value +1 holds independent of the magnitude of this slope. If the data points lie on a straight line with negative slope, y decreases as x increases, then r has the value -1. A value of r near zero indicates that the variables x and y are uncorrelated.

r is a way of summarizing the strength of a correlation which is known to be significant, but it is a poor statistic for deciding whether an observed correlation is statistically significant, and/or whether one observed correlation is significantly stronger than another. The reason is that r is ignorant of the individual distributions of x and y, so there is no universal way to compute its distribution in the case of the null hypothesis.

The STATISTICS\PEARSON command returns Pearson's r in the scalar variable r. It also returns scalar p, the significance level at which the null hypothesis of zero correlation is disproved. A small value of p indicates a significant correlation.

$$p = I_{\frac{N-2}{N-2+t^2}}(\frac{N-2}{2}, \frac{1}{2})$$

where I is the incomplete Beta function and t is defined by: $t = r\sqrt{\frac{N-2}{1-r^2}}$

Examples

Suppose you have a vector X=[1.2;2.1;3.2;4.5;5;6;7]. Entering STATISTICS X produces the following display:

```
7
Minimum =
                     1.2
                                       | Maximum =
Index of minimum = [1]
                                       | Index of maximum = [7]
Mean =
                     4.142857
                                       | Geometric mean =
                                                             4.13629
                                                             4.36619
Median not requested
                                       | Variance =
Standard deviation = 2.089543
                                       | Average deviation = 1.693878
                                       | Kurtosis =
Skewness =
                    -0.0696135
                                                             -1.708838
```

If you want to use the values for the maximum, minimum and mean of X, enter:

STATISTICS X XMEAN\MEAN XMIN\MIN XMAX\MAX

and you will have the scalars: XMAX=7, XMIN=1.2, and XMEAN=4.142857

If you also want the index values for the maximum and the minimum of X, enter:

STATISTICS X XMEAN\MEAN XMIN\MIN XMAX\MAX IMX\IMAX IMN\IMIN

and you will also have scalars: IMX=7 and IMN=1.

STATUS

Syntax STATUS

The STATUS command displays on the terminal screen most of the internal PHYSICA flags and settings. Example output from the STATUS command, is shown below.

```
PHYSICA version number:
                         2.10
       version date: January 16, 1998
Echoing
                 disabled | Confirm
                                               enabled
Prompting
                            Recall shell
                  enabled |
                                               enabled
Border
                  enabled |
                            Replot
                                               enabled
                             Autoscale
X replay
                  enabled |
                                          X and Y axes
History recording enabled |
Journaling enabled, file= PHYSICA.JOURNAL
                   macro journaling disabled
Stack
         disabled
Macro file extension default: pcm
Graphics hardcopy device: (bitmap) HPLASERJET 150 dpi
Colour: WHITE
                 (1)
World units type: CM
Window
         0 coordinates:
                          0.00
                                 0.00 100.00 100.00 percentages
                          0.00
                                 0.00 27.94 21.59 centimeters
Window
         0 coordinates:
                          0.00
                                 0.00 100.00 100.00 percentages
                                 0.00 27.94 21.59 centimeters
                          0.00
```

SURFACE

```
Syntax SURFACE m { xangle { zangle { vsf { psf }}}}

Qualifiers \COLOUR, \NORMAL, \HISTOGRAM, \XZLINES, \XLINES, \ZLINES

Defaults \NOCOLOUR, \NORMAL, \XZLINES

xangl = -45°, zangl = -45°, vsf = 1, psf = 0

Examples SURFACE M

SURFACE\COLOUR M

SURFACE\HIST M -30 -60

SURFACE\NORMAL\ZLINES M -45 -45 2
```

The SURFACE command displays a matrix in the form of a 3-dimensional figure which can be rotated in space. Figure 2.26 shows the coordinate system used in this command. The appearance of the figure is controlled by optional qualifiers and parameters.

Note: Unlike most other commands, once one of the optional parameters has been changed to a new value, that value becomes the default.

The qualifiers that control the appearance of the figure are:

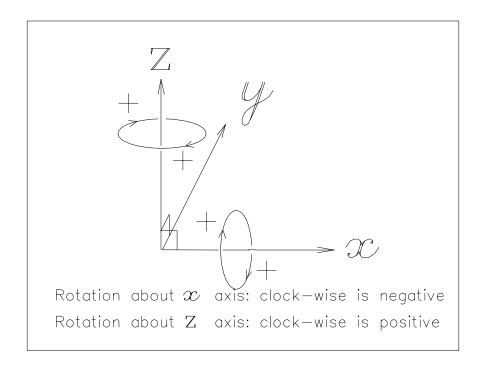


Figure 2.26: Surface coordinate system

| qualifier | results |
|------------|---|
| \NORMAL | a normal surface plot (default) |
| \HISTOGRAM | a histogram surface |
| | |
| \XZLINES | lines are drawn parallel to the x - and the z -axis (default) |
| \XLINES | lines are drawn parallel to the x-axis only |
| \ZLINES | lines are drawn parallel to the z -axis only |

The parameters that control the appearance of the figure are:

| parameter | results |
|-----------|---|
| xangl | rotation angle about the x -axis (default -45°) |
| zangl | rotation angle about the z -axis (default -45°) |
| vsf | vertical scale factor (default 1) |
| psf | pedestal size scale factor (default 0) |

Colour

If the \COLOUR qualifier is used, the height of the surface will be divided into seven (7) regions

and a different colour associated with each region. This allows the user to quickly identify areas of similar height.

Examples

The following script produced Figure 2.27.

```
R=MOD([0:143],4)+1
SORT\UP R
T=MOD([0:143],36)*10
GRID R*COSD(T) R*SIND(T) EXP(-R/2)*COSD(180*(R-1)) M
SURFACE M -15 -45 1 1.5
```

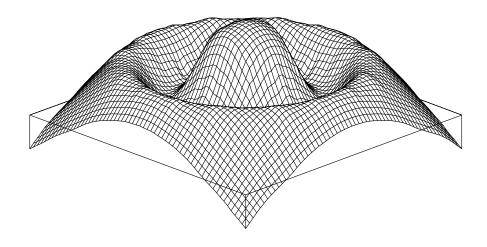


Figure 2.27: A SURFACE example

TERMINAL

If the TERMINAL command is encountered in a script file that is being executed, control is passed back to the terminal keyboard. Commands can then be entered interactively. To continue execution of the script file, type the <RETURN> key without entering anything. The script file will then continue execution with the command immediately following the TERMINAL command.

By default, the message 'type <RETURN> to continue' will be displayed. You can specify the message by entering a string with the command.

TEXT

```
Syntax TEXT { 'string' }
Qualifiers \GRAPH, \ERASE, \CONFIRM

Defaults \CONFIRM, \NOGRAPH, \NOERASE

Examples TEXT\NOCONFIRM 'This is a text string'
    TEXT 'The value of A is '//RCHAR(A)//' and B = '//RCHAR(B)
    TEXT '<alpha,^>2<_,gamma>=<Phi>'
    TEXT TIME//' '//DATE
    TEXT\ERASE\NOCONFIRM
```

By default, the TEXT command draws strings. The 'string' defaults to the last string that was entered with the TEXT command. Use the SET FONT command to set the font. The default font is TSAN.

Confirmation

If the TEXT command is entered interactively, the default is to request confirmation as to whether the text is acceptable as drawn. If you answer NO, the text will not appear on any hardcopies, or in an EDGR file. The text will not be entered into the hardcopy plot file, or into an open EDGR file, unless confirmed.

If the DISABLE CONFIRM command is entered before the TEXT command, the default will be that no such confirmation will be requested. The initial default is for CONFIRM to be enabled.

If the \backslash CONFIRM qualifier is used, the default, confirmation will be forced. If the \backslash NOCONFIRM qualifier is used, confirmation will be suppressed.

If the TEXT command is encountered in a script file, no confirmation is requested, even if ENABLE CONFIRM has been entered and the \backslash CONFIRM qualifier is used.

Stack files

If a stack file is open, via the STACK command, page 247, then the (x,y) coordinates of the text location will be written to the stack file, even if they are chosen by the graphics cursor. When the stack file is executed, using the EXECUTE command, page 76, the graphics cursor will not be used.

If confirmation is requested and the figure is not acceptable, then the command will *not* be written to the stack file.

Text characteristics

The font can be changed with the SET FONT command, page 228. The default font is TSAN.

The text height can be controlled with the keywords TXTHIT or %TXTHIT, using the SET command. The default is %TXTHIT = 3.

The text angle can be controlled with the keyword TXTANG, , using the SET command. The default is TXTANG = 0.

Justification and location

The justification of the text is controlled by the keyword CURSOR. See Table 2.64 and Table 2.65. The origin of the text is always the lower left corner of the string. The justification determines where this origin is placed with respect to a reference point, see Figure 2.28. The default value of CURSOR is ± 1 .

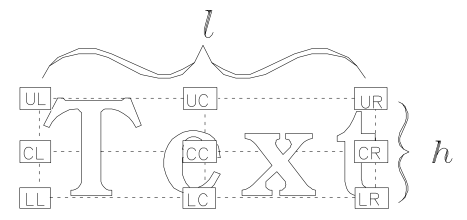


Figure 2.28: Text extent rectangle with two-character justification codes

If $CURSOR \le 0$, XLOC and YLOC will be used to determine the reference point for the text, and the justification is determined by the value of CURSOR.

If CURSOR > 0, the reference point is determined with the graphics cursor. A menu is available by typing key M, see Table 2.65. The justification of the text may be selected by typing key J and then selecting a justification, CC, LU, etc., or by simply typing one of the special justification keys, R, L, C, U, D, or V. No carriage return is needed after these special keys.

The value of CURSOR will be updated to the value corresponding to the justification that was chosen. For example, if CURSOR is currently 5 and the user chooses the justification code CV, then the value of CURSOR is changed to 6. The values of %XLOC and %YLOC will also be updated to the graphics cursor position that was chosen.

Text Formats

| | | Jus | tification | |
|---------------|----------------------|--|---|--------------------------|
| | (x_0, y_0) | = origin point of string (lower left corner) | | |
| | (x_{ref}, y_{ref}) | = reference poi | $\mathbf{nt} \; (x_{ref} = \mathtt{XLOC}, \mathbf{y})$ | $y_{ref} = 	exttt{YLOC}$ |
| CURSOR | h | = maximum he | eight of string (d | efault = TXTHIT) |
| | l | = length of stri | ng in world cooi | rdinates |
| | a | = angle of strin | g in degrees (de | efault = TXTANG) |
| > 0.0 | | See the fo | ollowing table | |
| 0.0 or -1.0 | LL | $x_0 = x_{ref}$ | $y_0 = y_{ref}$ | |
| -2.0 | LC | $x_0 = x_{ref} + l/2$ | $y_0 = y_{ref}$ | |
| -3.0 | LR | $x_0 = x_{ref} + l$ | $y_0 = y_{ref}$ | |
| -4.0 | LU | $x_0 = x_{ref}$ | $y_0 = y_{ref}$ | $a = 90^{\circ}$ |
| -5.0 | LD | $x_0 = x_{ref}$ | $y_0 = y_{ref}$ | $a = 270^{\circ}$ |
| -6.0 | CV | $x_0 = x_{ref} + l/2$ | $y_0 = y_{ref}$ | $a = 90^{\circ}$ |
| -7.0 | CL | $x_0 = x_{ref}$ | $y_0 = y_{ref} + h/2$ | |
| -8.0 | UL | $x_0 = x_{ref}$ | $y_0 = y_{ref} + h$ | |
| -9.0 | CC | $x_0 = x_{ref} + l/2$ | $y_0 = y_{ref} + h/2$ | |
| -10.0 | UC | $x_0 = x_{ref} + l/2$ | $y_0 = y_{ref} + h$ | |
| -11.0 | CR | $x_0 = x_{ref} + l$ | $y_0 = y_{ref} + h/2$ | |
| -12.0 | UR | $x_0 = x_{ref} + l$ | $y_0 = y_{ref} + h$ | |

Table 2.64: Text justification interaction with ${\tt CURSOR}$

| key | justification with respect to reference point at crosshair location | | | |
|-------|--|------------------------|--------------------|--|
| typed | when $\mathtt{CURSOR} > 0.0$ | | | |
| М | display the menu | | | |
| Q | quit, do not draw any text | | | |
| / | clear the alpha-numeric terminal screen, but not the graphics | | | |
| J | a menu of justifications will be displayed. To draw the text string, choose | | | |
| | one of the following two-character codes and type the RETURN key | | | |
| | | | | |
| | Two character code | Justificatio | | |
| | LL | lower left | (CURSOR set to 1) | |
| | CL | centre left | (CURSOR set to 7) | |
| | UL | upper left | (CURSOR set to 8) | |
| | LC | lower centre | (CURSOR set to 2) | |
| | CC | center centre | (CURSOR set to 9) | |
| | UC | upper centre | (CURSOR set to 10) | |
| | LR | lower right | (CURSOR set to 3) | |
| | CR | centre right | (CURSOR set to 11) | |
| | UR | upper right | (CURSOR set to 12) | |
| | LU | lower left at 90° | (CURSOR set to 4) | |
| | LD | lower left at 270° | (CURSOR set to 5) | |
| | CV | lower centre at 90° | (CURSOR set to 6) | |
| | | | | |
| L | lower left (LL) | | | |
| С | lower centre (LC) | | | |
| R | lower right (LR) | | | |
| U | lower left with an angle of 90° (LU) | | | |
| D | lower left with an angle of 270° (LD) | | | |
| V | lower centre with an angle of 90° (CV) | | | |
| X | lower left (\mathtt{LL}); using the y location selected by the crosshair | | | |
| | and the x location that is stored in <code>XLOC</code> | | | |
| Υ | lower left (\mathtt{LL}); using the x location selected by the crosshair | | | |
| | and the y location that is stored in YLOC | | | |
| other | use current value of CURSOR for justification | | | |
| | use the crosshair position | on for the reference p | oint | |

Table 2.65: Text menu and justification

The string may contain format commands and special characters which are included inside the string, and must be bracketed by the command delimiters, < and >. The special characters include all of the greek letters as well as some math symbols and other symbols. The names for the special characters can be seen by entering the DISPLAY SPECIAL command, page 58. See Figure 2.6 on page 61. The format commands are listed in Table 2.66.

| bolding | <bn></bn> |
|--------------------|--------------------------------------|
| colour | <cn></cn> |
| font | <ffontname></ffontname> |
| height | <Hnn.n> or $<$ Hnn.n%> |
| sub-scripts | <_> |
| super-scripts | <^>> |
| emphasis | |
| hexadecimal input | <x></x> |
| vertical spacing | $\forall nn.n > or \forall nn.n >$ |
| horizontal spacing | <znn.n> or <znn.n%></znn.n%></znn.n> |

Table 2.66: TEXT command text formatting commands

When <NOD> is included in a string, the bolding, colour, font, emphasis and hexadecimal mode will be left in their current state. This is a way of changing the defaults for these characteristics.

When <DEF> is included in a string, at the end of processing that string, bolding will be turned off, the colour will be reset to the colour chosen by the COLOUR command, the font will be reset to the font chosen by the SET FONT command, slant mode will be turned off, and hexadecimal mode will be turned off. This is the default action, so it is not necessary to include <DEF> in a string. It has been included for completeness.

Refer to Appendix A for explanations of CURSOR, XLOC, YLOC, TXTHIT, TXTANG.

Replotting text

When text strings are added to a graph, they will be replotted, along with any data curves, with the REPLOT command, page 208. The default is to store the text locations in world coordinates, that is, centimeters or inches.

If you want the text locations to be recorded in graph units, use the \GRAPH qualifier. In this case, the text will be replotted in the same location in terms of the graph, so that if you label a curve, the label will follow the curve on the new graph scales. However, the height of the text will be increased or decreased proportionally. To restore the text height after a REPLOT, you can get, using the GET command, the value of %TXTHIT before REPLOT and re-set %TXTHIT

after the REPLOT, using the SET command.

Drawing the Date and Time

The DATE function returns the current date as a string. The TIME function returns the current time as a string. For example:

```
=DATE
'16-APR-1993'
=TIME
'11:58:16'
```

So the current date and time could be drawn with: TEXT TIME//' '//DATE

Erasing text

Text can only be erased from the monitor screen and from a bitmap. Erasing will *not* work with plotter files and it will *not* work with EDGR. Erasing of rectangular regions is possible with PostScript graphics using the ERASEWINDOW command, page 76.

If the \ERASE qualifier is used, the last text drawn will be erased. If the TEXT \ERASE command is issued again, the next to last text drawn will be erased. And so on.

If the \NOCONFIRM qualifier is used, no confirmation will be requested. If the \CONFIRM qualifier is used, the string to be erased will be displayed on the screen, and you will be asked if this is the string to be erased. If the answer is NO, then the next string in reverse order of drawing is displayed, and so on. The default is \CONFIRM.

Example

The following script produced Figure 2.29.

```
%XLOC 50  ! x reference location
%YLOC 80  ! y reference location
CURSOR -2  ! lower centre justify at (x,y) reference location
! preceding blank line is necessary
TEXT 'This is a simple text string'
SET %YLOC 65
TEXT '<h10%,fscript.2>He<h8%>igh<h5%>t, font <alpha,beta,aleph>'
SET %YLOC 50
TEXT '<B22,froman.serif,h5%>DOT FILLED CHARACTERS'
SET %YLOC 35
TEXT '<INTEGRL>xdx = x<^>2<_>/2'
```

This is a simple text string

Height, font $\alpha\beta$ ×

DOT FILLED CHARACTERS

$$\int x dx = x^2/2$$

Figure 2.29: An example using the TEXT command

TILE

Syntax TILE file

The TILE command draws complicated bar graphs. The numbers which define the rectangular bars are read from the file. Strings may also be drawn. Comment lines are ignored, where a comment line is any line that begins with a !.

Bar definitions

A bar is defined by the following values:

| parameter | result |
|-----------|---|
| xlower | the x coordinate of the left edge of the bar, in graph units |
| xupper | the x coordinate of the right edge of the bar, in graph units |
| ymid | the y coordinate of the middle of the bar, in graph units |
| nh | the dot or hatch pattern with which to fill the bar |
| | $ \mathtt{nh} \geq 11$ then fill with grey scale dot pattern |
| | $1 \le \mathtt{nh} \le 10$ then fill with hatch pattern |
| | $ \mathtt{nh} = 0$ then no filling |
| | $\mathtt{nh} < 0$ then the outline of the bar is not drawn, |
| | but hatch pattern nh is drawn |
| height | the height of the bar, in graph units |
| colour | the colour of the bar outline and fill pattern |

Hatch patterns

A hatch pattern is composed of an angle and one to ten spacings. These spacings are simply cycled through as the region is being filled, that is, a line is drawn inside the region at the specified angle, then a parallel line is drawn at the first spacing, then another parallel line is drawn at the second spacing, and so on for the number of spacings in that pattern. This process is repeated until the region is filled. The hatch patterns can be redefined with the SET HATCH command and displayed with the DISPLAY FILL command. There are ten hatch patterns available.

Dot fill patterns

A dot pattern is of the form: uv, where the digit u is the increment number of dots to light up horizontally, $1 \le u \le 9$, and the digit v is the increment number of dots to light up vertically, $1 \le v \le 9$. For example, a dot pattern of 34 means to light up every third dot horizontally and every fourth dot vertically. If uv is negative, then the dots are erased instead of turned on.

PostScript output

For PostScript output, set the POSTRES keyword to the appropriate resolution for your hard-copy device, using the SET command. Use a combination of line thickness and resolution for a quicker resulting picture. For example, the Lexmark inkjet printer has a resolution of 360 dpi. A "good" picture can be obtained with POSTRES = 180 and LINTHK = 2.

String definitions

A string is defined by the following values:

| parameter | result |
|-----------|---|
| xlower | the x coordinate of the lower left corner of the string, in graph units |
| just | the justification to use for positioning the text, |
| | $	t L 	o 	extbf{left}$, $	t C 	o 	extbf{centre}$, $	t R 	o 	extbf{right}$ |
| ymid | the y coordinate of the middle of the text |
| font | the name of the font to use for drawing the text |
| height | the height of the text, in graph units |
| colour | the colour of the text $(1-8)$, |
| | see the COLOUR command for an explanation of the colour code |
| 'text' | the text itself, enclosed in quotes |

Example

Suppose you have a file, TILE.DAT, as below:

```
!
!xlo xup ymid nfill height colour
 0
      4
           1
                  2
                        .8
                                1
 5
     10
                        .8
                                2
           1
                 33
 0
     3
           2
                        .8
                                3
                 44
 4
     6.5 2
                 55
                        .8
                                4
 7
           2
     9
                        .8
                                5
                 10
 0
      2
           3
                  6
                        .8
                                2
 3
      5
           3
                 8
                        .8
                                3
                                5
 6
      8
           3
                        .8
                 66
 8.5 10
           3
                 77
                        .8
                                1
!xlo just ymid font height colour text
 -2
      R
            2
                TSAN
                            .3
                                    1 'Second label'
 -2
            1
                                        'First label'
      R
                TRIUMF.2
                            .3
 -2
      R
            3
                ROMAN.SERIF .3
                                    1 'Third label'
```

The following script produced Figure 2.30.

SCALES -10 12 11 0 4 4 ! manually set axes scales

TILE TILE.DAT ! draw the tiles

SET

BOX 0 ! turn off axis box YAXIS 0 ! turn off y-axis

NSXINC 2 ! number of small x increments = 2

GRAPH\AXES ! draw axes only

SET

XAXIS 0 ! turn off x-axis
YAXIS 1 ! turn on y-axis

YITICA -90 ! draw numbers on right side of y-axis
YTICA -90 ! draw tics on right side of y-axis
%XLAXIS 95 ! draw y-axis at right end of x-axis

GRAPH\AXES ! draw axes only

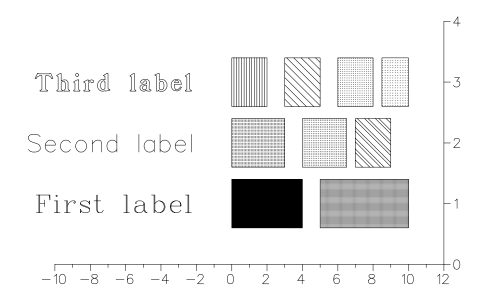


Figure 2.30: An example using the TILE command

TLEN

Syntax TLEN txt lenout

The TLEN command returns a scalar, lenout, which is the number of string elements of the array string variable, txt.

Example

```
Suppose that: T[1]='string 1'; T[2]='string 2'; T[3]='string 3' then TLEN T L would return the scalar L = 3.
```

TRANSFORM

```
Syntax TRANSFORM x p f(x,p) y

Example TRANSFORM X P EXP(-P*X^2) XT
```

The TRANSFORM command performs integral transforms of the function f(x,p). The function is integrated over the entire range of definition, as defined by the independent variable, vector x. The integral is evaluated for each value of the parameter vector p, with the result stored in the vector y.

$$y_i = \int f(x, p_i) dx$$

The function f(x,p), which forms the integrand of the transform integral, may contain other scalars, or other vectors dimensioned identically to x. Such vectors are considered, and integrated, as functions of the independent variable x. Vectors of mismatched size will generate an error.

This command can be used to perform standard transforms such as Laplace or Hankel transforms. Fourier transforms should be done using the FFT function, as this is much faster.

Example

The following script integrates the expression e^{-px^2} over x = [1:20] for each p = [1:3], and stores the results in output vector F.

```
X=[1:20]
P=[1:3]
TRANSFORM X P EXP(-P*X^2) F
```

after which: F=[0.1601404;0.05141661;0.01877862]

A check on this result is the following:

```
FI[1] = (INTEGRAL(X, EXP(-P[1] * X^2))) [LEN(X)]

FI[2] = (INTEGRAL(X, EXP(-P[2] * X^2))) [LEN(X)]

FI[3] = (INTEGRAL(X, EXP(-P[3] * X^2))) [LEN(X)]
```

after which: FI=[0.1601404;0.05141661;0.01877862]

UNIQUE

```
Syntax UNIQUE x y xout yout
UNIQUE\INDICES x y idx { (index_expression) }

Qualifiers \MESSAGES, \INDICES

Defaults \MESSAGES, \NOINDICES
```

The UNIQUE command eliminates adjacent duplicate points in the vectors x and y. A duplicate point means that x[i] = x[i+1] and y[i] = y[i+1].

The vectors xout and yout are created, and will contain no adjacent duplicate points. If none are found, xout = x and yout = y.

To disable informational messages from the UNIQUE command, use the \NOMESSAGES qualifier.

Indices

```
Syntax UNIQUE\INDICES x y idx { (index_expression) }
```

The UNIQUE\INDICES command finds adjacent duplicate points in the vectors x and y. A duplicate point means that x[i] = x[i+1] and y[i] = y[i+1]. The vector idx is created, and will contain the indices of x and y where there are no adjacent duplicate points. If none are found, idx will not be made.

If you wish to search a subset of x and y, use the optional parameter (index_expression). Enclosing the index_expression in parenthesis allows you to have blanks within the expression. The index_expression can be a simple range, for example, [1:10], or a mathematical expression, for example, where(z=n). The index_expression can be used directly on the input vectors x and y, but then the output idx index vector will contain indices relative to [1:len(x[index_expression])].

Examples

Suppose that:

```
X=[1;2;2;3;4;5;5;7;8;9;9;10]
Y=[-6;-5;-4;-3;-2;-1;-1;1;2;3;3;4]
```

After the command: UNIQUE X Y XO YO

```
X= [ 1; 2; 2; 3; 4; 5; 5;7;8;9;9;10]

Y= [-6;-5;-4;-3;-2;-1;-1;1;2;3;3; 4]

X0=[ 1; 2; 2; 3; 4; 5; 7;8;9; 10]

Y0=[-6;-5;-4;-3;-2;-1; 1;2;3; 4]
```

After the command: UNIQUE\INDICES X Y IXY

```
X= [ 1; 2; 2; 3; 4; 5; 5;7;8; 9;9;10]
Y= [-6;-5;-4;-3;-2;-1;-1;1;2; 3;3; 4]
IXY=[ 1; 2; 3; 4; 5; 6; 8;9;10; 12]
```

Suppose that:

After the command: UNIQUE\INDICES X Y IDX

```
IDX = [1;2;3;5;6;7;8;9;10;11;12;13;14;15;17;18;19;20]
```

After the command: UNIQUE\INDICES X[WHERE(Z=2)] Y[WHERE(Z=2)] IDX

```
IDX = [1;2;3;4;5;7;8;9;10]
```

After the command: UNIQUE\INDICES X Y IDX WHERE(Z=2)

```
IDX = [11;12;13;14;15;17;18;19;20]
```

USE

Syntax USE filename

The USE command causes program input to come from a file, instead of from the keyboard. When the end of file is reached, input will again be expected to be entered from the keyboard. Nesting is not allowed. Within USE files, labels, GOTO statements, IF statements and blocks, and DO loops are not allowed.

This command should be useful for entering blocks of commands when the TERMINAL command has been encountered while executing a macro script file, since another script cannot be executed from that mode.

Commands

The default file extension is .PCM If the filename is a text variable, it is first replaced by its value.

VMS: If filename is a logical name, it is replaced by its translation.

If filename does not contain a file name extension, the default file name extension is appended to filename.

If filename doesn't exist in the current directory, and if the logical name PHYSICA\$LIB has been defined, that location is checked.

UNIX: If filename is an environment variable, it is replaced by its translation. If filename cannot be found in the current location, filename with the default file extension appended is checked. If this file cannot be found, and if the environment variable PHYSICA_LIB has been defined, then PHYSICA_LIB/filename is tried.

If this file cannot be found, then PHYSICA_LIB/filename with the default file name extension appended is tried.

Environment variables in file names

For UNIX users, it is now possible to use an environment variable in a file name, if the environment variable is preceded by a \$. For example,

```
setenv FILE mystuff
physica
use $FILE
```

The environment variable can be just the first part of the filename, for example,

```
setenv FILE my
physica
use $FILEstuff
```

VECTOR

```
Syntax VECTOR x1 { x2 ... } n
```

The VECTOR command creates new vectors or changes the lengths of existing vectors.

If x1 is already a vector, it will be either trimmed down to the specified length or zero filled to expand it. If x1 exists but is not a vector, it will be destroyed first. If x1 does not exist, it will be created as a zero filled vector with length n. Other variable names, xI, may be entered. They will be treated just like x1.

VOLUME

By default, the VOLUME command calculates the volume under the scattered points given by vectors x, y and z=f(x,y). The output is an optional scalar, volm. If volm is not entered, the value is simply displayed. The region given by x and y is first triangulated, using a Thiessen triangulation, then the integral is approximated by integrating the piecewise linear interpolants of the data values.

If the \POLAR qualifier is used, the data is assumed to be in polar coordinates with x the radial components and y the angular components, in degrees.

Volume under a matrix

If the \MATRIX qualifier is used, a matrix is expected. The grid region is triangulated. The volume is approximated by summing the averages of the matrix values at the triangle vertices multiplied by the area of each triangle. If x and y are not entered then x defaults to $[1;2;\ldots;ncol]$ and y defaults to $[1;2;\ldots;nrow]$, where ncol is the number of columns of the matrix m and nrow is the number of rows. The \POLAR qualifier is not allowed with matrix data.

WAIT

```
Syntax WAIT n
```

The WAIT command causes a delay of n seconds. This is intended for use when EXECUTing a command script file.

Suppose some message is to be displayed on the terminal screen, via the DISPLAY command, and the user is expected to interactively enter some information. One could ring the bell, using the BELL command, display the message, and then cause a wait of a few seconds before clearing the screen. The user would then have a better opportunity to read the message and then act on it.

Window

```
Syntax WINDOW { n { lowx lowy { upx upy }}}
WINDOW\TILE nx ny nstart

Qualifiers \TILE, \MESSAGES

Defaults \NOMESSAGES, nstart = 1
```

The WINDOW command chooses a subset of the graphics page. The default window is the entire current graphics page, that is, window number zero (0).

If the WINDOW command is entered with no parameters, a listing of the currently defined windows will be displayed, along with their corner coordinates.

Use the \nomessages qualifier to turn off the display of informational messages to the terminal screen.

What are windows

Windows are an easy way to subdivide the graphics output page into rectangular regions, allowing multiple graphs and/or multiple figures and/or multiple text regions. A window is a subset of the page. A window, other than the default zero level window, has a smaller plotting unit range than the full page.

Commensurateness is never lost in a sub-window. Windows are transparent to EDGR.

Boundaries

There is usually at least one rectangle drawn on the monitor screen. The largest rectangle represents the world boundary, that is, the maximum extent of the hardcopy page. A smaller, inner, rectangle, drawn with a dashed line, represents a sub-window within the page. These rectangles are for the user's reference only and will not appear on a hardcopy. These boundary rectangles can be turned off with the DISABLE BORDER command, page 55.

Plotting units

Sub-windows have different plotting unit ranges than the full page. Thus, when a sub-window has been selected, it is possible that not all subsequent graphics will be contained within this window. However, commensurateness is always preserved within a sub-window. For example, circles, which appear as circles when drawn into the full page, will still be circular when drawn into any sub-window.

Defining a new window

```
Syntax WINDOW n { lowx lowy { upx upy }}
```

Window zero, the full page or world, cannot be redefined.

A new window can be defined by including the optional final four parameters with the command. The lower left corner will be (lowx, lowy) and the upper right corner will be (upx, upy). These four parameters should be expressed as percentages of the full page, for example, lowx = 50, lowy = 50 represents the centre of the world, while lowx = 100, lowy = 100 represents the upper right hand corner.

If a window number, n, is entered but not the final four parameters, and that window number is undefined, then the graphics cursor will be used to choose the lower left and upper right corners of the new window.

Pre-defined windows

The initial pre-defined windows are displayed in Table 2.67. See also Figure 2.31.

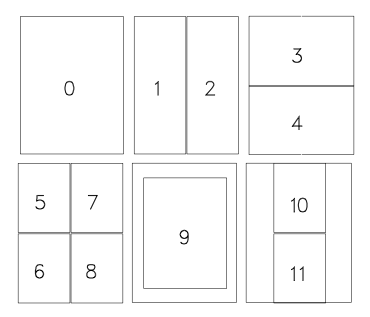


Figure 2.31: The initial pre-defined windows in PORTRAIT orientation

Windows and GPLOT

Choosing a window with the WINDOW command, which has its lower left corner at (lowx, lowy) and its upper right corner at (upx, upy), is equivalent to choosing the GPLOT window with the SET command. That is, the following command:

Commands

| | percentages | | | | | centir | neters | |
|---------------|-------------|------|-----|-----|-------|--------|--------|-------|
| window number | lowx | lowy | upx | upy | lowx | lowy | upx | upy |
| 0 | 0 | 0 | 100 | 100 | 0.00 | 0.00 | 25.00 | 19.00 |
| 1 | 0 | 0 | 50 | 100 | 0.00 | 0.00 | 12.50 | 19.00 |
| 2 | 50 | 0 | 100 | 100 | 12.50 | 0.00 | 25.00 | 19.00 |
| 3 | 0 | 50 | 100 | 100 | 0.00 | 9.50 | 25.00 | 19.00 |
| 4 | 0 | 0 | 100 | 50 | 0.00 | 0.00 | 25.00 | 9.50 |
| 5 | 0 | 50 | 50 | 100 | 0.00 | 9.50 | 12.50 | 19.00 |
| 6 | 0 | 0 | 50 | 50 | 0.00 | 0.00 | 12.50 | 9.50 |
| 7 | 50 | 50 | 100 | 100 | 12.50 | 9.50 | 25.00 | 19.00 |
| 8 | 50 | 0 | 100 | 50 | 12.50 | 0.00 | 25.00 | 9.50 |
| 9 | 10 | 10 | 90 | 90 | 2.50 | 1.90 | 22.50 | 17.10 |
| 10 | 25 | 50 | 75 | 100 | 6.25 | 9.50 | 18.75 | 19.00 |
| 11 | 25 | 0 | 75 | 50 | 6.25 | 0.00 | 18.75 | 9.50 |
| 15 | 0 | 75 | 50 | 100 | 0.00 | 14.25 | 12.50 | 19.00 |
| 16 | 50 | 75 | 100 | 100 | 12.50 | 14.25 | 25.00 | 19.00 |
| 17 | 0 | 50 | 50 | 75 | 0.00 | 9.50 | 12.50 | 14.20 |
| 18 | 50 | 50 | 100 | 75 | 12.50 | 9.50 | 25.00 | 14.20 |

Table 2.67: The initial pre-defined windows

WINDOW n lowx lowy upx upy

is equivalent to the following:

```
SET
%XLWIND lowx
%YLWIND lowy
%XUWIND upx
%YUWIND upy
```

Using the SET command to define a window is *not* recommended. When the CLEAR command is entered, the window chosen with the last WINDOW command is set up, and the values of XLWIND, XUWIND, YLWIND, and YUWIND are changed. Thus, if their values had been re-defined with the SET command, these values would be lost when the CLEAR command is entered.

Multiple window creation

```
Syntax WINDOW\TILE nx ny { nstart }
Default nstart = 1
```

The WINDOW\TILE command divides the graphics page up into nx horizontal by ny vertical windows, giving the first window the number nstart, which must be > 0. The total number of windows just defined will be nx*ny. The window numbered nstart will be in the upper left corner of the page, while the window numbered nx*ny-nstart+1 will be in the lower right corner.

WORLD

```
Syntax WORLD vxin vyin { vxout vyout }

Qualifiers \PERCENT

Defaults absolute coordinates returned
```

The WORLD command converts graph coordinates into world coordinates, or into percentages if the \PERCENT qualifier is used.

The world coordinates are either centimeters or inches depending on the units chosen with the SET UNITS command. The default units are centimeters.

The variables vxin and vyin may be scalars or vectors, but they must both be the same type. The resultant values are displayed on the terminal screen. If output variables vxout and vyout are present, the values are stored there. The type of variable created depends on the variable type of vxin and vyin.

WRITE

```
WRITE file { (format) } x1 { x2 ... }
Syntax
          WRITE\SCALAR file s1 { s2 ... }
          WRITE\MATRIX file matrix
          WRITE\TEXT file txtvar
Qualifiers \SCALAR, \MATRIX, \TEXT, \FORMAT, \APPEND
Examples WRITE FILE.DAT X Y Z
          WRITE\FORMAT FILE.DAT ('X=',F10.3,' Y=',F7.1,' Z=',F9.2) X Y Z
          WRITE\APPEND FILE.DAT X Y Z
          WRITE\SCALAR FILE.DAT X[2] Y[3] M[2,4]
          WRITE\SCALAR\FORMAT FILE.DAT ('String',3(F10.3)) A Y[3] C
          WRITE\MATRIX\APPEND FILE.DAT M[1:100,1:10]
          WRITE\MATRIX\FORMAT FILE.DAT (7F10.3,2X) M[1:100,1:10]
          WRITE\TEXT FILE.DAT 'String'
          WRITE\TEXT FILE.DAT T3
          WRITE\TEXT\APPEND FILE.DAT 'A = '//RCHAR(A, 'F10.3')
```

Commands

WRITE is a general purpose command for writing vectors, scalars, a matrix, or a string. The variable type that will be written is determined by command qualifier. The parameters that are expected also depend on this qualifier. By default, the WRITE command writes vectors to a file.

Environment variables in file names

For UNIX users, it is now possible to use an environment variable in a file name, if the environment variable is preceded by a \$. For example,

```
setenv FILE dum.dat
physica
write $FILE x y z
```

The environment variable can be just the first part of the filename, for example,

```
setenv FILE dum
physica
write $FILE.dat x y z
```

Appending to a file

By default, a new file is opened to receive the output data. If the \APPEND qualifier is used, and if the output file already exists, the data will be appended onto the end of the file.

Formats

```
Syntax WRITE\FORMAT file (format) x1 { x2 ... }
WRITE\SCALAR\FORMAT file (format) s1 { s2 ... }
WRITE\MATRIX\FORMAT file (format) matrix
```

By default, free format is used to write the data to the file. You must use the \FORMAT qualifier to indicate a format is present. The format must be enclosed in parentheses, (and). Any standard FORTRAN format is valid, but only REAL variables can be written. Do not use INTEGER, LOGICAL or CHARACTER formats.

You may use the PHYSICA defined format BINARY, minimum abbreviation B, to write binary unformatted files containing 8 byte numbers.

Vectors

The default is to write vectors. The WRITE command writes the vectors xI to a file. The minimum of the lengths of the vectors will be used. A maximum of twenty-nine (29) vectors can be written with one WRITE command.

If the same vector name is entered more than once, consecutive elements of that vector will be written to the same record. For example, if you enter:

```
WRITE FILE.DAT X X X
```

the following data will be written to the file:

```
X[1] X[2] X[3]
X[4] X[5] X[6]
...
```

or, if you enter: WRITE FILE.DAT X Y X Y X Y

the following data will be written to the file:

```
X[1] Y[1] X[2] Y[2] X[3] Y[3]
X[4] Y[4] X[5] Y[5] X[6] Y[6]
...
```

Examples

To write the vectors X, Y and Z to the file DUM.DAT using free format, that is, there will be three columns of numbers in the file, enter:

```
WRITE DUM.DAT X Y Z
```

To write the vectors X, Y and Z to DUM. DAT using a specified format, enter:

```
WRITE\FORMAT DUM.DAT (' X=',F6.2,' Y=',F6.2,' Z=',F6.2) X Y Z
```

and DUM.DAT will look like:

```
X=123.45 Y=-23.45 Z= 23.45

X=124.45 Y=-24.45 Z= 24.45

X=125.45 Y=-25.45 Z= 25.45

X=126.45 Y=-26.45 Z= 26.45

... ... ...
```

Scalars

The WRITE\SCALAR command writes scalars to a file. One line will be written to the file. A

Commands

maximum of twenty-nine (29) scalars can be written with one WRITE command.

Examples

To write scalars A and B to file FILE.DAT using some format, and then to append to the file the vectors X, Y and Z, enter:

```
WRITE\SCALAR\FORMAT FILE.DAT ('A = ',F10.3,' B = ',F10.3) A B
WRITE\APPEND FILE.DAT X Y Z
```

Matrix

The WRITE\MATRIX command writes a matrix to a file. Only one matrix can be written with each WRITE\MATRIX command.

String

The WRITE\TEXT command writes a string variable, or literal string, to a file. Only one text string can be written with each WRITE\TEXT command and only one record will be written.

Examples

Suppose you want to write a header line to a file and then write some data stored in vectors to that file. For example:

```
WRITE\TEXT FILE.DAT 'This is a header line' WRITE\APPEND FILE.DAT X Y Z
```

Suppose X is a vector, X=[1.1;2.2;3.3;4.4], and you want to write the values of X to a file, with some text. For example:

```
DO J = [1:LEN(X)]

WRITE\TEXT\APPEND FILE.DAT 'X['//RCHAR(J)//'] = '//RCHAR(X[J],'F4.1')

ENDDO
```

ZEROLINES

Syntax ZEROLINES

Qualifiers \HORIZONTAL, \VERTICAL

Defaults both horizontal and vertical lines drawn

The ZEROLINES command draws horizontal and/or vertical lines on a graph through (0,0) depending on the qualifier that is used. The line(s) will have the current line type, as set with the SET LINTYP command.

| qualifier | result |
|-------------|--|
| \HORIZONTAL | a line is drawn through $(0,0)$, parallel to the x -axis, and from the left edge, XMIN, to the right edge, XMAX |
| \VERTICAL | a line is drawn through $(0,0)$, parallel to the y -axis, and from the bottom edge, YMIN to the top edge, YMAX |
| none | both horizontal and vertical lines are drawn |

Example

The following script produces Figure 2.32.

WINDOW 5 ! choose pre-defined window

SCALES -5 5 2 -5 5 2 ! set graph scales GRAPH\AXESONLY ! just draw axes

ZEROLINES! overlay horiz. and vert. lines

WINDOW 6

GRAPH\AXESONLY

ZEROLINES\HORIZONTAL ! overlay horizontal line

WINDOW 12 50 0 75 50 ! define a window

GRAPH\AXESONLY

ZEROLINES\VERTICAL ! overlay vertical line

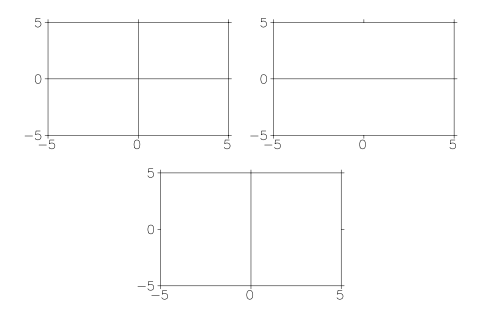


Figure 2.32: An example illustrating the ${\tt ZEROLINES}$ command

3 OPERATORS

Table 3.68 lists the Boolean operators that are recognized in PHYSICA expressions. Table 3.69 lists the other operators. Operators requiring some explanation are described individually in following sections.

| string form | symbolic form | description |
|-------------|---------------|-----------------------|
| "OR" | 1 | or |
| "XOR" | 11 | exclusive or |
| "AND" | & | and |
| "NOT" | ~ | not |
| "LT" | < | less than |
| "GT" | > | greater than |
| "EQ" | < | equal |
| "LE" | <= or ~> | less than or equal |
| "GE" | >= or ~< | greater than or equal |
| "NE" | ~= | not equal |

Table 3.68: Boolean operators

| operator | description | operator | description |
|----------|------------------|-----------------|---------------------|
| () | parentheses | ^ or ** | exponentiation |
| * | multiplication | / | division |
| + | addition | _ | subtraction |
| >< | outer product | <> | inner product |
| <- | matrix transpose | >- | matrix reflect |
| /1 | vector union | /& | vector intersection |
| // | append | | |

Table 3.69: Other operators

Boolean operators

The Boolean operators, listed in Table 3.68, all come in two forms, symbolic and string. The double quotes, " ", are required with the string form. Boolean operators return a value of zero (0) when false and a value of one (1) when true.

The Boolean operators can operate on scalars, vectors, or matrices, but both operands must be the same size and shape. The result of the operation is a variable with this size and shape.

Operators

All of the Boolean operators are binary, except for the not operator, "NOT" or ~, which is unary.

Examples

Suppose you have two vectors: X = [1;2;3;4;5;6;7] Y = [-2;-1;0;1;2;3;4] Then:

```
X"OR"Y = X|Y = [1;1;1;1;1;1]
X"XOR"Y = X||Y = [0;0;1;0;0;0;0]
X"AND"Y = X&Y = [1;1;0;1;1;1;1]
X"EQ"Y = X=Y = [0;0;0;0;0;0;0]
X"NE"Y = X~=Y = [1;1;1;1;1;1]
X"GT"Y = X>Y = [1;1;1;1;1;1]
X"LT"Y = X<Y = [0;0;0;0;0;0;0]
X"GE"Y = X>=Y = [1;1;1;1;1;1;1]
X"LE"Y = X<=Y = [0;0;0;0;0;0;0]
"NOT"(X"OR"Y) = ~(X|Y) = [0;0;0;0;0;0;0]</pre>
```

Transpose

The transpose operator, <-, is a unary operator that produces the transpose of a matrix, that is, the rows and columns are interchanged. Suppose the matrix m has n_r rows and n_c columns. $m[i,j] \longleftrightarrow m[j,i]$ for $i=1,2,\ldots,n_r$ and $j=1,2,\ldots,n_c$.

Example

$$\mathbf{M} = \begin{pmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \\ 10 & 11 & 12 \end{pmatrix}$$

$$< -\mathbf{M} = \begin{pmatrix} 1 & 4 & 7 & 10 \\ 2 & 5 & 8 & 11 \\ 3 & 6 & 9 & 12 \end{pmatrix}$$

Reflect

The reflect operator, >-, is a unary operator that interchanges the columns of a matrix. Suppose the matrix m has n_c columns and n_r rows. Column n_c is interchanged with column 1, that is, $m[1:n_r,1] \longleftrightarrow m[1:n_r,n_c]$; column n_c-1 is interchanged with column 2, that is, $m[1:n_r,2] \longleftrightarrow m[1:n_r,n_c-1]$; and so on.

Examples

$$\mathbf{M} = \begin{pmatrix} 1 & 10 & -1 \\ 2 & 20 & -2 \\ 3 & 30 & -3 \\ 4 & 40 & -4 \end{pmatrix} \rightarrow -\mathbf{M} = \begin{pmatrix} -1 & 10 & 1 \\ -2 & 20 & 2 \\ -3 & 30 & 3 \\ -4 & 40 & 4 \end{pmatrix}$$

You can reflect the rows of a matrix, by using the reflect operator and the transpose operator.

Union

The union operator, /|, is a binary operatore that only accepts vectors as operands, and returns a vector which contains the union of the elements of these two vectors.

All vectors have an order property. Vectors are either in ascending order, descending order, or un-ordered. The type is displayed in the SHOW command, where +0 means ascending order, -0 means descending order, and no symbol means un-ordered. For now, being ordered only has an affect on the vector union, /|, and the vector intersection, /&. These operations are much faster if the vector operands are ordered. The WHERE function produces an ascending order vector, as does the SORT\UP command. The SORT\DOWN command produces a descending order vector. This new vector property will be utilised more in the future to enhance speed and efficiency.

Example

To illustrate vector union, suppose you have two vectors:

$$X = [1;2;3;4;5;6;7]$$
 $Y = [-2;-1;0;1;2]$

Then: X/Y = [-2;-1;0;1;2;3;4;5;6;7]

Intersection

The intersection operator, /&, is a binary operatore that only accepts vectors as operands, and returns a vector which contains the intersection of the elements of these two vectors.

All vectors have an order property. Vectors are either in ascending order, descending order, or un-ordered. The type is displayed in the SHOW command, where +0 means ascending order, -0 means descending order, and no symbol means un-ordered. For now, being ordered only

Operators

has an affect on the vector union, /|, and the vector intersection, /&. These operations are much faster if the vector operands are ordered. The WHERE function produces an ascending order vector, as does the SORT\UP command. The SORT\DOWN command produces a descending order vector. This new vector property will be utilised more in the future to enhance speed and efficiency.

Example

To illustrate vector intersection, suppose you have two vectors:

```
X = [1;2;3;4;5;6;7] Y = [-2;-1;0;1;2]
Then: X/\&Y = [1:2]
```

Append

The meaning of the append operator, //, depends on its operands. If the operands are both vectors, the second vector is appended to the first. If the operands are strings, the second string is appended to the first.

Examples

To illustrate appending vectors, suppose you have two vectors:

```
X = [3;5;7] Y = [-2;-4;-5]
Then: X//Y = [3;5;7;-2;-4;-5]
```

To illustrate appending strings, suppose you have a scalar string variable:

```
T = 'this is a string'
hen: T//' and another' = 'this is a string and another'
```

Outer product

The outer product operator, ><, operates on two vectors and produces a matrix composed of all possible combinations of products of elements of the vectors.

If
$$x = [x_1; x_2; \dots; x_m]$$
 and $y = [y_1; y_2; \dots; y_n]$ then $x > < y = \begin{pmatrix} x_1y_1 & x_1y_2 & \dots & x_1y_n \\ x_2y_1 & x_2y_2 & \dots & x_2y_n \\ \vdots & & & \vdots \\ x_my_1 & x_my_2 & \dots & x_my_n \end{pmatrix}$

Example

Suppose you have two vectors: X = [1;3;5] Y = [2;4]

Then:
$$X > Y = \begin{pmatrix} 2 & 4 \\ 6 & 12 \\ 10 & 20 \end{pmatrix}$$

Inner product

The inner product, <>, operating on two vectors produces a scalar; operating on a vector and a matrix produces a vector, operating on two matrices produces a matrix.

Operating on two vectors

The inner product operating on two vectors produces a scalar, whose value is equal to the sum of the products of the vectors' elements. The two vectors *must* be the same length.

If
$$x = [x_1; x_2; ...; x_n]$$
 and $y = [y_1; y_2; ...; y_n]$ then $x \leftrightarrow y = \sum_{i=1}^n x_i y_i$

Example

Suppose you have two vectors: X = [1;3;5] Y = [2;4;6]

Then: $X \lt \gt Y = 44$

Operating on a vector and a matrix

The inner product operating on a vector and a matrix produces a vector. If the vector is the first operand, its length *must* be equal to the number of rows of the matrix. The resultant vector length will be the number of columns of the second operand matrix.

If
$$x = [x_1; x_2; \dots; x_m]$$
 and $a = \begin{pmatrix} a_{1,1} & a_{1,2} & \dots & a_{1,n} \\ a_{2,1} & a_{2,2} & \dots & a_{2,n} \\ \vdots & & & \vdots \\ a_{m,1} & a_{m,2} & \dots & a_{m,n} \end{pmatrix}$

then
$$(x \le a)_i = \sum_{j=1}^m x_j a_{j,i}$$
 for $i = 1, 2, ..., n$

Examples

Operators

The inner product of a vector and a matrix:

$$X = [1;3;5] \qquad M = \begin{pmatrix} 1 & 4 \\ 2 & 5 \\ 3 & 6 \end{pmatrix}$$

Then: X <> M = [22; 49]

Operating on a matrix and a vector

The inner product operating on a matrix and a vector produces a vector. If the vector is the second operand, its length *must* be equal to the number of columns of the matrix, and the resultant vector length will be the number of rows of the first operand matrix.

If
$$a = \begin{pmatrix} a_{1,1} & a_{1,2} & \dots & a_{1,n} \\ a_{2,1} & a_{2,2} & \dots & a_{2,n} \\ \vdots & & & \vdots \\ a_{m,1} & a_{m,2} & \dots & a_{m,n} \end{pmatrix}$$
 and $x = [x_1; x_2; \dots; x_n]$

then
$$(a > x)_i = \sum_{j=1}^n a_{i,j} x_j$$
 for $i = 1, 2, ..., m$

Examples

The inner product of a matrix and a vector:

$$X = [1;3;5]$$

$$M = \begin{pmatrix} 1 & 3 & 5 \\ 2 & 4 & 6 \end{pmatrix}$$

Then: M <> X = [35; 44]

Operating on two matrices

The inner product operating on two matrices produces a matrix. The number of columns of the first operand matrix *must* be equal to the number of rows of the second operand matrix. The resultant matrix will be a square matrix with the number of rows and the number of

columns equal to the number of rows of the first operand.

$$\mathbf{If} \quad a = \begin{pmatrix} a_{1,1} & a_{1,2} & \dots & a_{1,n} \\ a_{2,1} & a_{2,2} & \dots & a_{2,n} \\ \vdots & & & \vdots \\ a_{m,1} & a_{m,2} & \dots & a_{m,n} \end{pmatrix} \quad \mathbf{and} \quad b = \begin{pmatrix} b_{1,1} & b_{1,2} & \dots & b_{1,m} \\ b_{2,1} & b_{2,2} & \dots & b_{2,m} \\ \vdots & & & \vdots \\ b_{n,1} & b_{n,2} & \dots & b_{n,m} \end{pmatrix}$$

then
$$(a > b)_{i,j} = \sum_{k=1}^{n} a_{i,k} b_{k,j}$$
 for $i = 1, 2, ..., m$ and $j = 1, 2, ..., m$

Example

The inner product of two matrices:

$$A = \begin{pmatrix} 11 & 12 & 13 \\ 21 & 22 & 23 \\ 31 & 32 & 33 \end{pmatrix} \text{ and } B = \begin{pmatrix} 1 & 2 & 3 \\ 2 & 4 & 6 \\ 3 & 6 & 9 \end{pmatrix} \text{ Then: } A \Leftrightarrow B = \begin{pmatrix} 74 & 148 & 222 \\ 134 & 268 & 402 \\ 194 & 388 & 582 \end{pmatrix}$$

4 FUNCTIONS

PHYSICA supports many types of functions, both numeric and string. The basic numeric type of function operates on scalars, vectors, or matrices, but one number at a time. In other words, it performs its calculations on an element by element basis. These include the trigonometric functions, and the basic arithmetic functions such as the exponential and logarithmic functions. The resultant variable type of one of these element by element functions will be the same as the variable type of its argument.

PHYSICA also supports array functions, which operate on variables in their entirety, such as derivative, integral and smoothing functions. Some of these functions have a different resultant variable type than their arguments. There are also functions that operate on strings, such as case changing functions, and functions that have numeric arguments but result in strings.

A special type of the array functions are called looping functions, such as the sum and product functions. The looping functions all require a previously declared scalar dummy variable as second argument. The looping functions mimic standard mathematical notation, for example, the sum:

$$\mathrm{SUM}(f(j),j,1:N) \equiv \sum_{j=1}^N f(j)$$

Where j is the dummy variable. A dummy variable is different from other scalar variables in that its value is only defined while inside the looping function. The first argument of a looping function would normally be some function of the dummy variable, but it is not necessary that the dummy variable appear in the first argument.

Element by element functions

Table 4.70 on page 289 lists the trigonometric functions, while Table 4.71 on page 290 lists other basic numeric functions that are intrinsic to the program. These functions all expect numeric arguments, and operate on an element by element basis.

The trigonometric and basic single argument functions will always have a resultant type which will be the same as the type of its argument, that is, a scalar argument results in a scalar, a vector argument results in a vector with the same length as the argument, and a matrix argument results in a matrix with the same size and shape as the argument. Some of these element by element functions, such as MAX, DIM and ELTIME, do require some explanation, but the definitions of most are assumed to be obvious to the reader.

| function | description | function | description |
|-------------|--------------------------------|----------|--------------------------|
| SIN(x) | Sine (radians) | SINH(x) | Hyperbolic Sine |
| SIND(x) | Sine (degrees) | | |
| COS(x) | Cosine (radians) | COSH(x) | Hyperbolic Cosine |
| COSD(x) | Cosine (degrees) | | |
| TAN(x) | Tangent (radians) | TANH(x) | Hyperbolic Tangent |
| TAND(x) | Tangent (degrees) | | |
| COT(x) | Cotangent (radians) | COTH(x) | Hyperbolic Cotangent |
| SEC(x) | Secant (radians) | SECH(x) | Hyperbolic Secant |
| CSC(x) | Cosecant (radians) | CSCH(x) | Hyperbolic Cosecant |
| ASIN(x) | Arc Sine (radians) | ASINH(x) | Hyperbolic Arc Sine |
| ASIND(x) | Arc Sine (degrees) | | |
| ACOS(x) | Arc Cosine (radians) | ACOSH(x) | Hyperbolic Arc Cosine |
| ACOSD(x) | Arc Cosine (degrees) | | |
| ATAN(x) | Arc Tangent (radians) | ATANH(x) | Hyperbolic Arc Tangent |
| ATAND(x) | Arc Tangent (degrees) | | |
| ATAN2(y,x) | Arc Tangent of y / x (radians) | | |
| ATAN2D(y,x) | Arc Tangent of y / x (degrees) | | |
| ACOT(x) | Arc Cotangent (radians) | ACOTH(x) | Hyperbolic Arc Cotangent |
| ASEC(x) | Arc Secant (radians) | ASECH(x) | Hyperbolic Arc Secant |
| ACSC(x) | Arc Cosecant (radians) | ACSCH(x) | Hyperbolic Arc Cosecant |

Table 4.70: Trigonometric functions

| function | description |
|--------------|--|
| ABS(x) | absolute value $ x $ |
| EXP(x) | exponential e^x |
| FACTORIAL(x) | factorial $x!$ |
| INT(x) | integer portion of x |
| NINT(x) | nearest integer ($x+0.5 	imes \mathtt{SIGN}(1,x)$) |
| LOG(x) | base e logarithm |
| LN(x) | base e logarithm |
| LOG10(x) | base 10 logarithm |
| RAN(x) | random number |
| SQRT(x) | square root \sqrt{x} |
| ELTIME(x) | elapsed time in seconds |
| DIM(x,y) | positive difference function |
| MOD(x,y) | modulus function |
| SIGN(x,y) | transfer of sign |
| MAX(x1,x2,) | maximum of argument list |
| MIN(x1,x2,) | minimum of argument list |

Table 4.71: Basic element by element numeric functions

ATAN2

Syntax vout = ATAN2(v1, v2)

The ATAN2(v1,v2) function returns the Arc Tangent of v1/v2, with $-\pi < \text{ATAN2(v1,v2)} \le \pi$.

- If v1 > 0, the result is positive.
- If v1 = 0, the result is zero if v2 > 0 and π if v2 < 0.
- If v1 < 0, the result is negative. If the value of the second argument is zero, the absolute value of the result is $\pi/2$.

Both arguments must not have the value zero.

ATAN2D

```
Syntax vout = ATAN2D( v1, v2 )
```

The ATAN2D(v1,v2) function returns the Arc Tangent of v1/v2 in degrees, with $-180^{\circ} <$ ATAN2(v1,v2) $\leq 180^{\circ}$.

- If v1 > 0, the result is positive.
- If v1 = 0, the result is zero if v2 > 0 and 180° if v2 < 0.
- If v1 < 0, the result is negative. If the value of the second argument is zero, the absolute value of the result is 90° .

Both arguments must not have the value zero.

RAN

Syntax vout = RAN(v)

The RAN function is an element by element function that requires one (1) argument. The argument can be a scalar, vector or matrix. A scalar argument results in a scalar. A vector argument results in a vector with the same length as the argument, and a matrix argument results in a matrix with the same dimensions as the argument.

RAN uses the current value of the seed to generate the next random number, $0 \le \text{RAN}(v) < 1$. The initial value for the random number seed is 12345. Every time a random number is requested, either from the GENERATE\RANDOM command or from the RAN function, the seed is updated. You can change the seed value with the SET SEED command.

ELTIME

Syntax sout = ELTIME(s)

The ELTIME function is an element by element function that requires one (1) argument. The argument can be a scalar, vector or matrix, but it is intended to have a scalar argument. ELTIME returns the elapsed time in seconds. Initialize the time by calling ELTIME(0), which return 0, and then subsequent calls of ELTIME(s), with s>0, will give the elapsed time since initialization. If not initialized, ELTIME(s), with s>0, returns the elapsed time since midnight. ELTIME(s), with s<0, always returns the elapsed time since midnight. For example:

| call | result |
|------------|---|
| ELTIME(0) | returns 0 (initialization) |
| ELTIME(1) | returns elapsed time since initialization, or |
| | if not initialized, returns elapsed time since midnight |
| ELTIME(-1) | returns elapsed time since midnight |

Functions

DIM

```
Syntax v = DIM( v1, v2 )
```

The DIM function is an element by element function that requires two (2) arguments. The arguments can be scalars, vectors or matrices, but vectors and matrices cannot be mixed, and all arrays must be the same size. Scalar arguments result in a scalar. A vector argument result in a vector with the same length as the argument, and matrix arguments result in a matrix with the same dimensions as the arguments.

| argument 1 | argument 2 | result | | | |
|------------|------------|--------|-----------------------------|---|---------------------------|
| scalar | scalar | scalar | DIM(a,b) | = | max(0, a-b) |
| vector | vector | vector | DIM(x, y)[j] | = | max(0, x[j]-y[j]) |
| scalar | vector | vector | DIM(a, x)[j] | = | max(0, a-x[j]) |
| vector | scalar | vector | DIM(x, a)[j] | = | max(0, x[j]-a) |
| matrix | matrix | matrix | DIM(m1, m2)[i,j] | = | max(0, m1[i,j]-m2[i,j]) |
| scalar | matrix | matrix | <pre>DIM(a, m)[i,j]</pre> | = | max(0, a-m[i,j]) |
| matrix | scalar | matrix | <pre>DIM(m, a)[i,j]</pre> | = | max(0, m[i,j]-a) |

MOD

```
Syntax v = MOD( v1, v2)
```

The MOD function is an element by element function that requires two (2) arguments. The arguments can be scalars, vectors or matrices, but vectors and matrices cannot be mixed, and all arrays must be the same size. Scalar arguments result in a scalar. A vector argument result in a vector with the same length as the argument, and matrix arguments result in a matrix with the same dimensions as the arguments.

| argument 1 | argument 2 | result | | | |
|------------|------------|--------|--------------------|---|--------------------------------------|
| scalar | scalar | scalar | MOD(a,b) | = | a-b*INT(a/b) |
| vector | vector | vector | MOD(x, y)[j] | = | x[j]-y[j]*INT(x[j]/y[j]) |
| scalar | vector | vector | MOD(a, x)[j] | = | a-x[j]*INT(a/x[j]) |
| vector | scalar | vector | MOD(x, a)[j] | = | x[j]-a*INT(x[j]/a) |
| matrix | matrix | matrix | MOD(m1, m2)[i,j] | = | m1[i,j]-m2[i,j]*INT(m1[i,j]/m2[i,j]) |
| scalar | matrix | matrix | MOD(a, m)[i,j] | = | a-m[i,j]*INT(a/m[i,j]) |
| matrix | scalar | matrix | MOD(m, a)[i,j] | = | m[i,j]-a*INT(m[i,j]/a) |

SIGN

```
\overline{Syntax} v = SIGN( v1, v2)
```

The SIGN function is an element by element function that requires two (2) arguments. The arguments can be scalars, vectors or matrices, but vectors and matrices cannot be mixed, and all arrays must be the same size. Scalar arguments result in a scalar. A vector argument result in a vector with the same length as the argument, and matrix arguments result in a matrix with the same dimensions as the arguments.

| argument 1 | argument 2 | result | | | |
|------------|------------|--------|---------------------|---|---------------------------------|
| scalar | scalar | scalar | SIGN(a,b) | = | a (sign of b) |
| vector | vector | vector | SIGN(x, y)[j] | = | <pre> x[j] (sign of y[j])</pre> |
| scalar | vector | vector | SIGN(a, x)[j] | = | <pre> a (sign of x[j])</pre> |
| vector | scalar | vector | SIGN(x, a)[j] | = | <pre> x[j] (sign of a)</pre> |
| matrix | matrix | matrix | SIGN(m1, m2)[i,j] | = | m1[i,j] (sign of m2[i,j]) |
| scalar | matrix | matrix | SIGN(a, m)[i,j] | = | <pre> a (sign of m[i,j])</pre> |
| matrix | scalar | matrix | SIGN(m, a)[i,j] | = | m[i,j] (sign of a) |

MIN

```
Syntax v = MIN(v1, \{v2, ...\})
```

The MIN function is an element by element function that accepts from one (1) to a maximum of twenty (20) arguments. If only one argument is supplied, the minimum element of that argument is returned. If two or more arguments are supplied, the arguments are compared element by element, and the minimum values are returned. The arguments can be scalars, vectors or matrices, but cannot be mixed, and all arrays must be the same size. Scalar arguments result in a scalar. Vector arguments result in a vector with the same length as the arguments, and matrix arguments result in a matrix with the same dimensions as the arguments.

| arguments | result | | | |
|------------|--------|------------------|---|---------------------------|
| one vector | scalar | MIN(x) | = | minimum(x[1],x[2],,x[#]) |
| one matrix | scalar | MIN(m) | = | minimum(m[1,1],,m[#,#]) |
| scalars | scalar | MIN(a,b,) | = | minimum(a,b,) |
| vector | vector | MIN(x,y,)[j] | = | minimum(x[j],y[j],) |
| matrix | matrix | MIN(m1,m2,)[i,j] | = | minimum(m1[i,j],m2[i,j],) |

MAX

```
Syntax v = MAX(v1, \{v2, ...\})
```

The MAX function is an element by element function that accepts from one (1) to a maximum of twenty (20) arguments. If only one argument is supplied, the maximum element of that argument is returned. If two or more arguments are supplied, the arguments are compared

Functions

element by element, and the maximum values are returned. The arguments can be scalars, vectors or matrices, but cannot be mixed, and all arrays must be the same size. Scalar arguments result in a scalar. Vector arguments result in a vector with the same length as the arguments, and matrix arguments result in a matrix with the same dimensions as the arguments.

| arguments | result | | | |
|------------|--------|------------------|---|---------------------------|
| one vector | scalar | MAX(x) | = | maximum(x[1],x[2],,x[#]) |
| one matrix | scalar | MAX(m) | = | maximum(m[1,1],,m[#,#]) |
| scalars | scalar | MAX(a,b,) | = | maximum(a,b,) |
| vector | vector | MAX(x,y,)[j] | = | maximum(x[j],y[j],) |
| matrix | matrix | MAX(m1,m2,)[i,j] | = | maximum(m1[i,j],m2[i,j],) |

Special mathematical functions

Airy's functions

The Airy functions Ai and Bi occur in electromagnetic theory and in quantum mechanics.

Airy's functions of the first kind, AIRY, and second kind, BIRY are

$$\begin{aligned} \text{AIRY}(x) &= Ai(x) = c_1 f(x) - c_2 g(x) \\ \text{BIRY}(x) &= Bi(x) = \sqrt{3} \left[c_1 f(x) + c_2 g(x) \right] \\ \text{where} \quad c_1 &= \frac{3^{-2/3}}{\Gamma(\frac{2}{3})} \quad c_2 = \frac{3^{-1/3}}{\Gamma(\frac{1}{3})} \\ f(x) &= 1 + \frac{1}{3!} x^3 + \frac{1 \cdot 4}{6!} x^6 + \frac{1 \cdot 4 \cdot 7}{9!} x^9 + \cdots \\ g(x) &= x + \frac{2}{4!} x^4 + \frac{2 \cdot 5}{7!} x^7 + \frac{2 \cdot 5 \cdot 8}{10!} x^{10} + \dots \end{aligned}$$

Beta functions

Complete beta function

$$\mathtt{BETA}(a,b) = \beta(a,b) = \int_0^1 t^{a-1} (1-t)^{b-1} dt = \frac{\Gamma(a)\Gamma(b)}{\Gamma(a+b)}$$

Where a and b are both real and positive.

Incomplete beta function

$$\text{BETAIN}(x,a,b) = I_x(a,b) = \frac{\int_0^x t^{a-1} (1-t)^{b-1} dt}{\int_0^1 t^{a-1} (1-t)^{b-1} dt} = \frac{\int_0^x t^{a-1} (1-t)^{b-1} dt}{\beta(a,b)}$$

Bessel functions

First and second kinds

The Bessel functions of the first and second kinds, J_n and Y_n , are linearly independent solutions to the differential equation

$$x^{2} \frac{d^{2}}{dx^{2}} y + x \frac{d}{dx} y + (x^{2} - n^{2}) y = 0$$

Bessel functions arise in solving differential equations for systems with cylindrical symmetry.

BESJO(x) =
$$J_0(x) = \frac{1}{\pi} \int_0^{\pi} \cos(x \sin t) dt = \sum_{k=0}^{\infty} \frac{(-\frac{x^2}{4})^k}{k! k!}$$

BESJ1(x) =
$$J_1(x) = \frac{1}{\pi} \int_0^{\pi} \cos(x \sin t - t) dt = \frac{x}{2} \sum_{k=0}^{\infty} \frac{(-\frac{x^2}{4})^k}{k!(k+1)!}$$

$$\begin{split} \text{BESYO}(x) &= Y_0(x) = \frac{4}{\pi^2} \int_0^{\frac{\pi}{2}} \cos(x \cos t) [\gamma + \ln(2x \sin^2(t)] dt \\ &= \frac{2}{\pi} [-\frac{1}{x} + \ln(\frac{x}{2} + \gamma) J_0(x) - \sum_{k=1}^{\infty} (\sum_{j=1}^k \frac{1}{j}) \frac{(-\frac{x^2}{4})^k}{k! k!}] \\ &= \frac{2}{\pi} [J_0(x) \ln(\frac{\gamma x}{2}) + \sum_{k=1}^{\infty} \frac{(\frac{x^2}{4})^k}{k! k!} (\sum_{j=1}^k \frac{1}{j})] \end{split}$$

$$\text{BESY1}(x) = Y_1(x) = \frac{2}{\pi} \left[-\frac{1}{x} + \ln(\frac{x}{2})J_1(x) - \frac{x}{4} \sum_{k=0}^{\infty} [\psi(k+1) + \psi(k+2)] \frac{(-\frac{x^2}{4})^k}{k!(k+1)!} \right]$$

where ψ is the digamma function, and where γ is Euler's constant

$$\gamma = \lim_{m \to \infty} \left[\sum_{k=1}^{m} \frac{1}{k} - \ln(m) \right] = 0.5772156649...$$

Functions

Modified Bessel functions

The modified Bessel functions of the first and second kinds, I_n and K_n , are solutions to the differential equation

$$x^{2} \frac{d^{2}}{dx^{2}} y + x \frac{d}{dx} y - (x^{2} + n^{2}) y = 0$$

BESIO
$$(x) = I_0(x) = \sum_{k=0}^{\infty} \frac{(\frac{x}{2})^2 k}{k! k!}$$

BESI1(x) =
$$I_1(x) = \frac{x}{2} \sum_{k=0}^{\infty} \frac{(\frac{x}{2})^2 k}{k!(k+1)!}$$

$$\mathrm{BESKO}(x) = K_0(x) = -[\ln(\frac{x}{2} + \gamma]I_0(x) - \sum_{k=1}^{\infty} \sum_{i=1}^k \frac{1}{j} \frac{(\frac{x}{2})^2 k}{k! k!}$$

$$\mathrm{BESK1}(x) = K_1(x) = \frac{1}{x} + \ln(\frac{x}{2})I_1(x) - \frac{x}{4} \sum_{k=0}^{\infty} [\psi(k+1) + \psi(k+2)] \frac{(\frac{x}{2})^2 k}{k!(k+1)!}$$

where ψ is the digamma function, where γ is Euler's constant

$$\gamma = \lim_{m \to \infty} \left[\sum_{k=1}^{m} \frac{1}{k} - \ln(m) \right] = 0.5772156649...$$

Binomial coefficient

$$\mathtt{BINOM}(n,m) = \left(\begin{array}{c} n \\ m \end{array}\right) = \frac{n!}{(n-m)!m!}$$

Chebyshev polynomials

Series of Chebyshev polynomials are used in making numerical approximations to functions.

$$CHEBY(n, x) = T_n(x) = \cos(n\cos^{-1}x)$$

Probability functions

Bivariate normal probability function

$$BIVARNOR(h, k, r) = Probability(x > h, y > k)$$

$$= \int_{h}^{+\infty} \int_{k}^{+\infty} \frac{\exp(-\left[\frac{x^{2}-2rxy+y^{2}}{2(1-r^{2})}\right])}{2\pi\sqrt{1-r^{2}}} dx dy$$

Chi-square probability function

$$\mathtt{CHISQ}(x,n) = \chi^2(x|n) = [2^{\frac{n}{2}}\Gamma(\frac{n}{2})]^{-1} \int_x^{+\infty} t^{\frac{n}{2}-1} e^{-\frac{t}{2}} dt \quad \text{where} \quad 0 \leq x < \infty, \ n \geq 1$$

Inverse Chi-square

Given $y = \chi^2(x|n)$ and n, then x is found: CHISQINV(y,n) = x

Probability integral of Chi-square distribution

$$\mathrm{PROB}(\chi^2,n) = \frac{1}{2^{n/2}\Gamma(n/2)} \int_{\chi^2}^{\infty} t^{\frac{n}{2}-1} e^{-t/2} dt$$

where n is the number of degrees of freedom, $n \ge 1$. For more information, please refer to:

Handbook of Mathematical Functions

by Abramowitz and Stegun, 1964, pages 978ff.

Gaussian or normal probability function

The Gaussian probability function is also known as the normal probability function.

$$\mathrm{GAUSS}(x) = \mathrm{FREQ}(\mathbf{x}) = P(x) = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{x} e^{-t^2/2} dt$$

Normalized Gaussian distribution

Functions

$$\text{NORMAL(x,a,b)} = \frac{1}{b\sqrt{2\pi}}e^{-\frac{(x-a)^2}{2b^2}}$$

Inverse Gaussian

Given $y = \mathtt{GAUSS}(x)$, then x is found: $\mathtt{GAUSSIN}(y) = x$

Cosine integral

$$\mathtt{COSINT}(x) = Ci(x) = -\int_{x}^{\infty} \frac{\cos(t)}{t} dt = \gamma + \ln|x| + \int_{0}^{x} \frac{\cos(t) - 1}{t} dt$$

where γ is Euler's constant

$$\gamma = \lim_{m \to \infty} \left[\sum_{k=1}^{m} \frac{1}{k} - \ln(m) \right] = 0.5772156649...$$

Sine integral

$$SININT(x) = Si(x) = \int_0^x \frac{\sin(t)}{t} dt$$

Dawson's integral

$$\mathtt{DAWSON}(x) = e^{-x^2} \int_0^x e^{t^2} dt$$

Digamma Psi function

$$\mathrm{DIGAMMA}(x) = \psi(x) = \frac{d}{dx}[\ln\Gamma(x)] = \frac{\Gamma'(x)}{\Gamma(x)}$$

Note that
$$\psi(1) = -\gamma$$
, $\psi(n) = -\gamma + \sum_{k=1}^{n-1} k^{-1}$

where γ is Euler's constant

$$\gamma = \lim_{m \to \infty} \left[\sum_{k=1}^{m} \frac{1}{k} - \ln(m) \right] = 0.5772156649...$$

Dilogarithm

The dilogarithm, Li₂, occurs in Feynman diagram integrals in particle physics.

$$\mathrm{DILOG}(x) = Li_2(x) = -\int_0^x \frac{\ln|1-t|}{t} dt = \sum_{k=1}^\infty \frac{x^k}{k^2}$$

 $Li_2(1-x)$ is sometimes known as Spence's integral.

Elliptic integrals

Elliptic integrals have the form $\int \mathcal{F}(x,y)dx$ where \mathcal{F} is a rational function of x and y, and y^2 is a cubic or quartic polynomial in x. Any elliptic integral can be expressed in terms of the three canonical forms. The elliptic integrals are said to be "complete" when the amplitude is $\frac{\pi}{2}$.

First kind

For $|x| \leq 1$ and $|p| \leq \pi/2$

FINELLIC
$$(x,p) = F(p,x) = \int_0^p \frac{dt}{\sqrt{1 - x^2 \sin^2 t}} = \int_0^y \frac{dy}{\sqrt{(1 - y^2)(1 - k^2 y^2)}}$$

For |x| < 1, ELLICK(x) = $F(\pi/2, x)$

Second kind

For $|x| \leq 1$ and $|p| \leq \pi/2$

EINELLIC
$$(x, p) = E(p, x) = \int_0^p \sqrt{1 - x^2 \sin^2 t} dt = \int_0^y \frac{\sqrt{1 - x^2 y^2}}{\sqrt{1 - y^2}} dy$$

where $y = \sin(p)$. For $|x| \le 1$ ELLICE $(x) = E(\pi/2, x)$

Error function

For more information, please refer to:

Functions

Algorithm 610, A Portable FORTRAN Subroutine for Derivatives of the Psi Function, **D.E. Amos**, **ACM Transactions on Mathematical Software**, **December 1983**, Vol. 9, No. 4, pages 494–502.

<u>Handbook of Mathematical Functions</u>, M. Abramowitz, I.A. Stegun, New York, Dover Publications Inc., 1965.

The error function, ERF(x), is the integral of the Gaussian distribution.

$$\mathrm{ERF}(x) = \frac{2}{\sqrt{\pi}} \int_0^x e^{-t^2} dt$$

Inverse

Given y = ERF(x), then x is found: AERF(y) = x

Complementary error function

$$\begin{aligned} \mathtt{ERFC}(x) &=& \frac{2}{\sqrt{\pi}} \int_{x}^{\infty} e^{-t^{2}} dt \\ &=& 1 - \mathtt{ERF}(x) \end{aligned}$$

Inverse

Given y = ERFC(x), then x is found: AERFC(y) = x

Exponential integrals

$$\mathtt{EXPINT}(x) = \int_{x}^{\infty} \frac{e^{-t}}{t} dt$$

$$EI(x) = \int_{-\infty}^{x} \frac{e^{-t}}{t} dt = -EXPINT(-x)$$

Exponential integrals of order n

$$\text{EXPN}(x,n) = \int_{1}^{\infty} \frac{e^{-xt}}{t^n} dt \quad \text{where} \quad n = 0, 1, 2, \dots \quad \text{and} \quad x > 0$$

EXPN(x, n) is related to EXPINT(x) via EXPN(1, 1) = EXPINT(1).

Fermi-Dirac function

$$\mathrm{FERDIRAC}(x,p) = \int_0^\infty \frac{t^p}{e^{t-x}+1} dt \quad \text{ only for } \quad p = -\frac{1}{2}, \; \frac{1}{2}, \; \frac{3}{2}$$

Fisher's F-distribution function

Fisher's *F*-distribution function is also known as the variance-ratio distribution function.

$$\text{FISHER}(m,n,x) = \frac{\Gamma(\frac{m+n}{2})}{\Gamma(\frac{m}{2})\Gamma(\frac{n}{2})} \int_0^{\frac{m}{n}x} \frac{t^{\frac{m}{2}-1}}{(t+1)^{\frac{m+n}{2}}} dt = \frac{m^{\frac{m}{2}}n^{\frac{n}{2}}}{\beta(\frac{m}{2},\frac{n}{2})} \int_0^x t^{\frac{m-2}{2}} (n+mt)^{-\frac{m+n}{2}} dt$$

where $x \ge 0$ and β is the complete β function.

Fresnel integrals

Fresnel and associated Fresnel integrals are related to the error function and occur in diffraction theory.

$$FREC1(x) = C(x) = \int_0^x \cos(\frac{\pi}{2}t^2)dt$$

$$FRES1(x) = S(x) = \int_0^x \sin(\frac{\pi}{2}t^2)dt$$

$$FREC2(x) = \frac{1}{\sqrt{2\pi}} \int_0^x \frac{\cos(t)}{\sqrt{t}} dt = FREC1(\sqrt{\frac{2x}{\pi}})$$

$$FRES2(x) = \frac{1}{\sqrt{2\pi}} \int_0^x \frac{\sin(t)}{\sqrt{t}} dt = FRES1(\sqrt{\frac{2x}{\pi}})$$

Gamma function

$$\operatorname{GAMMA}(x) = \Gamma(x) = \int_0^\infty t^{x-1} e^{-t} dt$$

Natural logarithm of the Gamma function

Functions

$$\operatorname{GAMMLN}(x) = \operatorname{LOGAM}(x) = \ln \Gamma(x)$$

Incomplete Gamma functions

$$\begin{aligned} \operatorname{GAMMACIN}(x,a) &= \Gamma(x,a) = \int_a^\infty t^{x-1} e^{-t} dt \\ \operatorname{GAMMAIN}(x,a) &= \gamma(x,a) = \int_0^a t^{x-1} e^{-t} dt \\ \operatorname{GAMMATIN}(x,a) &= \gamma^*(x,a) = \frac{a^{-x} \gamma(x,a)}{\Gamma(x)} \\ \operatorname{GAMMQ}(x,a) &= \frac{1}{\Gamma(x)} \int_a^\infty t^{x-1} e^{-t} dt \end{aligned}$$

Hermite polynomials

Hermite polynomials arise as the quantum mechanical wave functions for a harmonic oscillator.

HERMITE
$$(n,x) = H_n(x) = (-1)^n e^{x^2} \frac{d^n}{dx^n} (e^{-x^2})$$

Hypergeometric function

The hypergeometric function is also called the Gauss series.

$$\mathrm{HYPGEO}(a,b,c,x) = \ _2F_1(a,b,c,x) = \frac{\Gamma(c)}{\Gamma(a)\Gamma(b)} \sum_{n=0}^{\infty} \frac{\Gamma(a+n)\Gamma(b+n)}{\Gamma(c+n)} \frac{x^n}{n!}$$

Logarithmic confluent hypergeometric function

$$\begin{split} & \text{CHLOGU}(a,b,x) = U(a,b,x) = x^{-a} \, _2\!F_0(a,1+a-b;;-1/x) \\ & = \frac{\pi}{\sin(\pi b)} \left[\frac{M(a,b,x)}{\Gamma(1+a-b)\Gamma(b)} - x^{1-b} \frac{M(1+a-b,2-b,x)}{\Gamma(a)\Gamma(2-b)} \right] \end{split}$$

Where M(a,b,x) is Kummer's Function or the regular confluent hypergeometric function

$$M(a,b,x) = {}_{1}F_{1} = 1 + \frac{a}{b}x + \frac{(a)_{2}}{(b)_{2}}\frac{x^{2}}{2!} + \dots + \frac{(a)_{n}}{(b)_{n}}\frac{x^{n}}{n!} + \dots$$

where
$$(p)_n = p(p+1)(p+2)\cdots(p+n-1) = \Gamma(p+n)$$
 and $(p)_0 = 1$

Note: This function fails when 1 + a - b is close to zero for small x.

Jacobi polynomials

Jacobi polynomials occur in studies of the rotation group, particularly in quantum mechanics. Legendre and Chebyshev are special cases of Jacobi polynomials.

$$\mathtt{JACOBI}(a,b,n,x) = P_n^{(a,b)}(x) = \frac{(-1)^n}{2^n n!} (1-x)^{-a} (1+x)^{-b} \frac{d^n}{dx^n} [(1-x)^{a+n} (1+x)^{b+n}]$$

Kelvin functions

Kelvin functions occur in electrical engineering.

Kelvin functions of the first kind

The Kelvin functions of the first kind, order 0

$$BER(x) = ber_0 x$$
 $BEI(x) = bei_0 x$

where
$$ber_0 x + i \ bei_0 x = J_0(xe^{3\pi i/4})$$

and where J_0 is Bessel's function of the first kind, order 0.

Kelvin functions of the second kind

Kelvin functions of the second kind, order 0, for non-negative x.

$$KER(x) = ker_0x$$
 $KEI(x) = kei_0x$

where
$$ker_0x + i \ kei_0x = e^{-\frac{\nu\pi i}{2}}|_{\nu=0} \ K_{\nu}(xe^{i\pi/4})|_{\nu=0} = K_0(xe^{i\pi/4})$$

and K_0 is the modified Bessel function of the second kind, of order 0.

Laguerre polynomials

Laguerre polynomials are related to hydrogen atom wave functions in quantum mechanics.

$$\mathtt{LAGUERRE}(n,x) = e^x \frac{d^n}{dx^n} [x^n e^{-x}]$$

Legendre functions and polynomials

The most elementary of the Legendre functions, the Legendre polynomials, $P_n(x)$ can be defined by the generating function:

$$(1 - 2xr + r^2)^{-\frac{1}{2}} = \sum_{n=0}^{\infty} P_n(x)r^n$$

More explicit representations are:

$$\text{LEGENDRE}(n,x) = P_n(x) \equiv P_n^0(x) = \frac{1}{2^n n!} \frac{d^n}{dx^n} (x^2 - 1)^n = \,_2F_1(-n,n+1,1,(1-x)/2)$$

where $_2F_1$ is the hypergeometric function.

Unnormalized associated Legendre functions of degree n

$$\begin{array}{lcl} {\rm PLMU}(n,m,x) & = & P_n^m(x) \\ & = & (1-x^2)^{m/2} \frac{d^m}{dx^m} P_n(x) \quad {\rm for} \quad m \geq 0 \\ \\ & = & \frac{(n+m)!}{(n-m)!} P_n^{-m}(x) \quad {\rm for} \quad m < 0 \end{array}$$

Normalized associated Legendre functions

$$\mathtt{PLM}(n,m,x) = \mathtt{PLMN}(n,m,x) = \overline{P_n^m}(x) = \sqrt{\frac{2n+1}{2}\frac{(n-m)!}{(n+m)!}}P_n^m(x)$$

where -1 < x < 1, n is a non-negative integer and $m = -n, \ldots, -1, 0, 1, \ldots, n$ satisfy the differential equation:

$$(1-x^2)\frac{d^2y}{dx^2} - 2x\frac{dy}{dx} + \left[n(n+1) - \frac{m^2}{1-x^2}\right]y = 0$$

Note: PLMU(n, 0, x) = LEGENDRE(n, x).

Poisson-Charlier polynomial

$$\operatorname{POICA}(a,n,x) = \rho_{\eta}^{\alpha}(x) = a^{n/2}(n!)^{-1/2} \sum_{k=0}^{n} (-1)^{n-k} \left(\begin{array}{c} n \\ k \end{array} \right) k! a^{-k} \left(\begin{array}{c} x \\ k \end{array} \right)$$

Rademacher function

$$RADMAC(k, x) = \Upsilon_k(x) = \mathbf{sign}[\sin(2^{k+1}\pi x)]$$

Alternatively, find m such that $m \le 2^{k+1}x < (m+1)$; where $m=0, \pm 1, \pm 2, \ldots$

then
$$\Upsilon_k(x) = \begin{cases} +1 & \text{if } m = \text{even} \\ -1 & \text{if } m = \text{odd} \end{cases}$$

Struve functions

First order

STRUVEO
$$(x) = \frac{2}{\pi} [x - \frac{x^3}{(1 \cdot 3)^2} + \frac{x^5}{(1 \cdot 3 \cdot 5)^2} - \cdots]$$

Second order

$$\mathtt{STRUVE1}(x) = \frac{2}{\pi} [\frac{x^2}{1^2 \cdot 3} - \frac{x^4}{(1 \cdot 3)^2 \cdot 5} + \frac{x^6}{(1 \cdot 3 \cdot 5)^2 \cdot 7} - \cdots]$$

Student's t-distribution

$$\mathtt{STUDENT}(t,n) = P(\frac{t}{n}) = [\beta(\frac{1}{2},\frac{n}{2})\sqrt{n}]^{-1} \int_{-\infty}^{t} (1+\frac{x^2}{n})^{-(n+1)/2} dx$$

where β is the complete β function. Since $\Gamma(\frac{1}{2}) = \sqrt{\pi}$

$$\mathtt{STUDENT}(t,n) = \frac{\Gamma(\frac{n+1}{2})}{\Gamma(\frac{n}{2})\sqrt{n\pi}} \int_{-\infty}^t (1+\frac{x^2}{n})^{-(n+1)/2} dx$$

Inverse

Given $y = P(\frac{t}{n})$ and n, then t is found: STUDENTI(y, n) = t.

Normalized tina resolution

$$\begin{split} \text{TINA}(x,a,b,c) &= \frac{1}{2ce^{b^2/4c^2}}e^{\frac{x-a}{c}}(1-\text{ERF}(\frac{x-a}{b}) \\ &= \frac{1}{2ce^{b^2/4c^2}}e^{(1-\frac{2}{\sqrt{\pi}})(x-a)/c}\int_0^{(x-a)/b}e^{-t^2}dt \end{split}$$

If (x-a)/b > 80 or (x-a)/c > 80 then TINA(x,a,b,c) is returned as zero (0). This is done to circumvent floating overflow problems for large x.

Vector coupling coefficients

Clebsch-Gordan coefficients, Wigner's 3j, 6j, and 9j symbols, Jahn's U-function, and Racah coefficients are the vector coupling coefficients in the theory of angular momentum in quantum mechanics. For more information, please refer to:

Group Theory and its Application to the Quantum Mechanics of Atomic Spectra by Eugene P. Wigner, Academic Press, 1959

Elementary Theory of Angular Momentum

by M.E. Rose, John Wiley & Sons, Inc., 1957

Angular Momentum in Quantum Mechanics

by A.R. Edmonds, Princeton University Press, 1960

The Clebsch-Gordan vector-addition coefficient, $(j_1j_2m_1m_2|jm)$ is defined as:

$$(j_1 j_2 m_1 m_2 | j m) = \delta(m, m_1 + m_2) \sqrt{\frac{(j_1 + j_2 - j)!(j + j_1 - j_2)!(j + j_2 - j_1)!(2j + 1)}{(j + j_1 + j_2 + 1)!}}$$

$$\times \sum_{k} \frac{(-1)^{k} \sqrt{(j_{1} + m_{1})!(j_{1} - m_{1})!(j_{2} + m_{2})!(j_{2} - m_{2})!(j + m)!(j - m)!}}{k!(j_{1} + j_{2} - j - k)!(j_{1} - m_{1} - k)!(j_{2} + m_{2} - k)!(j - j_{2} + m_{1} + k)!(j - j_{1} - m_{2} + k)!}$$

where
$$\delta(i,k) = \begin{cases} 1, & i = k \\ 0, & i \neq k \end{cases}$$

and with the following restrictions:

$$j_1, j_2, j = +n$$
 or $+\frac{n}{2}$ $(n = \text{integer})$ $j_1 + j_2 + j = n$ $+j_1 + j_2 - j$ $+j_1 - j_2 + j$ $-j_1 + j_2 + j$ ≥ 0 $m_1, m_2, m = \pm n$ or $\pm \frac{n}{2}$ $|m_1| \leq j_1, \ |m_2| \leq j_2, \ |m| \leq j$ $(j_1 j_2 m_1 m_2 | j_1 j_2 j_m) = 0$ for $m_1 + m_2 \neq m$

Clebsch-Gordan coefficient function

CLEBSG
$$(j1, j2, j, m1, m2, m) = (j_1 j_2 m_1 m_2 | jm)$$

and

CLEBSG
$$(j1, j2, j) = (j_1 j_2 00 | j0)$$

Note: $(j_1j_200|j0) = 0$ when $j_1 + j_2 + j = 2n + 1$.

Wigner 3 - j function

$$\mathtt{WIGN3J}(j1,j2,j,m1,m2,m) = \left(\begin{array}{ccc} j_1 & j_2 & j \\ m_1 & m_2 & m \end{array} \right) = \frac{(-1)^{j_1-j_2-m}}{\sqrt{2j+1}} (j_1 j_2 m_1 m_2 | j - m)$$

and

$$\mathtt{WIGN3J}(j1,j2,j) = \left(\begin{array}{ccc} j_1 & j_2 & j \\ 0 & 0 & 0 \end{array} \right) = \frac{(-1)^{j_1-j_2}}{\sqrt{2j+1}} (j_1 j_2 00 | j0)$$

Racah coefficients

To define the Racah coefficients we first define the "triangle" coefficient:

$$\Delta(abc) = \sqrt{\frac{(a+b-c)!(a-b+c)!(-a+b+c)!}{(a+b+c+1)!}}$$

Racah's *W*-function is defined as:

$$\begin{split} \text{RACAHC}(a,b,c,d,e,f) &= W(abcd;ef) = \\ &\Delta(abe)\Delta(cde)\Delta(acf)\Delta(bdf) \\ &\times \sum_{k} \frac{(-1)^{k+a+b+c+d}(k+1)!}{(k-a-b-e)!(k-c-d-e)!(k-a-c-f)!(k-b-d-f)!} \\ &\times \frac{1}{(a+b+c+d-k)!(a+d+e+f-k)!(b+c+e+f-k)!} \end{split}$$

Wigner's 6 - j function

$$\mathtt{WIGN6J}(j1,j2,j,m1,m2,m) = \left\{ \begin{array}{ccc} j_1 & j_2 & j \\ m_1 & m_2 & m \end{array} \right\} = (-1)^{j_1 + j_2 + m_1 + m_2} W(j_1 j_2 m_2 m_1; jm)$$

Wigner's 9-j function

$$\mathtt{WIGN9J}(j1,j2,j,m1,m2,m,n1,n2,n) = \left\{ \begin{array}{ccc} j_1 & j_2 & j \\ m_1 & m_2 & m \\ n_1 & n_2 & n \end{array} \right\}$$

please refer to: Group Theory and its Application to the Quantum Mechanics of Atomic Spectra by Eugene P. Wigner, Academic Press, 1959

Jahn's U function

$$JAHNUF(j1, j2, m2, m1, j, m) = U(j_1 j_2 m_2 m_1; jm)$$

$$= (-1)^{j_1+j_2+m_1+m_2} \sqrt{(2j+1)(2m+1)} \left\{ \begin{array}{ccc} j_1 & j_2 & j \\ m_1 & m_2 & m \end{array} \right\}$$

Walsh functions

The Walsh functions form a complete set of orthogonal, normalized, rectangular periodic functions with a period of one.

Let m be found so that k can be written as a binary number:

$$k = \sum_{i=0}^m a_i 2^i;$$
 where $a_i = 0$ or 1 then $\mathrm{WALSH}(k,x) = \prod_{i=0}^m \Upsilon_i^{a_i}(x)$

where Υ_i is the Rademacher function. WALSH can assume the values of 1 or -1 only.

Voigt profile

The Voigt profile function is the convolution integral of a Gaussian and a Lorentzian function. We wish to evaluate

$$G(E) = \int_{-\infty}^{\infty} f(E - E')g(E' - E_1)dE'$$

where f(E) is the Gaussian function

$$f(E) = e^{-E^2/2\sigma^2}$$

and g(E) is the Lorentzian

$$g(E) = \frac{1}{E^2 + \Gamma^2}$$

Putting

$$x = (E' - E)/\sqrt{2}\sigma$$

$$a = (E_1 - E)/\sqrt{2}\sigma$$

$$b = |\Gamma/\sqrt{2}\sigma$$

we obtain

$$G = \frac{1}{\sqrt{2}\sigma} \int_{-\infty}^{\infty} \frac{e^{-x^2}}{b^2 + (a-x)^2} dx$$

This integral has been evaluated for b > 0 in terms of the complex error function $\omega(z)$.

$$G = \frac{1}{\sqrt{2}\sigma} \frac{\pi}{b} \Re[\omega(a+ib)]$$
$$= \frac{1}{\sqrt{2}\sigma} - \Re[e^{b+ia} erfc(b-ia)]$$

The Voigt function implemented here is: $VOIGT(E, E_1, \Gamma/2, \sigma) = G$

Functions that return a string

DATE

The DATE function has no arguments. It returns a string which contains the current date with the format \mathtt{dd} -mmm-yyyy.

Example

| function | result | |
|----------|--------------|--|
| DATE | '6-MAY-1993' | |

TIME

The TIME function has no arguments. It returns a string which contains the current time with the format hh:mm:ss.

Example

| function | result |
|----------|----------|
| TIME | 10:36:25 |

UCASE

The UCASE function converts a string into upper case.

Example

| function | result |
|--------------------------------------|--------------------|
| <pre>UCASE('this is a string')</pre> | 'THIS IS A STRING' |

LCASE

The LCASE function converts a string into lower case.

Example

| function | result |
|------------------|-----------------------------|
| LCASE('THIS IS A | STRING') 'this is a string' |

TCASE

Syntax string = TCASE(string)

The TCASE function toggles the case for each character of a string.

Example

| function | | result |
|----------|--------------------|--------------------|
| TCASE('T | hIs iS A StRiNg') | 'tHiS Is a sTrInG' |
| | | CHAR |
| Syntax | string = CHAR(scal | ar) |
| | string = CHAR(vect | or) |

The CHAR function accepts either a numeric vector or a numeric scalar as argument. It converts these ASCII decimal codes to the equivalent characters and returns these characters as a string. The inverse of this function is the ICHAR function.

Syntax string = EXPAND(string)

The EXPAND function accepts a string as argument. The result is also a character string. It parses the argument, expanding any expression variables it finds. If an expression variable, contained in the argument, also contains expression variables then they are also expanded, and so on until all such expression variables have been expanded. Syntax checking is done during the expansion.

The maximum length of a completely expanded expression is two thousand five hundred (2500) characters.

Example

```
! define a scalar A
A=2
B=3
                 ! define a scalar B
                 ! define a string variable FC1
FC1='(A+B)/A'
FC2='SQRT(A/B)'
                 ! define a string variable FC2
FC3='FC1*FC2'
                 ! define a string variable FC3
FC4='FC3+4*FC2'
                ! define a string variable FC4
=FC4
                 ! displays 'FC3+4*FC2'
=EXPAND(FC4)
                 ! displays '(((A+B)/A)*(SQRT(A/B)))+4*(SQRT(A/B))'
=EVAL(FC4)
                 ! displays 5.307228
```

VARNAME

```
Syntax string = VARNAME(variable)
```

The VARNAME function accepts a variable, either string or numeric, as its argument, and converts that variable name into a string. This function can be useful in scripts where a variable is passed to the script as one of the generalized? parameter. You could then convert the name to a string for display or manipulation.

Examples

The following script shows one way in which the VARNAME function could be used.

```
t1=varname(?1) ! 1st variable name passed to the script converted to a string t2=varname(?2) ! 2nd variable name passed to the script converted to a string graph ?1 ?2 ! plot graph of 2nd variable versus 1st variable text 'graph of '//t1//' vs '//t2 ! lable the plot with the command
```

VARTYPE

```
Syntax string = VARTYPE(name)
```

The VARTYPE function returns the type of the argument as a character string. If the argument is undefined, VARTYPE returns the string unknown. For example, if S is a scalar, X is a vector, M is a matrix, T is a string variable, and TA is a string array variable, then:

returned string

| VARTYPE(1.0) | 'number' |
|-----------------|------------------|
| VARTYPE(S) | 'scalar' |
| VARTYPE(X) | 'vector' |
| VARTYPE(M) | 'matrix' |
| VARTYPE(T) | 'string' |
| VARTYPE(TA) | 'string array' |
| VARTYPE('abc') | 'literal string' |

STRING

```
Syntax quote_string = STRING(some_string)
```

The STRING function is useful in command scripts, where you want to pass a string to the script as a parameter, without enclosing it in quotes. For example, @script test where script.pcm is as follows:

```
x = string(?1)//' is a quote string'
display x
```

would display

```
test is a quote string
```

The key element here is that the user does not have to supply the quotes around the parameter test.

Examples

```
STRING(test case) produces 'test case'
STRING(test) produces 'test'
STRING('test') produces 'test'
```

RCHAR

```
Syntax string = RCHAR(scalar)
string = RCHAR(scalar,string)
```

The RCHAR function accepts a numeric scalar as first argument and returns a string. It converts the numeric value to a string using PHYSICA's own format. You can specify your own format string by including it as the optional second argument, a string.

Examples

| function | result |
|----------------------|--------------|
| RCHAR(-1.234) | '-1.234' |
| RCHAR(PI) | '3.141593' |
| RCHAR(2*PI, 'E10.4') | '0.6283E+01' |

Strings may be appended together using the append operator, //. Scalar values can also be appended to strings using the RCHAR function. For example, suppose A = -1.234, and T is a string variable with T = ' units'.

| function | result |
|--|----------------------------------|
| 'The value of A is '//RCHAR(A)//T | 'The value of A is -1.234 units' |
| The value of A is $'//RCHAR(A, 'F4.1')//T$ | 'The value of A is -1.2 units' |

TRANSLATE

Syntax out_string = TRANSLATE(in_string)

The TRANSLATE function accepts a string as argument, and returns a string which is:

VMS: the translation of the logical name 'in_string'

UNIX: the translation of the environment variable 'in_string'

If 'in_string' has no translation, TRANSLATE simply returns it unchanged.

Examples

The following shows one way in which the TRANSLATE function could be used.

```
read translate('data_dir')//'file.dat' x y z ! pre-define data_dir
```

Numeric functions with string arguments

CLEN

```
Syntax scalar = CLEN(string)
```

The CLEN function accepts only a string as argument. It returns the number of characters in the string. The string can be a literal quote string or a string variable. It cannot be a string

array variable.

ICHAR

Syntax vector = ICHAR(string)

The ICHAR function accepts a string as argument. The string cannot be a complete array string variable, but it can be an element of an array string variable. It returns a numeric vector whose elements are the equivalent ASCII decimal codes for the characters. If the string is only one character, the resultant vector will be of length one. The inverse of this function is the CHAR function.

EQS

Syntax scalar = EQS(string1,string2)

The EQS function accepts two strings as arguments. If the two strings are exactly equal it returns one (1), otherwise it returns zero (0).

NES

Syntax scalar = NES(string1,string2)

The NES function accepts two strings as arguments. If the two strings are exactly equal it returns zero (0), otherwise it returns one (1).

SUB

Syntax scalar = SUB(string1,string2)

The SUB function accepts two strings as arguments. If string1 is a subset of string2 it returns one (1), otherwise it returns zero (0).

SUP

Syntax scalar = SUP(string1,string2)

The SUP function accepts two strings as arguments. If string2 is a subset of string1 it returns one (1), otherwise it returns zero (0).

INDEX

Syntax scalar = INDEX(string1,string2)

The INDEX function accepts two strings as arguments and returns a scalar value. If string2

is a subset of string1 it returns the substring's starting position. If string2 occurs more than once in string1, the starting position of the first (leftmost) occurrence is returned. If string2 does not occur in string1, the value zero (0) is returned.

Example

| function | result |
|-------------------------------------|--------|
| <pre>INDEX('abc', 'abc')</pre> | 1 |
| <pre>INDEX('abcd', 'abc')</pre> | 1 |
| <pre>INDEX('abc','abcd')</pre> | 0 |
| <pre>INDEX('xxabcabc', 'abc')</pre> | 3 |

EVAL

Syntax numeric = EVAL(string)

string variables can be used in numeric expressions, as so called expression variables, to shorten or to simplify an expression. Parentheses around the expression variable are assumed during numeric evaluation. For example:

```
T='A+B'
Y=X*T ! this is equivalent to Y=X*(A+B)
```

A string variable will be numerically evaluated if it is a numeric operand or the argument of a numeric function. Otherwise, a string variable is treated as a string. Use the EVAL function to force numeric evaluation. The type of result, that is, scalar, vector, or matrix, depends on the evaluated expression.

Example

Suppose that string variable T='3+2'.

| input | result |
|----------|-----------------------|
| =T | the string '3+2' |
| =EVAL(T) | the numeric value 5 |

Numeric analysis functions

AREA

```
Syntax scalar = AREA(vector1, vector2)
```

The AREA function calculates the area enclosed in the polygon specified by the vertex coor-

dinates given in the vector arguments. Both vector arguments must be of the same length. The polygon need not be closed, that is, the last point will be assumed to connect to the first point.

DERIV

```
Syntax vector = DERIV(vector1, vector2)
vector = DERIV(vector1, vector2, 'keyword')

Keywords SMOOTH, INTERP, FC, LAGRANGEn (n=3,5,7,9)

Default SMOOTH
```

The DERIV function evaluates the first derivatives of the vector y, the dependent variable, with respect to the vector x, the independent variable, at the x locations. The vector x must be strictly monotonically increasing. The result of this function is a vector with the same length as the vectors x and y. The algorithm that is employed depends on the keyword that is used. By default, the derivatives are calculated using smoothing cubic splines.

Smoothing splines

```
Syntax vector = DERIV(vector1, vector2)
vector = DERIV(vector1, vector2, 'SMOOTH')
```

By default, or if the SMOOTH keyword is used, then smoothing cubic splines, which may not pass through the data points, are used. The spline tension will be the current value of TENSION, which may be changed with the SET command. A tension of zero gives the loosest splines, while a large tension gives linear interpolation. The default tension is 1.

Interpolating splines

```
Syntax vector = DERIV(vector1, vector2, 'INTERP')
```

Cubic splines, which always pass through the data points, are used. The spline tension will be the current value of TENSION, which may be changed with the SET command. A tension of zero gives the loosest splines, while a large tension gives linear interpolation. The default tension is 1.

Lagrange polynomials

```
Syntax     vector = DERIV(vector1, vector2, 'LAGRANGE3')
     vector = DERIV(vector1, vector2, 'LAGRANGE5')
     vector = DERIV(vector1, vector2, 'LAGRANGE7')
     vector = DERIV(vector1, vector2, 'LAGRANGE9')
```

If the LAGRANGEn keyword is used, where n can be 3, 5, 7, or 9, the derivatives are calculated using the method of Lagrange interpolating polynomials.

Monotone piecewise cubic interpolation

```
Syntax vector = DERIV(vector1, vector2, 'FC')
```

If the FC keyword is used, the derivatives are calculated using the Fritsch and Carlson method of monotone piecewise cubic interpolation. This algorithm produces a visually pleasing interpolant, that is, the interpolating curve has no extraneous "bumps" or "wiggles". For an explanation of this method, see:

SIAM Journal of Numerical Analysis, volume 17, number 2, April 1980.

Example

Suppose that you want to see the derivatives, from 0 to π , of $cos(x)^3 + sin(x)^4$. The following commands produce Figure 4.33.

```
LEGEND ON

LEGEND TRANSPARENCY OFF

LEGEND\GRAPH FRAME .25 -2.25 2 -1

SET PCHAR 1

X=[0:PI:.1]

GRAPH 'cos(x)<^>3<_>+sin(x)<^>4<_>' X COS(X)^3+SIN(X)^4

SET PCHAR 2

GRAPH\NOAXES 'derivative' X DERIV(X,COS(X)^3+SIN(X)^4,'LAGRANGE5')

REPLOT
```

INTEGRAL

```
Syntax vector = INTEGRAL(vector1, vector2)
vector = INTEGRAL(vector1, vector2, 'keyword')

Keywords SMOOTH

Default SMOOTH
```

The INTEGRAL function integrates the vector, vector2, the dependent variable, with respect to vector vector1, the independent variable. vector1 must be strictly monotonically increasing. The output of this function is a vector with the same length as vector1 and vector2. The last element of the output vector is the integral over the full range of vector1. The method used depends on the keyword. Currently, there is only one type of integration available.

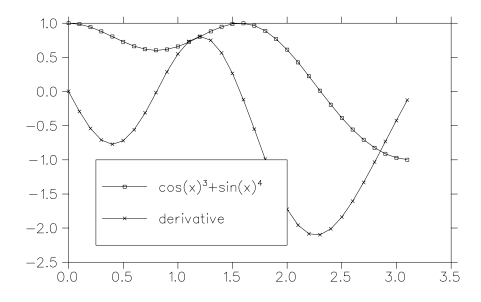


Figure 4.33: An example illustrating the DERIV function

Smoothing splines

```
Syntax vector = INTEGRAL(vector1, vector2)
vector = INTEGRAL(vector1, vector2, 'SMOOTH')
```

By default, the integration method utilizes an interpolating spline under tension. The spline tension used is the current value of <code>TENSION</code>, which may be changed with the <code>SET</code> command. The nature of the interpolating curve varies continuously from pure cubic splines, for <code>TENSION</code> = 0, to a piecewise linear curve, that is, points joined by straight line segments, for large <code>TENSION</code>.

Example

Suppose you need to see the integral values, from 0 to π , of $\cos(x)^3 + \sin(x)^4$. The following sequence of commands could be used. See Figure 4.34

```
LEGEND ON

LEGEND TRANSPARENCY OFF

LEGEND\GRAPH FRAME .25 -.75 2 .25

SET PCHAR 1

X=[0:PI:.1]

GRAPH 'cos(x)<^>3<_>+sin(x)<^>4<_>' X COS(X)^3+SIN(X)^4

SET PCHAR 2

GRAPH\NOAXES 'integral' X INTEGRAL(X,COS(X)^3+SIN(X)^4)

REPLOT
```

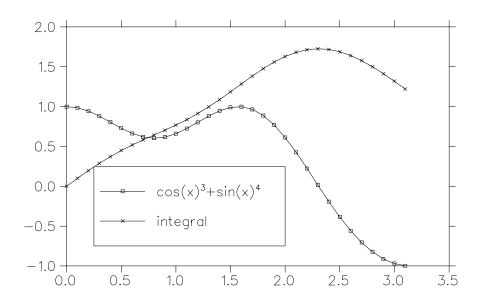


Figure 4.34: An example illustrating the INTEGRAL function

GAUSSJ

Syntax vector = GAUSSJ(matrix, vector)

The GAUSSJ function solves the system of equations m <> x = b, where m is a square matrix and b is a vector, and returns x, the vector of solutions. This function uses the Gauss-Jordan method of elimination with full pivoting. The matrix must be square. The length of the input vector must be the same as the row dimension of the matrix. The function returns a vector with the same length.

Example 1

To solve the following three equations for the three unknowns x_1 , x_2 , and x_3 :

```
3x_1 + 4x_2 + 15x_3 = 26
10x_1 + 2x_2 + 3x_3 = 14
5x_1 - 4x_2 + 3x_3 = 22
You could use the script: m = [[3;1;5]; [4;2;4]; [5;3;3]]
b = [26;14;22]
soln = gaussj(m,b)
```

The answer: soln=[1.238095;-2.365079;2.116402] can be checked by using the outer product operator: m <> soln and comparing this result with the original b vector.

Example 2

The following script, INVERSE.PCM, will find the inverse of a square matrix. If you have a square matrix M, you could find it's inverse, INV_M, with the command: @INVERSE M

```
if ( vlen(?1)[1] "NE" vlen(?1)[2] ) then
  display 'input matrix must be square'
  return
endif
n = vlen(?1)[1]
! identity(n) is the identity matrix of order n
do k = [1:n]
  inv_m[1:n,k] = gaussj(?1,identity(n)[1:n,k])
enddo
```

INVERSE

```
Syntax matrix = INVERSE(matrix)
```

The function INVERSE(m) returns the inverse of the matrix m, which must be a square matrix. The output is a matrix with the same shape as the argument. The answer can be checked by using the inner product operator, for example:

```
invm=INVERSE(m) ! find the inverse of m
=m<>invm ! this should be close to the identity matrix
```

Method

Suppose that the matrix A has n rows and n columns. Let X represent the inverse of A, and let I be the identity matrix:

$$I_{i,j} = \begin{cases} 1 & \text{if} \quad i = j \\ 0 & \text{otherwise} \end{cases}$$

The LU decomposition method² is used for finding the inverse matrix X. Write A as the product of two matrices: A = L <> U where L is lower triangular and U is upper triangular. A lower triangular matrix has elements only on the diagonal and below, while an upper triangular matrix has elements only on the diagonal and above. This decomposition is used to solve n sets of n linear equations. The matrix subscript *,j represents the entire j_{th} column of that matrix.

$$A <\!\!\!\!\!> X_{*,j} = (L <\!\!\!\!> U) <\!\!\!\!> X_{*,j} = L <\!\!\!\!> (U <\!\!\!\!> X_{*,j}) = I_{*,j} \ \ \text{for each} \ \ j = 1,2,\ldots,n$$

Solve for the y vectors, there will be n of them, such that $L > y = I_{*,j}$ and then solve for the j_{th} column of X: $U < > X_{*,j} = y$ for each j = 1, 2, ..., n

Since L and U are triangular

$$y_1 = I_{1,j}/L_{1,1}$$

 $y_i = \left[I_{i,j} - \sum_{k=1}^{i-1} L_{i,k} y_k\right]/L_{i,i}$ for $i = 2, 3, ..., n$

and

$$X_{n,j} = y_n/U_{n,n}$$

 $X_{i,j} = \left[y_{i,j} - \sum_{k=i+1}^{n} U_{i,k} X_{k,j}\right]/U_{i,i}$ for $i = n-1, n-2, \dots, 1$

DET

Syntax scalar = DET(matrix)

The function DET(m) returns the determinant of the matrix m, which *must* be a square matrix. The output is a scalar.

Beware: The determinant of a reasonably sized matrix can get very large, or very small, leading to over/underflows.

The method used to find the determinant uses the LU decomposition of the matrix argument. Please refer to the discussion on LU decomposition of a matrix in the INVERSE function section, page 321.

²The definitions used here are taken from "Numerical Recipes – The Art of Scientific Computing" by W.H. Press, B.P. Flannery, S.A. Teukolsky, W.T. Vetterling, Cambridge University Press, 1986.

IDENTITY

```
Syntax matrix = IDENTITY(scalar)
```

The function IDENTITY(n) returns the identity matrix of order n. That is, the $n \times n$ matrix with 1's on the diagonal and zeros elsewhere. For example:

$$\texttt{IDENTITY(4)} = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}$$

EIGEN

```
Syntax matrix = EIGEN(matrix)
```

If matrix m is an $n \times n$ symmetric matrix, then EIGEN(m) returns a matrix with n rows and n+1 columns. Column n+1 contains the eigenvalues, while columns 1 to n are the eigenvectors of the symmetric matrix m. The eigenvector x and the eigenvalue s of matrix m satisfy the equation: $m \le x = s x$.

One way to check a result is with the following script:

```
e=EIGEN(m)
n=vlen(m)[1]
D0 j = [1:n]
! these should be all zero (or close to zero)
    =m<>e[*,j]-e[j,n+1]*e[*,j]
ENDDO
```

Example

The following script:

```
m=[[2;-1;0;0];[-1;2;-1;0];[0;-1;2;-1];[0;0;-1;2]]
e=eigen(m)
display 'matrix m'
write\matrix\format sys\soutput (4f9.5) m
display 'matrix e'
write\matrix\format sys\soutput (5f9.5) e
```

produces:

```
matrix m

2.00000 -1.00000 0.00000 0.00000

-1.00000 2.00000 -1.00000 0.00000

0.00000 -1.00000 2.00000 -1.00000

0.00000 0.00000 -1.00000 2.00000

matrix e

0.37175 0.60150 0.60150 -0.37175 0.38197

0.60150 0.37175 -0.37175 0.60150 1.38197

0.60150 -0.37175 -0.37175 -0.60150 2.61803

0.37175 -0.60150 0.60150 0.37175 3.61803
```

The eigenvalues are e[*,5] = [0.38197;1.38197;2.61803;3.61803]

The eigenvectors are e[*,j] for j = [1:4]

PFACTORS

```
Syntax vector = PFACTORS(scalar)
```

The PFACTORS(n) function returns a vector containing the prime factors of the scalar n. For example, if you enter X=PFACTORS(420) then X=[2;2;3;5;7].

This function can be usefull in determining a good length for the input vector for the FFT function.

FFT

```
Syntax matrix = FFT(vector)
matrix = FFT(vector, 'keyword')

Keywords AMP&PHASE, COS&SIN

Default AMP&PHASE
```

The FFT function calculates the discrete fast Fourier transform of the input variable, vector. By default, FFT returns the amplitudes and the phases, where the phases are in degrees. If the COS&SIN keyword is used, FFT returns the Fourier coefficients.

Note: The reason that the amplitudes and phases are returned by default is historical. Actually, the Fourier coefficients, that is, the cosine and sine coefficients, are calculated and the amplitudes and phases are just derived from them, as described below. It is a simple matter for the user to request the cosine and sine coefficients, and then to calculate the amplitudes and phases him/herself.

Suppose that the length of the input vector is 2N. The output of this function is a matrix with N+1 rows and 2 columns. The first column contains the amplitudes (or the cosine coefficients), and the second column contains the phases (or the sine coefficients).

The IFFT function calculates the inverse fast Fourier transform.

Fourier coefficients

If the COS&SIN keyword is used, then the FFT function returns the actual Fourier coefficients. Let that the cosine coefficients be called \mathcal{H} 's and the sine coefficients be called \mathcal{G} 's. $\mathcal{H}_0/2$ is the mean value of the input data.

As shown in the example subsection below, these coefficients can be used for smooth interpolation. Suppose x_i is the interpolation location, and 2N is the number of original data points.

$$y_i = \frac{\mathcal{H}_0}{2} + \sum_{k=1}^{N} \left(\mathcal{H}_k \cos(kx_i) + \mathcal{G}_k \sin(kx_i) \right)$$

Discrete Fourier series

Given 2N samples of real data y_j (where $j=0,1,2,\ldots,2N-1$) taken at equally spaced intervals $\Delta t=T/(2N)$, where T is the period, the corresponding Fourier series is:

$$y(t) = \frac{\mathcal{H}_0}{2} + \sum_{k=1}^{N} \left[\mathcal{H}_k \cos(2\pi kt/T) + \mathcal{G}_k \sin(2\pi kt/T) \right]$$

where $G_0 = G_N = 0$ and $H_0/2$ is the mean value of y.

From the original 2N data points, we have exactly 2N calculated coefficients, that is, we have (N+1) \mathcal{H} 's and (N-1) \mathcal{G} 's with "real" information.

If t = 0 at y_0 , then for each y_j we have t = jT/N, for j = 0, 1, ..., 2N - 1, and so

$$y_j = \frac{\mathcal{H}_0}{2} + \sum_{k=1}^{N} \left[\mathcal{H}_k \cos(jk\pi/N) + \mathcal{G}_k \sin(jk\pi/N) \right]$$

The amplitude, A, and the phase, P, are calculated as follows:

$$A_0 = \mathcal{H}_0$$
 $A_N = \mathcal{H}_N$

$$A_j = \sqrt{\mathcal{H}_j^2 + \mathcal{G}_j^2}$$
 for $j = 1, 2, ..., N - 1$

$$P_0 = P_N = 0$$

$$P_j = \frac{180}{\pi} \operatorname{arctan}(\mathcal{G}_j/\mathcal{H}_j)$$
 for $j = 1, 2, \dots, N-1$ and if $P < 0^{\circ}$ then $P = P + 360^{\circ}$

Restrictions

The input vector y is assumed to be periodic with an *even* number of points, that is,

$$LEN(y) = 2N > 2$$
, and $y_0 = y_{2N} = \cdots$

Prime factors

Suppose that $\mathtt{LEN}(y) = 2N$. The largest prime factor of 2N must be ≤ 23 , and there must be no more than 11 distinct prime factors. The product of the square-free prime factors of 2N must be ≤ 210 . The calculation speed is enhanced by using a value of 2N with small prime factors, particularly powers of 2. For reference, the prime numbers less than 1000 are listed below.

```
2 3 5 7 11 13 17 19 23 29 31 37 41 43 47 53 59 61 67 71 73 79 83 89 97 101 103 107 109 113 127 131 137 139 149 151 157 163 167 173 179 181 191 193 197 199 211 223 227 229 233 239 241 251 257 263 269 271 277 281 283 293 307 311 313 317 331 337 347 349 353 359 367 373 379 383 389 397 401 409 419 421 431 433 439 443 449 457 461 463 467 479 487 491 499 503 509 521 523 541 547 557 563 569 571 577 587 593 599 601 607 613 617 619 631 641 643 647 653 659 661 673 677 683 691 701 709 719 727 733 739 743 751 757 761 769 773 787 797 809 811 821 823 827 829 839 853 857 859 863 877 881 883 887 907 911 919 929 937 941 947 953 967 971 977 983 991 997
```

For example, $2N=202=2\times 101$ is not allowed, and $2N=402=2\times 3\times 67$ is also not allowed. If 2N does not satisfy the above restrictions, the input vector can be *padded out*, usually with zeroes, to an even length whose prime factors do not exceed 23. The PFACTORS function, page 324, returns the prime factors of a constant or scalar.

Example

The following script demonstrates how you can use the FFT function to smooth data. Note that when all the Fourier coefficients are used, the smoothed curve must pass through the original data points. See Figure 4.35.

```
! even number of points
N = 16
X = [0:N-1]*2*PI/N
                    ! generate some "data"
Y = SIN(X) + 5*RAN(X)
M = FFT(Y, 'COS&SIN') ! calculate Fourier coefficients
H = M[*,1]
                     ! extract column 1 as a vector
G = M[*,2]
                     ! extract column 2 as a vector
Z = [0:2*PI:.05]
SCALAR\DUMMY K
                    ! define K to be dummy variable for SUM function
SCALE 0 6 0 -2 7 0
                    ! set axis scales
SET %XLABSZ 5
                    ! increase the size of the x-axis label
DO J = [3:N/2+1:2]
 WINDOW (J-3)/2+15
                    ! choose a graphics window
                    ! choose plotting symbol
 SET PCHAR -1
 LABEL\XAXIS 'Number of harmonics used = '//RCHAR(J)
 GRAPH X Y
                     ! plot original data
 SET PCHAR 0
                     ! choose no plotting symbol
 ENDDO
GRAPH\NOAXES [0;2*PI] [H[1]/2;H[1]/2] ! overlay the mean value
             Number of harmonics used = 3
                                          Number of harmonics used =
```

Figure 4.35: An FFT example showing data smoothing

Number of harmonics used = 9

Number of harmonics used = 7

IFFT

Syntax vector = IFFT(matrix)

vector = IFFT(matrix, 'keyword')

Keywords AMP&PHASE, COS&SIN

Default AMP&PHASE

The IFFT function calculates the inverse discrete Fourier transform of the two column input matrix. This matrix is usually calculated by the FFT function, thus reconstructing the original data.

By default, IFFT expects amplitudes and phases, where the phases are in degrees. The first column of the matrix should contain the amplitudes and the second column the phases. If the COS&SIN keyword is used, IFFT expects the Fourier coefficients, that is, the cosine coefficients in the first column and the sine coefficients in the second column. If the input matrix has N rows, the function returns a vector with length 2(N-1).

The principle usage of the IFFT function is to modify some of the amplitudes returned from the FFT function and note their effect on the original data. A typical application would be one of data smoothing, in which the user would zero out the amplitudes of the higher order harmonics.

CONVOL

Syntax vector = CONVOL(vector1, vector2, scalar)

The CONVOL function accepts vectors as the first two arguments. It convolutes or deconvolutes the input vector, vector1, with the specified blurring vector, vector2. The result is a vector the same length as vector1. The third argument should be a scalar, and indicates which operation is to be performed.

Convolution of an odd number of points

```
Syntax vector = CONVOL(x,b,0)
```

The blurring vector, b, must contain an odd number of points. Suppose that N is the length of b and b is the length of b. The convolution of b with b is:

$$y_i = \sum_{j=1}^{N} x_{i-\frac{N}{2}+j-1} \cdot b_{N-j+1}$$

$$= x_{i-\frac{N}{2}}b_N + x_{i-\frac{N}{2}+1}b_{N-1} + \dots + x_{i+\frac{N}{2}-1}b_2 + x_{i+\frac{N}{2}}b_1$$

for $i = 1, 2, 3, \dots, M$. References to subscripts out of the range of x are not summed. The

blurring vector is normalized to 1 to insure that the integrals of the y and x are identical. This normalization is internal, the blurring vector is returned unchanged. To ensure proper convolution, x should be padded at it's upper and lower ends with zeros so it's length is at least the minimum of:

the non-zero length of b; and $\frac{1}{2}$ the length of b

Note: The lengths of b and x can differ. To avoid centroid shifts in the output, centre the blurring vector properly. For example, suppose that b has 2N-1 elements containing a gaussian, then it's peak should be at N.

Convolution of an even number of points

```
Syntax vector = CONVOL(x,b,1)
```

The blurring vector, b, should contain an even number of points. The preferred lengths are powers of 2. The convolution is done using fast Fourier transforms. The following restrictions apply:

- x must be padded at it's lower end with zeros with the number of elements which are non-zero in b, for example, if x and b are of length 128, and b is zero in the range 30 128, then x must contain zeros in locations 1 29
- x and b must have the same length
- the end points of *b* must not be equal. A difference of less than 0.0001 produces oscillations in the deconvoluted result. The usual way is to shift *b* to the left so that the first point has a non-zero value. Together with the first restriction, this ensures that the right point has the value zero, leaving the ends unequal.

Noise in b produces a change in the output, which, due to averaging, has a small effect. Noise effects depend on the shape of the deconvoluted peak.

Convolution noise effects

The narrower this peak, the more effect the noise in b has. This occurs because each noisy point becomes a greater percentage of the total number in the convoluting or deconvoluting function, thus reducing the average effect. In many applications, the noise in the measured data is statistical in nature and so, to reduce the sensitivity to this noise on the deconvolution, apply smoothing filters on the measured data before deconvolution.

Deconvolution of an even number of points

```
Syntax 	 vector = CONVOL(x,b,-1)
```

The blurring vector, *b*, should contain an even number of points. The preferred lengths are powers of 2. The deconvolution is done using fast Fourier transforms. The following restrictions apply:

- 1. x must be padded at it's lower end with zeros with the number of elements which are non-zero in b, for example, if x and b are of length 128, and b is zero in the range 30 128, then x must contain zeros in locations 1 29
- 2. x and b must have the same length
- 3. the end points of *b* must not be equal. A difference of less than 0.0001 produces oscillations in the deconvoluted result. The usual way is to shift *b* to the left so that the first point has a non-zero value. Together with the first restriction, this ensures that the right point has the value zero, leaving the ends unequal.

Noise in b produces a change in the output, which, due to averaging, has a small effect. Noise effects depend on the shape of the deconvoluted peak. The narrower this peak, the more effect the noise in b has. This occurs because each noisy point becomes a greater percentage of the total number in the convoluting or deconvoluting function, thus reducing the average effect. In many applications, the noise in the measured data is statistical in nature and so, to reduce the sensitivity to this noise on the deconvolution, apply smoothing filters on the measured data before deconvolution.

INTERP

The INTERP function interpolates the data contained in vector1, the independent variable, and vector2, the dependent variable. vector1 must be strictly monotonically increasing. The interpolant locations are given in vector3. The INTERP function will return the interpolated values as a vector with the same length as vector3. The algorithm that is employed depends on the keyword that is used. The default is use interpolating splines.

An interpolated curve will always pass through the original data points. If it is not important that the curve pass through the original data, use the SMOOTH function. If your independent variable is not monotonically increasing, use the SPLINTERP function.

Spline interpolation

```
Syntax vector = INTERP(x,y,xi)
vector = INTERP(x,y,xi,'SPLINE')
```

By default, or if the SPLINE keyword is used, the interpolant is calculated by the method of cubic splines under tension. The tension factor corresponds to the "curviness", and must be greater than zero. If it is close to zero, each interpolated function is almost a cubic spline and the resulting curve is quite "loose". If the tension is large, then the resultant is almost linear. The tension used is the current value of TENSION, which may be changed with the SET command.

Linear interpolation

```
Syntax vector = INTERP(x,y,xi,'LINEAR')
```

If the LINEAR keyword is used, the interpolating method used is linear interpolation.

Lagrange interpolation

```
Syntax vector = INTERP(x,y,xi,'LAGRANGE')
```

If the LAGRANGE keyword is used, the interpolating method used is general Lagrange interpolation. The degree of the interpolating polynomial depends on the number of input data points. Suppose that N is the number of points, equal to the lengths of x and y. If N=2, then a simple straight line is used for interpolating. If N=3 or N=4, then a quadratic is used. If $N\geq 5$, then a polynomial of degree 4 is used for the interpolation.

Interpolation by monotone piecewise cubic polynomials

```
Syntax vector = INTERP(x,y,xi,'FC')
```

If the FC keyword is entered, the interpolant is calculated using the Fritsch and Carlson method of monotone piecewise cubic interpolation. This algorithm produces a visually pleasing interpolant, that is, the interpolating curve has no extraneous "bumps" or "wiggles".

For an explanation of this method, see:

SIAM Journal of Numerical Analysis, volume 17, number 2, April 1980.

SPLINTERP

```
Syntax vector = SPLINTERP(vector1, vector2, scalar)
```

The SPLINTERP function interpolates the data contained in vector1, the independent variable, and vector2, the dependent variable. vector1 need not be monotonically increasing. The interpolated curve will always pass through the original data points. The number of output interpolant locations is given in scalar. Suppose scalar = n. The output of this function is a matrix with n rows and 2 columns. The first column will contain the output locations and the second column the interpolated values.

The points are first parameterized in terms of normalized arc length. The normalized length of x is the real length divided by the range of x, that is, the maximum value minus the minimum value. The arclength at a point is approximated by the sum of the lengths of straight line segments connecting all points up to that point. A spline under tension is calculated for vector1 versus arc length and vector2 versus arc length. The vector1 and vector2 values are interpolated separately and then combined to form the output interpolant.

The tension factor corresponds to the "curviness", and must be greater than zero. If it is close to zero, each interpolated function is almost a cubic spline and the resulting curve is quite "loose". If the tension is large, then the resultant is almost linear. The tension used is the current value of TENSION, which may be changed with the SET command.

For monotonically increasing data, use the INTERP function.

SMOOTH

```
Syntax vector = SMOOTH(vector1, vector2, vector3)
vector = SMOOTH(vector1, vector2, vector3, vector4)
```

The SMOOTH function calculates a smooth curve through the data contained in vector1, the independent variable, and vector2, the dependent variable. vector1 *must* be strictly monotonically increasing. The output locations are given in vector3, which must also be monotonically increasing. This function returns the smoothed values as a vector with the same length as vector3. The smooth curve is calculated by the method of cubic splines under tension.

For another smoothing method, using Savitzky-Golay filters, use the SAVGOL function. Depending on the tension, a smoothed curve may not pass through the original data points. If you want the curve to always pass through the original data, use the INTERP function. If your data is not monotonically increasing, use the SPLSMOOTH function.

Weights

```
Syntax vector = SMOOTH(x,y,xout,w)
```

If no weights, w, are entered, the weight at each data point defaults to 1. The weights control

the amount of smoothing at each data point. As the weight at a point decreases, the spline fits that data point more closely.

Spline tension

The tension factor corresponds to the "curviness", and must be greater than zero. The tension used is the current value of TENSION, which may be changed with the SET command.

If the tension is set to zero, the result will be an interpolating cubic spline. If the tension is large, the result will be the least-squares line through the data. As the tension decreases, the amount of smoothing decreases and the data points are fit more exactly. As the tension increases, the fit straightens and has less curvature at peaks, valleys and endpoints.

Suppose that N is the length of y and that the weights are the standard deviations of y. Values of tension t in the range $N-\sqrt{2N} \le t \le N+\sqrt{2N}$ give the most natural looking results. To obtain the most suitable fit, the user may wish to do several runs with different values of t. By observing the spline fits plotted on a graph, the fit with the most suitable amount of smoothing can be selected.

Method

Given a set of abscissae: $x_1 < x - 2 < \ldots < x_N$,

a cubic spline function over the region (x_1, x_N) is composed of cubic parabolas

$$G(x) = a_i + b_i \cdot (x - x_i) + c_i \cdot (x - x_i)^2 + d_i \cdot (x - x_i)^3$$

where $x_i \le x < x_{i+1}$, which join at the endpoints x_i such that \mathcal{G} , \mathcal{G}' , and \mathcal{G}'' are continuous.

The smoothing function is constructed by minimizing $\int_{x_1}^{x_N} \mathcal{G}''(x)^2 dx$ subject to the constraint $\sum_{i=1}^{N} \left((\mathcal{G}(x_i - y_i)/w_i)^2 \le t \right)$ where $w_i > 0$ are the weights and $t \ge 0$ is the spline tension.

The solution proceeds by the standard methods of minimizing the functional

$$\int_{x_1}^{x_N} \mathcal{G}''(x)^2 dx + p \cdot \left\{ \sum_{i=1}^N \left((\mathcal{G}(x_i) - y_i) / w_i \right)^2 + z^2 - t \right\}$$

where z and p are auxiliary parameters. The functional is minimized with respect to z and p by setting the partial derivatives with respect to z and p equal to zero.

SPLSMOOTH

The SPLSMOOTH function calculates a smooth curve through the data contained in vector1, the independent variable, and vector2, the dependent variable. vector1 need not be monotonically increasing. The number of output locations is given in scalar. Suppose scalar = n. The output of this function is a matrix with n rows and 2 columns. The first column will contain the output locations and the second column the smoothed values.

The points are first parameterized in terms of normalized arc length. The normalized length of x is the real length divided by the range of x, that is, the maximum value minus the minimum value. The arclength at a point is approximated by the sum of the lengths of straight line segments connecting all points up to that point. A spline under tension is calculated for vector1 versus arc length and vector2 versus arc length. The vector1 and vector2 values are interpolated separately and then combined to form the output interpolant.

The tension factor corresponds to the "curviness", and must be greater than zero. If it is close to zero, each interpolated function is almost a cubic spline and the resulting curve is quite "loose". If the tension is large, then the resultant is almost linear. The tension used is the current value of TENSION, which may be changed with the SET command.

For monotonically increasing data, use the SMOOTH function.

Weights

```
Syntax matrix = SPLSMOOTH(x,y,n,w)
```

If m is the length of w, x and y, the goal is to achieve: $\sum_{i=1}^{m} ((G_i - y_i)/w_i)^2 = s$ where G_i is the cubic spline function at x_i and s is the spline tension. As w_i decreases, the spline fits the data point (x_i, y_i) more closely.

SAVGOL

```
Syntax vector = SAVGOL(scalar1,scalar2,vector)
```

The SAVGOL function calculates a smooth curve through the data contained in vector, the dependent variable, using the Savitzky-Golay smoothing filter method. The order of the filter is given in scalar1, and can be 2 or 4. The filter width is given in scalar2. These filters preserve the area under the data, the zeroth moment, but also the higher moments.

There is no input independent variable, as the data is assumed to be equally spaced.

As a rough guideline, best results are obtained when the filter width, m, of the order 4 Savitzky-Golay filter is between one and two times the full width half maximum (FWHM) of desired features in the data.

The Savitzky-Golay smoothing method

A nonrecursive filter is defined by the convolution formula: $yout_i = \sum_{j=-N_L}^{N_U} c_j y_{i+j}$

The c_j 's are the coefficients of the filter, the y's are the input data, and the yout's are the outputs. The set of points y_{i-N_L} to y_{i+N_U} define the moving window of the filter, where $N_L = m/2$ and $N_U = m-N_L-1$. The Savitzky-Golay smoothing method finds filter coefficients that preserve the 0^{th} , the 1^{st} , the 2^{nd} , and higher moments. The idea is to approximate the underlying function within a moving filter window by a polynomial of order 2, or of order 4. For each data point, y_i , least-squares fit a polynomial to all m points within the filter window, and then set $yout_i$ to be the value of that polynomial at position i. No use is made of the value of that polynomial at any other point. For the next point, y_{i+1} , a whole new least-squares fit is done using the shifted window.

Since the process of least-squares fitting involves only a linear matrix inversion, the coefficients of a fitted polynomial are themselves linear in the values of the data. All the least-squares fitting can be done in advance, for fictitious data consisting of all zeros except for a single 1, and then the fits on the real data are done by taking linear combinations. Thus, there are particular sets of filter coefficients, c_j , for which the convolution formula automatically accomplishes the process of polynomial least-squares fitting inside a moving window.

The Savitzky-Golay filters provide smoothing without loss of resolution when the underlying function can be locally well fitted by a polynomial. This is true for smooth line profiles not too much narrower than the filter width. When this is not true, these filters have no advantage over other smoothing methods.

If the data is irregularly sampled, that is, the y values are not equally spaced, one can pretend that the data points are equally spaced. This amounts to virtually shifting, within each moving window, the data points to equally spaced positions. Such a shift introduces the equivalent of an additional source of noise into the function values. In those cases where smoothing is useful, this noise will often be much smaller than the noise already present. Specifically, if the location of the points is approximately random within the window, then a rough criterion is this: If the change in y across the full width of the window, m, is less than $\sqrt{m}/2$ times the measurement noise on a single point, then this implementation of the Savitzky-Golay filter can be used.

For more information on Savitzky-Golay filters, see:

Computers in Physics, volume 4, number 6, November/December 1990, pages 669 - 672.

JOIN

Syntax scalar = JOIN(vector1, vector2)

The arguments of the JOIN function must both be vectors. JOIN produces a matrix with 3 columns. The first column is the intersection of vector1 and vector2, that is, if you enter m=join(x,y) then m[*,1] is the same as x/&y. m[i,2] is the index of x from which m[i,1] was taken, and m[i,3] is the index of y from which m[i,1] was taken. If the vector arguments are ordered, the JOIN function will proceed much faster than if they are unordered.

Example

Suppose that you have two vectors:

$$X = [0;1;2;3;4;5;6;7;8;9;10], Y = [1;3;5;7;9]$$

then
$$JOIN(X,Y) = \begin{pmatrix} 1 & 2 & 1 \\ 3 & 4 & 2 \\ 5 & 6 & 3 \\ 7 & 8 & 4 \\ 9 & 10 & 5 \end{pmatrix}$$

Functions that return a variable's characteristics

EXIST

Syntax scalar = EXIST(string)

The EXIST function accepts one string as its argument and returns a one (1) if the string is an existing variable name, and zero (0) is it is not.

Example

If you enter X2=3, the function EXIST('X2') would have the value 1.

You can even enter: N=2 and T='X'//RCHAR(N) and the function EXIST(T) would have the value 1.

LEN

Syntax scalar = LEN(vector)

The LEN function only accepts a vector as argument. It returns the length of the vector as a scalar.

VLEN

Syntax vector = VLEN(vector)
vector = VLEN(matrix)

The VLEN function accepts either a vector or a matrix as argument. It returns the length of each dimension of the argument. If the argument is a vector, the result is a vector of length one. If the argument is a matrix, the result is a vector of length two, with the first element being the number of rows and the second the number of columns.

FIRST

Syntax scalar = FIRST(vector)

The FIRST function returns the starting index for vector. The result of the FIRST function is a scalar. The length of vector x is LAST(x)-FIRST(x)+1, which is equal to LEN(x).

LAST

Syntax scalar = LAST(vector)

The LAST function returns the final index for vector. The result of the LAST function is a scalar. The length of vector x is LAST(x)-FIRST(x)+1, which is equal to LEN(x).

ICLOSE

Syntax scalar = ICLOSE(vector, scalar)

The ICLOSE function returns the index of vector which corresponds to where vector is closest to the value of scalar, that is, ICLOSE returns the *first* j where |vector[j] - scalar| is a minimum.

IEQUAL

Syntax scalar = IEQUAL(vector, scalar)

The IEQUAL function returns the index of vector which corresponds to where vector is equal to the value of scalar, that is, IEQUAL returns the *first* j where |vector[j] = scalar|. If scalar is not in the vector, IEQUAL returns zero (0).

WHERE

Syntax vector = WHERE(vector)

The WHERE function only accepts a vector as its argument. It returns the indices where this vector is not equal to zero.

Examples

| function | result |
|------------------------|---------------|
| WHERE([-5:5]>0) | [7;8;9;10;11] |
| WHERE($[-5:5] <= 0$) | [1;2;3;4;5;6] |

Suppose you have two vectors X and Y and you want to graph only those points that lie within the unit circle, that is, that satisfy $SQRT(X^2+Y^2) \le 1$

```
IDX=WHERE(SQRT(X^2+Y^2)<=1)
GRAPH X(IDX) Y(IDX)</pre>
```

Shape changing functions

FOLD

Syntax matrix = FOLD(vector, scalar)

The FOLD function has two arguments. The first must be a vector, and the second a scalar. The result is a matrix formed by folding the data in the vector into the columns of a matrix. Suppose that vector x has m elements. $FOLD(x,n)_{i,j} = x_{i+(j-1)m}$ for $i=1,2,\ldots,n$ and $j=1,2,\ldots,m/n$. Note that m must be divisible by n.

Example

| function | result |
|--------------------------|--|
| | $ \left(\begin{array}{ccccc} 1 & 4 & 7 & 10 \\ 2 & 5 & 8 & 11 \\ 3 & 6 & 9 & 12 \end{array}\right) $ |
| FOLD([1:12],3) | 2 5 8 11 |
| | $\begin{pmatrix} 3 & 6 & 9 & 12 \end{pmatrix}$ |
| | $\begin{pmatrix} 1 & 5 & 9 \end{pmatrix}$ |
| EOI D ([4 , 4 O] / A) | $\begin{pmatrix} 1 & 5 & 9 \\ 2 & 6 & 10 \end{pmatrix}$ |
| FOLD([1:12],4) | 3 7 11 |
| | $\begin{pmatrix} 4 & 8 & 12 \end{pmatrix}$ |

UNFOLD

Syntax vector = UNFOLD(matrix)

The UNFOLD function has one argument, which must be a matrix. The result is a vector formed by unfolding the data in the rows of matrix. Suppose that matrix m has n_c columns and n_r rows. Then $UNFOLD(m)_{j+(i-1)n_c}=m_{i,j}$ for $i=1,2,\ldots,n_r$; and $j=1,2,\ldots,n_c$.

Example

Suppose that matrix
$$M = \begin{pmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{pmatrix}$$

| function | result |
|-----------|---|
| UNFOLD(M) | [1; 2; 3; 4; 5; 6; 7; 8; 9; 10; 11; 12] |

ROLL

The ROLL function accepts either a vector or a matrix as its first argument. It shifts the elements of a vector or the rows of a matrix by the specified step size, scalar.

- If scalar = n > 0, the last n elements of the vector or the last n rows of the matrix are rolled around to the beginning.
- If scalar = n = 0, the vector or matrix is returned unchanged.
- If scalar = n < 0, the first n elements of the vector or the first n rows of the matrix are rolled around to the end.

If scalar is non-integral, then linear interpolation is used to generate new values.

| function | result | |
|--|---|--|
| ROLL([1:10],2) | [9;10;1;2;3;4;5;6;7;8] | |
| ROLL([1:10],-2) | [3;4;5;6;7;8;9;10;1;2] | |
| ROLL([1:10],1.7) | [9.3;7.3;1.3;2.3;3.3;4.3;5.3;6.3;7.3;8.3] | |
| | | |
| Suppose matrix $M = \begin{pmatrix} 1 \\ 5 \\ 9 \end{pmatrix}$ | $\begin{bmatrix} 2 & 3 & 4 \end{bmatrix}$ | |
| Suppose matrix $M = \begin{bmatrix} 5 \end{bmatrix}$ | 6 7 8 | |
| (9) | 10 11 12 | |

Functions

| function | result |
|-------------|--|
| ROLL(M,2) | $ \left(\begin{array}{cccc} 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \\ 1 & 2 & 3 & 4 \end{array}\right) $ |
| ROLL(M,-1) | $ \left(\begin{array}{ccccc} 9 & 10 & 11 & 12 \\ 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \end{array}\right) $ |
| ROLL(M,1.5) | $ \left(\begin{array}{cccc} 7 & 8 & 9 & 10 \\ 5 & 6 & 7 & 8 \\ 3 & 4 & 5 & 6 \end{array}\right) $ |

STEP

The STEP function accepts either a vector or a matrix as its first argument. It shifts the elements of a vector or the rows of a matrix by the specified step size, scalar.

- If scalar = n > 0, the last n elements of the vector or the last n rows are lost.
- ullet If scalar = n = 0, the vector or matrix is returned unchanged.
- If scalar = n < 0, the first n elements of the vector or the first n rows are lost.

If scalar is non-integral, then linear interpolation is used to generate new values.

| function | result |
|--|---------------------------------------|
| STEP([1:10],2) | [1;1;1;2;3;4;5;6;7;8] |
| STEP([1:10],-2) | [3;4;5;6;7;8;9;10;10] |
| STEP([1:10],1.7) | [1;1;1.3;2.3;3.3;4.3;5.3;6.3;7.3;8.3] |
| | |
| Suppose matrix $M = \begin{pmatrix} 1 \\ 5 \\ 9 \end{pmatrix}$ | 2 3 4 |
| Suppose matrix $M = \begin{bmatrix} 5 \end{bmatrix}$ | 6 7 8 |
| (9) | 10 11 12 / |

| function | result |
|-------------|---|
| STEP(M,2) | $\left(\begin{array}{cccc} 1 & 2 & 3 & 4 \\ 1 & 2 & 3 & 4 \\ 1 & 2 & 3 & 4 \end{array}\right)$ |
| STEP(M,-1) | $ \left(\begin{array}{ccccc} 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \\ 9 & 10 & 11 & 12 \end{array}\right) $ |
| STEP(M,1.5) | $\left(\begin{array}{cccc} 1 & 2 & 3 & 4 \\ 1 & 2 & 3 & 4 \\ 3 & 4 & 5 & 6 \end{array}\right)$ |

WRAP

Syntax matrix = WRAP(matrix,scalar)

The WRAP function accepts a matrix as its first argument. It wraps the column elements of a matrix by the specified step size, scalar.

- If scalar = n > 0, the column elements of the matrix are shifted down. The first n elements in the first column are zero filled, then the last n elements of each column are brought into the next column. The last n elements in the last column are lost.
- If scalar = n = 0, the matrix is returned unchanged.
- If scalar = n < 0, the column elements of the matrix are shifted up. The first n elements in the first column are lost. The first n elements of each column are brought into the preceding column. The last n elements in the last column are zero filled.

If scalar is non-integral, it is truncated to an integer.

Suppose matrix
$$M = \begin{pmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{pmatrix}$$

Functions

| function | result |
|------------|--|
| WRAP(M,2) | $ \left(\begin{array}{cccc} 0 & 5 & 6 & 7 \\ 0 & 9 & 10 & 11 \\ 1 & 2 & 3 & 4 \end{array}\right) $ |
| | $\left(\begin{array}{cccc} 1 & 2 & 3 & 4 \end{array}\right)$ |
| | $ \left(\begin{array}{ccccc} 9 & 10 & 11 & 12 \\ 2 & 3 & 4 & 0 \end{array}\right) $ |
| WRAP(M,-2) | $\left(\begin{array}{cccc}2&3&4&0\\6&7&8&0\end{array}\right)$ |

Looping functions

Looping functions mimic standard mathematical notation, for example, the sum:

$$\mathrm{SUM}(f(j),j,1:N) \equiv \sum_{j=1}^N f(j)$$

Where j is the dummy variable.

The looping functions, SUM, PROD, RSUM, RPROD, and LOOP, require a previously declared scalar dummy variable as second argument. A dummy variable is declared with the SCALAR\DUMMY command. A dummy variable is different from other scalar variables in that its value is only defined while inside the looping function. The third argument of a looping function is always the range of this dummy variable, and must be a vector. The first argument of a looping function would normally be some function of the dummy variable, but it is not necessary that the dummy variable appear in the first argument.

SUM

```
Syntax scalar = SUM(scalar,dummy,scalar)
vector = SUM(vector,dummy,vector)
matrix = SUM(matrix,dummy,matrix)
```

The SUM function requires a previously declared scalar dummy variable as second argument. A dummy variable is declared with the SCALAR\DUMMY command. The third argument is always the range of this dummy variable, and must be a vector. The first argument would normally be some function of the dummy variable, but it is not necessary that the dummy variable appear in the first argument.

The SUM function accepts a scalar, a vector, or a matrix as its first argument. The SUM function is equivalent to $\sum_{i \in \mathcal{R}} \mathcal{F}$ where i is the dummy variable, \mathcal{R} is the range of the dummy variable, and \mathcal{F} is some function of the dummy variable.

If $\mathcal F$ is a scalar, the result is a scalar. If $\mathcal F$ is a vector, the result is a vector. If $\mathcal F$ is a matrix, the result is a matrix.

Examples

Suppose X=[1;2;3;4;5], Y=[2;3;4;5;6], M= $\begin{pmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{pmatrix}$ and I and J have been declared to be dummy variables with SCALAR\DUMMY I J.

| function | result |
|------------------------------|-----------------|
| SUM(X[I],I,1:5) | 15 |
| SUM(X,I,1:5) | [5;10;15;20;25] |
| $SUM((X^2+Y^2)[I],I,1:5)$ | 145 |
| SUM(SUM(M[I,J],I,1:3),J,1:4) | 78 |

PROD

The PROD function requires a previously declared scalar dummy variable as second argument. A dummy variable is declared with the SCALAR\DUMMY command. The third argument is always the range of this dummy variable, and must be a vector. The first argument would normally be some function of the dummy variable, but it is not necessary that the dummy variable appear in the first argument.

The PROD function accepts a scalar, a vector, or a matrix as its first argument. The PROD function is equivalent to $\prod_{i \in \mathcal{R}} \mathcal{F}$ where i is the dummy variable, \mathcal{R} is the range of the dummy variable, and \mathcal{F} is some function of the dummy variable.

If \mathcal{F} is a scalar, the result is a scalar. If \mathcal{F} is a vector, the result is a vector. If \mathcal{F} is a matrix, the result is a matrix.

Suppose X=[1;2;3;4;5], Y=[2;3;4;5;6], M=
$$\begin{pmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{pmatrix}$$
 and I and J have been declared to be dummy variables with SCALAR\DUMMY I J.

Functions

| function | | result |
|---|-------------------------|----------------------|
| PROD(X[I] |],I,1:5) | 120 |
| PROD(X,I | ,1:5) | [1;32;243;1024;3125] |
| PROD((X^2 | 2+Y^2)[I],I,1:5) | 4064125 |
| PROD (PROI | D(M[I,J],I,1:3),J,1:4) | 479001600 |
| RSUM | | |
| Syntax vector = RSUM(scalar,dummy,vector) | | |
| | matrix = RSUM(vector,du | ummy,matrix) |

The RSUM function requires a previously declared scalar dummy variable as second argument. A dummy variable is declared with the SCALAR\DUMMY command. The third argument is always the range of this dummy variable, and must be a vector. The first argument would normally be some function of the dummy variable, but it is not necessary that the dummy variable appear in the first argument.

The RSUM function accepts either a scalar or a vector as its first argument. The RSUM function calculates the running sums. If the first argument is a scalar, the last element of the RSUM output vector is the total sum. If the first argument is a vector, the last column of the RSUM output matrix is the total sum.

Suppose X=[1;2;3;4;5], Y=[2;3;4;5;6], M=
$$\begin{pmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{pmatrix}$$
 and I and J have been declared to be dummy variables with SCALAR\DUMMY I J.

| function | result |
|--------------------------------|--|
| RSUM(X[I],I,1:5) | [1;3;6;10;15] |
| RSUM(X,I,1:3) | $ \begin{pmatrix} 1 & 2 & 3 \\ 2 & 4 & 6 \\ 3 & 6 & 9 \\ 4 & 8 & 12 \\ 5 & 10 & 15 \end{pmatrix} $ |
| RSUM((X^2+Y^2)[I],I,1:5) | [5;18;43;84;145] |
| RSUM(RSUM(M[I,J],I,1:3),J,1:4) | $ \left(\begin{array}{cccc} 1 & 3 & 6 & 10 \\ 6 & 14 & 24 & 36 \\ 15 & 33 & 54 & 78 \end{array}\right) $ |

RPROD

```
Syntax vector = RPROD(scalar,dummy,vector)
matrix = RPROD(vector,dummy,matrix)
```

The RPROD function requires a previously declared scalar dummy variable as second argument. A dummy variable is declared with the SCALAR\DUMMY command. The third argument is always the range of this dummy variable, and must be a vector. The first argument would normally be some function of the dummy variable, but it is not necessary that the dummy variable appear in the first argument.

The RPROD function accepts either a scalar or a vector as its first argument. The RPROD function calculates the running products. If the first argument is a scalar, the last element of the RPROD output vector is the total product. If the first argument is a vector, the last column of the RPROD output matrix is the total product.

Examples

Suppose X=[1;2;3;4;5], Y=[2;3;4;5;6], M= $\begin{pmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{pmatrix}$ and I and J have been declared to be dummy variables with SCALAR\DUMMY I J.

| function | result |
|----------------------------------|---|
| RPROD(X[I],I,1:5) | [1;2;6;24;120] |
| RPROD(X,I,1:3) | $ \begin{pmatrix} 1 & 1 & 1 \\ 2 & 4 & 8 \\ 3 & 9 & 27 \\ 4 & 16 & 64 \\ 5 & 25 & 125 \end{pmatrix} $ |
| RPROD((X^2+Y^2)[I],I,1:5) | [5;65;1625;66625;4064125] |
| RPROD(RPROD(M[I,J],I,1:3),J,1:4) | $ \left(\begin{array}{cccc} 1 & 2 & 6 & 24 \\ 5 & 60 & 1260 & 40320 \\ 45 & 5400 & 1247400 & 479001600 \end{array}\right) $ |
| | LOOP |

The LOOP function requires a previously declared scalar dummy variable as second argument. A dummy variable is declared with the SCALAR\DUMMY command. The third argument

Functions

is always the range of this dummy variable, and must be a vector. The first argument would normally be some function of the dummy variable, but it is not necessary that the dummy variable appear in the first argument.

The LOOP function accepts either a scalar or a vector as its first argument. The LOOP function simply loops over the range of the dummy variable, filling the output with the appropriate value from the first argument. The output of the LOOP function will always have one dimension more than the first argument. If the first argument is a scalar, the output will be the elements of a vector. If the first argument is a vector, the output will be the columns of a matrix.

Examples

Suppose X=[1;2;3;4;5], Y=[2;3;4;5;6], M=
$$\begin{pmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \\ 13 & 14 & 15 & 16 \end{pmatrix}$$
 and I and J have been de-

clared to be dummy variables with SCALAR\DUMMY I J.

| function | result |
|--------------------------------|--|
| LOOP(X[I]*Y[(6-I)],I,1:5) | [6;10;12;12;10] |
| | $\left(\begin{array}{ccc}1&1&1\\2&2&2\end{array}\right)$ |
| LOOP(X,I,1:3) | 3 3 3 |
| | $ \begin{pmatrix} 1 & 1 & 1 \\ 2 & 2 & 2 \\ 3 & 3 & 3 \\ 4 & 4 & 4 \\ 5 & 5 & 5 \end{pmatrix} $ |
| LOOP(M[I,I],I,1:4) | [1;6;11;16] |
| LOOP(LOOP(M[I,J],I,1:J),J,1:4) | $ \left(\begin{array}{ccccc} 1 & 2 & 3 & 4 \\ 0 & 6 & 7 & 8 \\ 0 & 0 & 11 & 12 \\ 0 & 0 & 0 & 16 \end{array}\right) $ |
| LOOP(LOOP(M[J,I],I,1:J),J,1:4) | $ \left(\begin{array}{ccccc} 1 & 5 & 9 & 13 \\ 0 & 6 & 10 & 14 \\ 0 & 0 & 11 & 15 \\ 0 & 0 & 0 & 16 \end{array}\right) $ |

Suppose M=
$$\begin{pmatrix} 11 & 12 & 13 \\ 21 & 22 & 23 \\ 31 & 32 & 33 \end{pmatrix}$$
 and J has been declared to be a dummy variable with SCALAR\DUMMY J.

To invert the matrix, that is, to flip the matrix upside down, use:

| function result | |
|-------------------------|--|
| | (31 32 33) |
| <-LOOP(M[I,*],I,3:1:-1) | 21 22 23 |
| | $\begin{pmatrix} 11 & 12 & 13 \end{pmatrix}$ |

You must use the transpose operator, since M[I,*] is a vector, and each time LOOP iterates on I, it produces a column of the output matrix.

GPLOT KEYWORDS

This chapter contains detailed descriptions of the GPLOT graph and text plot characteristic keywords, which are controlled by the SET and GET commands.

The values associated with every name are converted internally to floating point real, and all angles are in degrees.

A.1 Summary

Following is a list of plot characteristic keywords, with very terse descriptions and default values.

General

| name | description | default value | |
|--------|---|---------------|---|
| PTYPE | controls whether pixels are turned on, off, or complemented | 0 | |
| LINTYP | line type | 1 | |
| LINTHK | line thickness | 1 | |
| COLOUR | colour | 1 | |
| NUMBLD | fill number for the axis numbers | 0 | |
| CLIP | controls whether data curves are clipped at the box edge | 1 | |
| HISTYP | histogram type | 0 | |
| CHARA | plotting symbol angle | horizontal | |
| CHARSZ | plotting symbol size | 1 | % |

Text

| name | description | default value |
|--------|--|---------------|
| CURSOR | text justification | 1 |
| TXTANG | text angle | 0 |
| TXTHIT | text height | 3 % |
| XLOC | horizontal reference location for text positioning | 50 % |
| YLOC | vertical reference location for text positioning | 50 % |

x-axis

| name | description | default value | |
|--------|---|---------------------|---|
| XAXIS | controls whether or not to draw the x-axis | 1 | |
| XLABSZ | height of the x -axis text label | 3 | % |
| XLOG | base of the <i>x</i> -axis numbers | 0 | |
| NXGRID | number of grid lines to draw parallel to the y -axis | 0 | |
| XCROSS | controls where the y -axis will cross the x -axis | 0 | |
| XZERO | controls whether zero is forced to appear on the x -axis | 0 | |
| XTICTP | type of tic marks to place on the x-axis | 1 | |
| XTICA | angle of the x -axis tic marks | $\parallel y$ -axis | |
| NLXINC | number of long x -axis tic marks | 2 | |
| XTICL | length of the long tic marks on the <i>x</i> -axis | 2 | % |
| NSXINC | number of short x -axis tic marks | 1 | |
| XTICS | length of the short tic marks on the <i>x</i> -axis | 1 | % |
| XAMX | maximum value for the <i>x</i> -axis | 10 | |
| XVMAX | virtual maximum for the x -axis | 10 | |
| XMIN | minimum value for the x -axis | 0 | |
| XVMIN | virtual minimum for the x -axis | 0 | |
| XMOD | base of the modulus for x -axis numbering | 0 | |
| XOFF | offset added to the numbers labeling the x -axis | 0 | |
| XLEADZ | controls whether x -axis leading zeros are displayed | 0 | |
| XPAUTO | controls the automatic x -axis scale factor | 1 | |
| XPOW | x-axis numbers scale factor | 0 | |
| NXDIG | number of digits to display in the x -axis numbers | 5 | |
| NXDEC | number of decimal places to display in the <i>x</i> -axis numbers | -1 | |
| XNUMSZ | height of the numbers labeling the <i>x</i> -axis | 3 | % |
| XNUMA | angle of the numbers labeling the x -axis | horizontal | |
| XITICA | angle at which to position the x -axis numbers | $\parallel y$ -axis | |
| XITICL | distance from the x -axis to the numbers labeling the x -axis | 3 | % |

y-axis

| name | description | default value | |
|--------|---|---------------------|---|
| YAXIS | controls whether or not to draw the y-axis | 1 | |
| YLABSZ | height of the y -axis text label | 3 | % |
| YLOG | controls whether the y -axis is to be linear or logarithmic | 0 | |
| NYGRID | number of grid lines to draw parallel to the x-axis | 0 | |
| YCROSS | controls where the x -axis will cross the y -axis | 0 | |
| YZERO | controls whether zero is forced to appear on the y -axis | 0 | |
| YTICTP | type of tic marks to place on the y -axis | 1 | |
| YTICA | angle of the y -axis tic marks | $\parallel x$ -axis | |
| NLYINC | number of long y -axis tic marks | 2 | |
| YTICL | length of the long tic marks on the y -axis | 2 | % |
| NSYINC | number of short y -axis tic marks | 1 | |
| YTICS | length of the short tic marks on the y-axis | 1 | % |
| YMAX | maximum value for the y -axis | 10 | |
| XAMVY | virtual maximum for the y -axis | 10 | |
| YMIN | minimum value for the y -axis | 0 | |
| YVMIN | virtual minimum value for the y -axis | 0 | |
| YMOD | base of the modulus for the <i>y</i> -axis numbering | 0 | |
| YOFF | offset added to the numbers labeling the y -axis | 0 | |
| YLEADZ | controls whether y -axis leading zeros are displayed | 0 | |
| YPAUTO | controls the automatic y -axis scale factor | 1 | |
| YPOW | y-axis numbers scale factor | 0 | |
| NYDIG | number of digits to display in the <i>y</i> -axis numbers | 5 | |
| NYDEC | number of decimal places to display in the y -axis numbers | -1 | |
| YNUMSZ | height of the numbers labeling the <i>y</i> -axis | 3 | % |
| YNUMA | angle of the numbers labeling the y -axis | horizontal | |
| YITICA | angle at which to position the y -axis numbers | $\parallel y$ -axis | |
| YITICL | distance from the y -axis to the numbers labeling the y -axis | 3 | % |

Axis Box

| name | description | default value | |
|--------|---|---------------|---|
| XLWIND | the left edge of the window | 0 | % |
| XUWIND | the right edge of the window | 100 | % |
| YLWIND | the bottom edge of the window | 0 | % |
| YUWIND | the top edge of the window | 100 | % |
| BOX | controls whether or not to draw an axis box around the graph | 1 | |
| XLAXIS | location of the lower end of the x -axis | 15 | % |
| XUAXIS | location of the upper end of the x -axis | 95 | % |
| XAXISA | angle of the x-axis | 0 | |
| YLAXIS | location of the bottom of the y -axis | 15 | % |
| YUAXIS | location of the top of the y -axis | 90 | % |
| YAXISA | angle of the y -axis | 90 | |
| BOTNUM | controls the height of the numbers on the bottom of the box | 0 | |
| BOTTIC | controls the length of the tic marks on the bottom of the box | 1 | |
| RITNUM | controls the height of the numbers on the right side of the box | 0 | |
| RITTIC | controls the length of the tic marks on the right side of the box | -1 | |
| TOPNUM | controls the height of the numbers on the top of the box | 0 | |
| TOPTIC | controls the length of the tic marks on the top of the box | -1 | |
| LEFNUM | controls the height of the numbers on the left side of the box | 0 | |
| LEFTIC | length of the tic marks on the left side of the box | 1 | |

A.2 General Characteristics

PTYPE

Default: PTYPE = 0

PTYPE controls whether graphics pixels are turned on, off, or toggled. This can be used to selectively erase graphics, by drawing with PTYPE =0 and redrawing with PTYPE =1; or by drawing and redrawing with PTYPE =2. This only works on the monitor screen and on bitmap hardcopy output. Values other than 0, 1, 2 are ignored.

- 0 pixels turned on (draw)
- 1 pixels turned off (erase)
- 2 pixels complemented

Complemented pixels means that a pixel is turned off if it is on, or turned off if it is on.

LINTYP

Default: LINTYP = 1

LINTYP controls the type of line to use when drawing a data curve. LINTYP should be between 1 and 10, inclusive. Table 2.8 on page 30 shows the specifications for the default line types. The length of the dashes in the dashed line types, the default types 3 to 10, are constant and do not depend on the separation between data points. The definition of a line type can be changed via the SET LINE command.

LINTHK

Default: LINTHK = 1

LINTHK controls the line thickness for bitmap hardcopy output and for PostScript hardcopy output. LINTHK has no affect on monitor screen output or on pen plotter hardcopy output.

COLOUR

Default: COLOUR = 1

COLOUR sets a colour code which is used to control the monitor screen colour and the hard-copy colour. See Table 2.6 on page 20 for a list of the colour names and associated colour numbers.

NUMBLD

Default: NUMBLD = 0

NUMBLD is the fill number for the axis numbers.

```
\begin{array}{ll} 0 & \text{no filling} \\ 1 \leq \texttt{NUMBLD} \leq 10 & \text{use hatch pattern} \\ 11 \leq \texttt{NUMBLD} \leq 99 & \text{use dot fill pattern} \end{array}
```

CLIP

Default: CLIP = 1

CLIP controls whether or not data curves that are plotted are clipped at the boundaries of the axis box.

```
0 do not clip \neq 0 clip at the boundaries of the axis box
```

HISTYP

Default: HISTYP = 0

HISTYP controls whether a normal line graph or a histogram is drawn. Histograms can have tails or no tails and the profile can be along the x-axis or along the y-axis. See Figure 2.16 on page 123.

- 0 normal line graph, not a histogram
- 1 histogram with no tails and profile along the x-axis.
 - Can control the width and colour of each individual bar
- 2 histogram with tails to y = 0 and profile along the x-axis. Can control the filling pattern, width and colour of each individual bar
- 3 histogram without tails and profile along the y-axis. Can control the height and colour of each individual bar
- 4 histogram with tails to x=0 and profile along the y-axis. Can control the filling pattern, height and colour of each individual bar

No plotting symbol will be drawn at the data points when $\mathtt{HISTYP} > 0$. The fill pattern, width and colour of each histogram bar can be controlled by using the optional arrays in the SET PCHAR command.

CHARA

Default: %CHARA = XAXISA

CHARA is the angle, in degrees, of the plotting symbols that are drawn at the data points, measured counterclockwise between a horizontal line and a base line through the plotting symbol. CHARA will not be used when plotting if the optional angle array is included with the SET PCHAR command.

If CHARA is set as a percentage, the actual value to which %CHARA has been set is ignored and the angle is set to XAXISA. This allows the user to change the angle of the x-axis and keep the plotting symbols base line parallel to the x-axis. By default, CHARA is set as a percentage.

CHARSZ

Default: %CHARSZ = 1

CHARSZ is the size of the plotting symbols that are drawn at the data points. $\mbox{\ensuremath{\texttt{CHARSZ}}}$ is a percentage of the height of the window, that is, $\mbox{\ensuremath{\texttt{CHARSZ}}} = \mbox{\ensuremath{\texttt{\texttt{CHARSZ}}}} \times (\mbox{\ensuremath{\texttt{YUWIND}}} - \mbox{\ensuremath{\texttt{YLWIND}}}) \div 100$ CHARSZ will be the base size of the plotting symbols if the optional size array is included in the SET PCHAR command.

A.3 Text

CURSOR

Default: CURSOR = 1

CURSOR controls the justification of the text that is drawn when using TEXT command. The origin of the text is always the lower left corner of the string. The justification determines where this origin is placed with respect to a reference point. Refer to Figure 2.28 on page 258 and to Table 2.64 on page 259.

- ≤ 0 then XLOC and YLOC will be used to determine the reference point for the text, and the justification will be determined by |CURSOR|.
- > 0 the user selects a reference point with the graphics cursor. The justification of the text is selected interactively. Refer to Table 2.65 on page 260.

The value of CURSOR will be updated to the value corresponding to the justification that was chosen. The values of %XLOC and %YLOC will be updated to the new reference point location.

TXTANG

Default: TXTANG = 0

TXTANG controls the angle at which text will be drawn, when using the TEXT command. TXTANG is the angle, in degrees, measured counterclockwise, between the base line of the text and a horizontal line. The value of TXTANG will be ignored if |CURSOR| = 4, 5, or 6.

TXTHIT

Default: %TXTHIT = 3

TXTHIT is the height of text to be drawn, when using the TEXT command. %TXTHIT is a percentage of the height of the window, that is, TXTHIT = $%TXTHIT \times (YUWIND - YLWIND) \div 100$

XLOC

Default: %XLOC = 50

XLOC is the horizontal reference position of a text drawn using the TEXT command. This reference point is used for text justification. See the explanation of CURSOR for how XLOC will be used. XLOC will be used if CURSOR < 0, or if the X key is typed when drawing text in interactive mode. The value of %XLOC is updated after each string is drawn with the TEXT command. %XLOC is a percentage of the width of the window, that is, XLOC = XLWIND + %XLOC \times (XUWIND - XLWIND) \div 100

YLOC

Default: %YLOC = 50

YLOC controls the vertical reference position of a text string drawn with the TEXT command. This reference point is used for text justification. See the explanation of CURSOR for how YLOC will be used. YLOC will be used if CURSOR < 0, or if the Y key is typed when drawing text in interactive mode. The value of %YLOC is updated after each string is drawn with the TEXT command. %YLOC is a percentage of the height of the window, that is, YLOC = YLWIND + %YLOC \times (YUWIND - YLWIND) \div 100

A.4 x-axis

Figure A.36 illustrates the definitions of some of the *x*-axis characteristics.

XAXIS

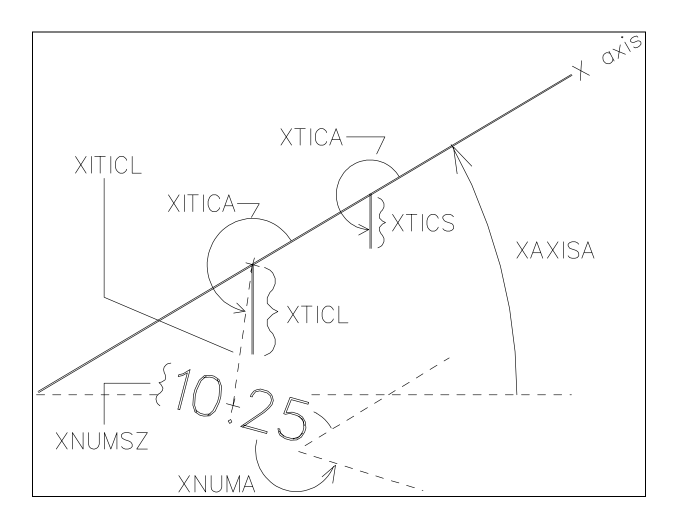


Figure A.36: Some *x*-axis characteristics

Default: XAXIS = 1

XAXIS controls whether or not the x-axis is drawn.

0 do not draw the x-axis $\neq 0$ draw the x-axis

If BOX $\neq 0$, then the bottom of the axis box will be drawn, even if XAXIS = 0.

XLABSZ

Default: %XLABSZ = 3

XLABSZ controls the height of the automatic text label for the x-axis, set by the LABEL\XAXIS command. %XLABSZ is a percentage of the height of the window, that is, XLABSZ = %XLABSZ \times (YUWIND – YLWIND) \div 100

XLOG

Default: XLOG = 0

XLOG determines whether the *x*-axis is to be linear or logarithmic. See Figure A.37 for examples of various logarithmic axes.

the x-axis will have a logarithmic scale. The numbers labeling the x-axis will be in exponent form, for example, $10^1\ 10^2\ 10^3\ 10^4$.

the x-axis will have a linear scale <-1 the x-axis will have a logarithmic scale. The numbers labeling the x-axis will be in full digit form, for example, $10\ 100\ 1000\ 10000$.

Suppose that $|\mathtt{XLOG}| > 1$. The base will be the *integer part* of $|\mathtt{XLOG}|$, except for the special case: $1.05 \times e > \mathtt{XLOG} > 0.95 \times e$, where e is the base of the natural logarithms, $e \approx 2.718281828$, in which case the base will be e. XMIN and XMAX will be the *exponents* for the minimum and maximum values on the x-axis. The maximum value displayed on the x-axis is $|\mathtt{XLOG}|^{\mathtt{XMAX}}$ and the minimum value displayed on the x-axis is $|\mathtt{XLOG}|^{\mathtt{XMIN}}$. Integer exponents are always used for the axis numbering, so the x-axis may not begin and/or end on a large tic mark.

If $|\mathtt{XLOG}|=10$, the small tic marks on the x-axis can be specified exactly with NSXINC. If small tic marks are desired at the locations $j_m \times 10^n$, where $2 \le j_m \le 9$, then set NSXINC $= -j_1 \dots j_m$. For example, if you want small tic marks at 2×10^n , 4×10^n , 5×10^n , and 8×10^n , then set NSXINC = -2458.

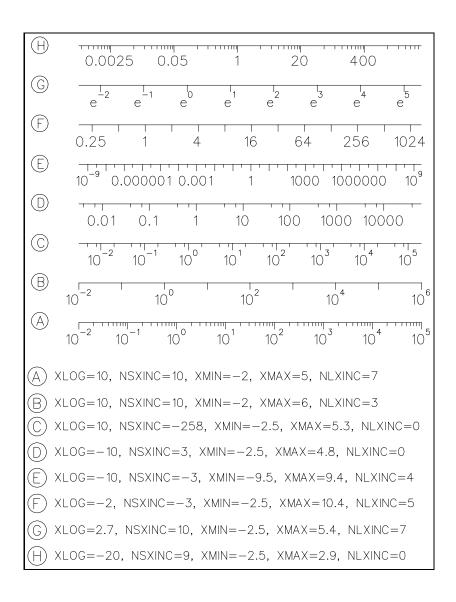


Figure A.37: Logarithmic *x*-axis examples

NXGRID

Default: NXGRID = 0

NXGRID controls the number of grid lines parallel to the y-axis. NXGRID specifies the number of large x-axis tic marks between each grid line.

- >0 grid lines parallel to the y-axis at every NXGRID large tic mark, starting with the first
- 0 suppress all grid lines parallel to the *y*-axis
- -1 grid lines parallel to the y-axis at every x-axis tic mark, large and small. This option applies only if the axes are orthogonal.

XCROSS

Default: XCROSS = 0

XCROSS controls where the y-axis is to cross the x-axis. The axes always cross at a large tic mark.

- 0 the y-axis will cross the x-axis at XMIN
- $\neq 0$ the y-axis will cross the x-axis at the large tic mark closest to x=0

XZERO

Default: XZERO = 0

XZERO controls whether or not to force zero to be displayed on the x-axis. XZERO is set to zero after each graph is drawn.

- $\neq 0$ if XMIN ≥ 0 then set XMIN to zero, or, if XMAX ≤ 0 then set XMAX to zero
- 0 do not force zero to be displayed on the *x*-axis

XTICTP

Default: XTICTP = 1

XTICTP controls the type of tic marks to place on the x-axis.

- 2 tic marks on both sides of the x-axis
- $\neq 2$ tic marks on only one side of the x-axis. XTICA controls the side on which the tic marks will appear

XTICA

Default: $XTICA = YAXISA - XAXISA + 180^{\circ}$

XTICA is the angle, in degrees, measured counterclockwise, between a horizontal line and a tic mark, both large and small, on the x-axis. See Figure A.36. By default, XTICA is set as a percentage. If XTICA is set as a percentage, then XTICA is set to YAXISA — XAXISA — XAXISA + 180. The actual value of %XTICA will be ignored. This allows the user to change the angle of the axes and keep the x-axis tic marks parallel to the y-axis.

NLXINC

Default: NLXINC = 2

NLXINC controls the number of large tic marks to be displayed on the x-axis.

- -1000 drop the first and the last numbers on the x-axis and determine the number of large x-axis tic marks
- <0 drop the first and the last numbers on the $x\mbox{-axis}$. The number of large tic marks on the $x\mbox{-axis}$ will be $|\mbox{\tt NLXINC}|+1$, unless $|\mbox{\tt XLOG}|>1$, in which case the number of tic marks will be determined to avoid fractional powers
- 0 the number of large tic marks on the x-axis will be automatically determined
- >0 the number of large tic marks on the $x\mbox{-axis}$ will be $\mbox{NLXINC}+1,$ unless $|\mbox{XLOG}|>1,$ in which case NLXINC will be determined to avoid fractional powers

XTICL

Default: %XTICL = 2

XTICL is the length of the long tic marks on the x-axis. See Figure A.36 on page 356. %XTICL is a percentage of the height of the window, that is, XTICL = %XTICL \times (YUWIND – YLWIND) \div 100

NSXINC

Default: NSXINC = 1

NSXINC controls the number of small tic marks to be displayed on the x-axis. The small tic marks appear between the large tic marks.

- < 1 no small tic marks on the x-axis
- ≥ 2 the number of small tic marks, on the *x*-axis, between each pair of large tic marks, will be NSXINC -1

If $|\mathtt{XLOG}| = 10$, the small tic marks on the x-axis can be specified exactly with NSXINC. If small tic marks are desired at the locations $j_m \times 10^n$, where $2 \le j_m \le 9$, then set NSXINC $= -j_1 \dots j_m$. For example, if you want small tic marks at 2×10^n , 4×10^n , 5×10^n , and 8×10^n , then set NSXINC = -2458.

XTICS

Default: %XTICS = 1

XTICS controls the length of the short tic marks on the x-axis. See Figure A.36 on page 356. %XTICS is a percentage of the height of the window, that is, XTICS = %XTICS \times (YUWIND – YLWIND) \div 100

XMAX

Default: XMAX = 10

XMAX controls the maximum value for the x-axis.

If $|\mathtt{XLOG}| > 1$, then XMAX is assumed to be the exponent for the maximum value on the x-axis. The maximum value displayed on the x-axis is $|\mathtt{XLOG}|^{\mathtt{XMAX}}$. Integer exponents are always used for the axis numbering, so the x-axis may not end on a large tic mark.

If $|XLOG| \le 1$, then XMAX is the actual maximum value for the x-axis. If XVMAX is equal to XMAX then the x-axis always ends on a large, labeled, tic mark. If XVMAX is not equal to XMAX then the x-axis will not end at a large, labeled, tic mark.

When the value of XMAX is changed, the value of XVMAX is simultaneously changed to the same value.

XVMAX

Default: XVMAX = 10

XVMAX controls the virtual maximum value for the x-axis. See Figure A.38 on page 363.

If |XLOG| > 1, the virtual maximum is ignored.

If $|\mathtt{XLOG}| \leq 1$, then XVMAX is the virtual maximum value of the labels for the x-axis. This value will not be displayed if XVMAX is greater than XMAX, but it will be displayed if XVMAX is less than or equal to XMAX. This virtual maximum is used to determine "nice" numbers for labeling the large tic marks on the x-axis. If XVMAX is not equal to XMAX then the x-axis will not end on a large, labeled, tic mark.

The value of XVMAX is changed to the value of XMAX when the value of XMAX is changed. So, if you want to make XVMAX different from XMAX, it must be changed *after* XMAX is changed.

If NLXINC is set to zero, then the x-axis will begin at XMIN and end at XMAX, but if these are not 'nice' numbers, the virtual minimum and maximum will be set to values outside the actual minimum and maximum so that the large tic marks can be labeled with "nice" numbers.

XMIN

Default: XMIN = 0

XMIN controls the minimum value for the x-axis.

If $|\mathtt{XLOG}| > 1$, then XMIN is assumed to be the exponent for the minimum value on the x-axis. The minimum value displayed on the x-axis is $|\mathtt{XLOG}|^{\mathtt{XMIN}}$. Integer exponents are always be used for the axis numbering, so the x-axis may not begin on a large tic mark.

If $|XLOG| \le 1$, XMIN is the actual minimum value for the x-axis. If XVMIN is equal to XMIN then the x-axis always begins on a large, labeled, tic mark. If XVMIN is not equal to XMIN then the x-axis will not begin at a large, labeled, tic mark.

When the value of XMIN is changed, the value of XVMIN is simultaneously changed to the same value.

XVMIN

Default: XVMIN = 0

XVMIN controls the virtual minimum value for the *x*-axis. See Figure A.38 on page 363.

If |XLOG| > 1, the virtual minimum is ignored.

If $|XLOG| \le 1$, then XVMIN is the virtual minimum value of the labels for the x-axis. This value will not be displayed if XVMIN is less than XMIN, but it will be displayed if XVMIN is greater than

or equal to XMIN. This virtual minimum is used to determine "nice" numbers for labeling the large tic marks on the x-axis. If XVMIN is not equal to XMIN then the x-axis will not begin on a large, labeled, tic mark.

The value of XVMIN is changed to the value of XMIN when the value of XMIN is changed. So, if you want to make XVMIN different from XMIN, it must be changed *after* XMIN is changed.

If NLXINC is set to zero, then the x-axis will begin at XMIN and end at XMAX, but if these are not "nice" numbers, the virtual minimum and maximum will be set to values outside the actual minimum and maximum so that the large tic marks can be labeled with "nice" numbers.

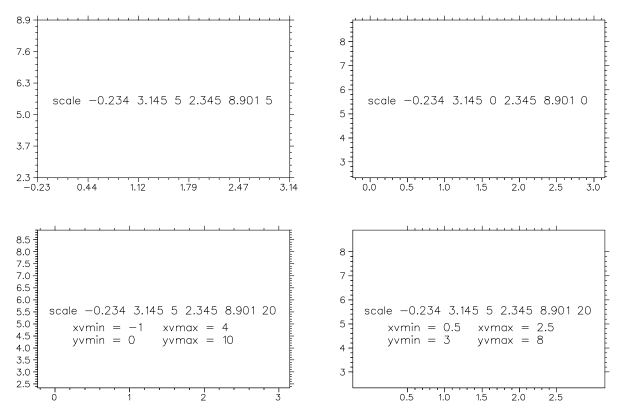


Figure A.38: Virtual axes examples

XMOD

Default: XMOD = 0

XMOD is another control on the numbering for the x-axis.

 $> 1 \qquad \qquad \text{the numbers displayed on the x-axis are modulo XMOD and positive.}$ If the number to be displayed on the axis is \$x\$ then the number that is actually displayed is $|x-{\tt XMOD}\times(x/{\tt XMOD})| + {\tt XOFF}$ $-1 \leq {\tt XMOD} \leq 1 \qquad \text{the numbers will be displayed normally}$ $< -1 \qquad \text{the numbers will be modulo } |{\tt XMOD}| \text{ and may be negative}$

XOFF

Default: XOFF = 0

XOFF is another control on the numbering for the x-axis. If |XMOD| > 1, then XOFF is added to the numbers to be displayed on the x-axis. If $|XMOD| \le 1$, then XOFF is ignored.

XLEADZ

Default: XLEADZ = 0

XLEADZ controls whether leading zeros are displayed on the x-axis numbers.

- 1 numbers will have leading zeros if they are between 0 and 1
- 0 numbers will *not* have leading zeros if they are between 0 and 1

The default is to not display these leading zeros. Numbers that are between -1 and 0 always have the leading zero displayed.

XPAUTO

Default value: XPAUTO = 1

XPAUTO, along with XPOW, controls the x-axis scale factor that is appended to the x-axis text label. This scale factor is needed when there are not enough digits allowed for the numbers labeling the x-axis. XPOW is only appended to the text label if XPOW $\neq 0$.

- 2 determine XPOW, but do *not* append the scale factor to the text label
- determine XPOW, and append the scale factor to the text label
- 0 use the present value of XPOW

If the user wishes to completely specify the appearance of the x-axis, XPAUTO must be set to 0, otherwise the number of digits and decimal places, NXDIG and NXDEC, may be changed.

XPOW

Default: XPOW = 0

XPOW controls the x-axis scale factor that is appended to the x-axis text label. This scale factor is a power of ten, that is, 10^{XPOW} , and the numbers labeling the x-axis should be multiplied by this scale factor to get the correct graph units. XPOW is only appended to the text label if XPOW $\neq 0$.

NXDIG

Default: NXDIG = 5

NXDIG controls the total number of digits to be displayed in each of the numbers labeling the x-axis. If NXDIG is smaller than required to display the x-axis numbers, NXDIG will not be increased, but a scale factor, $(\times 10^n)$, will be appended to the x-axis text label. If NXDIG is larger than required, NXDIG will be reduced to the minimum value required. The value of NXDIG is updated after each graph is drawn.

NXDEC

Default: NXDEC = -1

NXDEC controls the number of digits to be displayed in the fractional parts of the numbers labeling the x-axis. The value of NXDEC is updated after each graph is drawn.

- -1 display the numbers labeling the x-axis as integers, with no decimal point
- display the numbers labeling the x-axis with no fractional part, but with a decimal point
- >0 display the numbers labeling the x-axis with NXDEC digits in the fractional part

XNUMSZ

Default: %XNUMSZ = 3

XNUMSZ controls the size of the numbers labeling the x-axis. See Figure A.36. %XNUMSZ is a percentage of the height of the window, that is, XNUMSZ = %XNUMSZ \times (YUWIND – YLWIND) \div 100

XNUMA

Default: XNUMA = -XAXISA

XNUMA controls the angle of the base line for the numbers labeling the x-axis. XNUMA is the angle, in degrees, measured counterclockwise, between a line parallel to the x-axis and the base line of a number. Refer to Figure A.36 on page 356. By default, XNUMA is set as a percentage. If XNUMA is set as a percentage, then XNUMA is set to -XAXISA. The actual value of %XNUMA will be ignored. This allows the user to change the angle of the x-axis and keep the base line of the x-axis numbers horizontal.

XITICA

Default: XITICA = YAXISA - XAXISA + 180

XITICA, along with XITICL, controls the location of the numbers labeling the x-axis at the large tic marks. XITICA is the angle, in degrees, measured counterclockwise, between the x-axis and a line joining the base of each large tic mark on the x-axis to the centre of the number labeling that tic mark. See Figure A.36. By default, XITICA is set as a percentage. If XITICA is set as a percentage, then XITICA is set to YAXISA — XAXISA + 180. The actual value of %XITICA will be ignored. This allows the user to change the angle of the axes and keep the relative locations of the x-axis numbers the same.

XITICL

Default: %XITICL = 3

XITICL, along with XITICA, controls the location of the numbers labeling the x-axis at the large tic marks. XITICL is the distance from the base of each large tic mark on the x-axis, to the centre of the number labeling that tic mark. See Figure A.36 on page 356. %XITICL is a percentage of the height of the window, that is, XITICL = %XITICL \times (YUWIND – YLWIND) \div 100

A.5 y-axis

Figure A.39 illustrates the definitions of some of the *y*-axis characteristics.

YAXIS

Default: YAXIS = 1

YAXIS controls whether or not the y-axis is drawn.

 $\neq 0$ draw the y-axis

0 do not draw the *y*-axis

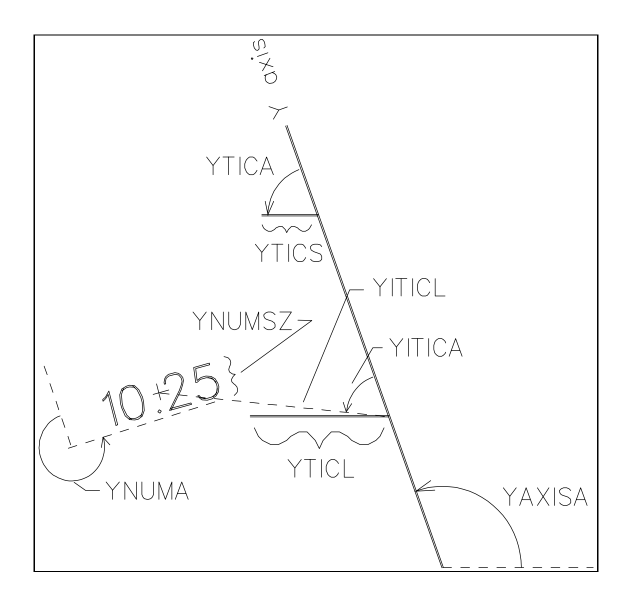


Figure A.39: Some *y*-axis characteristics

If BOX $\neq 0$, then the left side of the axis box will be drawn, even if YAXIS = 0.

YLABSZ

Default: %YLABSZ = 3

YLABSZ controls the size of the automatic text label for the y-axis, set by the LABEL\YAXIS command. %YLABSZ is a percentage of the height of the window, that is, YLABSZ = %YLABSZ \times (YUWIND - YLWIND) \div 100

YLOG

Default: YLOG = 0

YLOG determines whether the *y*-axis is to be linear or logarithmic. See Figure A.40 for examples of various logarithmic axes.

the y-axis will have a logarithmic scale. The numbers labeling the y-axis will be in exponent form, for example, $10^1\ 10^2\ 10^3\ 10^4$.

the y-axis will have a linear scale <-1 the y-axis will have a logarithmic scale. The numbers labeling the y-axis will be in full digit form, for example, $10\ 100\ 1000\ 10000$.

Suppose that $|\mathtt{YLOG}| > 1$. The base will be the *integer part* of $|\mathtt{YLOG}|$, except for the special case: $1.05 \times e > \mathtt{YLOG} > 0.95 \times e$, where e is the base of the natural logarithms, $e \approx 2.718281828$, in which case the base will be e. YMIN and YMAX will be the *exponents* for the minimum and maximum values on the y-axis. The maximum value displayed on the y-axis is $|\mathtt{YLOG}|^{\mathtt{YMAX}}$ and the minimum value displayed on the y-axis is $|\mathtt{YLOG}|^{\mathtt{YMIN}}$. Integer exponents are always used for the axis numbering, so the y-axis may not begin and/or end on a large tic mark.

If |YLOG| = 10, the small tic marks on the y-axis can be specified exactly with NSYINC. If small tic marks are desired at the locations $j_m \times 10^n$, where $2 \le j_m \le 9$, then set NSYINC $= -j_1 \dots j_m$. For example, if you want small tic marks at 2×10^n , 4×10^n , 5×10^n , and 8×10^n , then set NSYINC = -2458.

NYGRID

Default: NYGRID = 0

NYGRID controls the number of grid lines parallel to the x-axis. NYGRID specifies the number of large y-axis tic marks between each grid line.

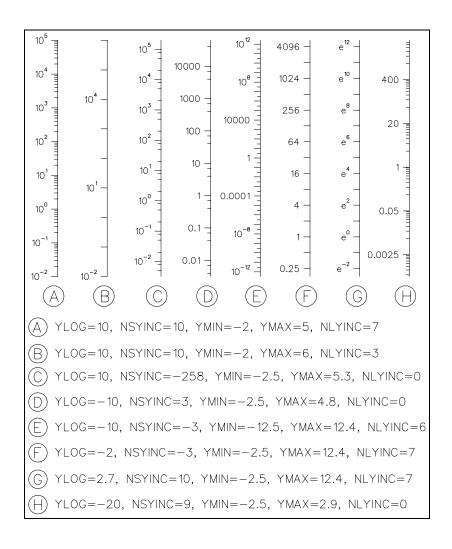


Figure A.40: Logarithmic *y*-axis examples

- >0 draw a grid line parallel to the x-axis at every NYGRID large tic mark, starting with the first
- 0 suppress all grid lines parallel to the *x*-axis
- -1 draw a grid line parallel to the x-axis at every y-axis tic mark, large and small. This option applies only if the axes are orthogonal.

YCROSS

Default: YCROSS = 0

YCROSS controls where the x-axis is to cross the y-axis. The axes always cross at a large tic mark.

- 0 the x-axis will cross the y-axis at YMIN
- $\neq 0$ the x-axis will cross the y-axis at y = 0, or at the large tic mark closest to y = 0

YZERO

Default: YZERO = 0

YZERO controls whether or not to force zero to be displayed on the y-axis. YZERO is set to zero after each graph is drawn.

- $\neq 0$ if YMIN ≥ 0 then set YMIN to zero, or, if YMAX ≤ 0 then set YMAX to zero
- 0 do not force zero to be displayed on the y-axis

YTICTP

Default: YTICTP = 1

YTICTP controls the type of tic marks to place on the y-axis.

- $\neq 2$ tic marks are drawn on only one side of the y-axis. YTICA controls the side on which the tic marks will appear
- 2 tic marks are drawn on both sides of the *y*-axis

YTICA

Default: YTICA = XAXISA - YAXISA + 180

YTICA controls the angle of the tic marks, both large and small, on the y-axis. YTICA is the angle, in degrees, measured counterclockwise, between the y-axis and a tic mark. See Figure A.39 on page 367. By default, YTICA is set as a percentage. If YTICA is set as a percentage, for example, SET %YTICA 0, then YTICA is set to XAXISA — YAXISA — YAXISA — The actual value of %YTICA will be ignored. This allows the user to change the angle of the axes and keep the y-axis tic marks parallel to the x-axis.

NLYINC

Default: NLYINC = 2

NLYINC controls the number of large tic marks to be displayed on the y-axis.

- -1000 drop the first and the last numbers on the y-axis and determine the number of large y-axis tic marks
- <0 drop the first and the last numbers on the $y\mbox{-axis}.$ The number of large tic marks on the $y\mbox{-axis}$ will be $|\mbox{\tt NLYINC}|+1,$ unless $\mbox{\tt YLOG}>1,$ in which case the number of tic marks will be determined to avoid fractional powers
- 0 the number of large tic marks on the y-axis will be automatically determined
- > 0 the number of large tic marks on the y-axis will be NLYINC+1, unless YLOG > 1, in which case NLYINC will be determined to avoid fractional powers

YTICL

Default: %YTICL = 2

YTICL is the length of the long tic marks on the y-axis. See Figure A.39 on page 367. %YTICL is a percentage of the height of the window, that is, $YTICL = \%YTICL \times (YUWIND - YLWIND) \div 100$

NSYINC

Default: NSYINC = 1

NSYINC controls the number of small tic marks to be displayed on the y-axis. The small tic marks appear between the large tic marks. See Figure A.39 on page 367.

- ≤ 1 no small tic marks on the *y*-axis
- ≥ 2 the number of small tic marks, on the y-axis, between each pair of large tic marks, will be NSYINC -1

If |YLOG| = 10, the small tic marks on the y-axis can be specified exactly with NSYINC. If small tic marks are desired at the locations $j_m \times 10^n$, where $2 \le j_m \le 9$, then set NSYINC $= -j_1 \dots j_m$. For example, if you want small tic marks at 2×10^n , 4×10^n , 5×10^n , and 8×10^n , then set NSYINC = -2458.

YTICS

Default: %YTICS = 1

YTICS is the length of the short tic marks on the y-axis. See Figure A.39 on page 367. %YTICS is a percentage of the height of the window, that is, $YTICS = \%YTICS \times (YUWIND - YLWIND) \div 100$

YMAX

Default: YMAX = 10

YMAX controls the maximum value for the y-axis.

If |YLOG| > 1, then YMAX is assumed to be the *exponent* for the maximum value on the *y*-axis. The maximum value displayed on the *y*-axis is $|YLOG|^{YMAX}$. Integer exponents are always used for the axis numbering, so the *y*-axis may not end on a large tic mark.

If $|YLOG| \le 1$, then YMAX is the actual maximum value for the y-axis. If YVMAX is equal to YMAX then the y-axis always ends on a large, labeled, tic mark. If YVMAX is not equal to YMAX then the y-axis will not end at a large, labeled, tic mark.

When the value of YMAX is changed, the value of YVMAX is simultaneously changed to the same value.

YVMAX

Default: YVMAX = 10

YVMAX controls the virtual maximum value for the *y*-axis. See Figure A.38 on page 363.

If |YLOG| > 1, the virtual maximum is ignored.

If $|\text{YLOG}| \leq 1$, then YVMAX is the virtual maximum value of the labels for the y-axis. This value will not be displayed if YVMAX is greater than YMAX, but it will be displayed if YVMAX is less than or equal to YMAX. This virtual maximum is used to determine "nice" numbers for labeling the large tic marks on the y-axis. If YVMAX is not equal to YMAX then the y-axis will not end on a large, labeled, tic mark.

The value of YVMAX is changed to the value of YMAX when the value of YMAX is changed. So, if you want to make YVMAX different from YMAX, it must be changed *after* YMAX is changed.

If NLYINC is set to zero, then the y-axis will begin at YMIN and end at YMAX, but if these are not "nice" numbers, the virtual minimum and maximum will be set to values outside the actual minimum and maximum so that the large tic marks can be labeled with "nice" numbers.

YMIN

Default: YMIN = 0

YMIN controls the minimum value for the *y*-axis.

If |YLOG| > 1, then YMIN is assumed to be the *exponent* for the minimum value on the *y*-axis. The minimum value displayed on the *y*-axis is $|YLOG|^{YMIN}$. Integer exponents are always be used for the axis numbering, so the *y*-axis may not begin on a large tic mark.

If $|YLOG| \le 1$, YMIN is the actual minimum value for the y-axis. If YVMIN is equal to YMIN then the y-axis always begins on a large, labeled, tic mark. If YVMIN is not equal to YMIN then the y-axis will not begin at a large, labeled, tic mark.

When the value of YMIN is changed, the value of YVMIN is simultaneously changed to the same value.

YVMIN

Default: YVMIN = 0

YVMIN controls the virtual minimum value for the y-axis. See Figure A.38 on page 363.

If |YLOG| > 1, the virtual minimum is ignored.

If $|YLOG| \le 1$, then YVMIN is the virtual minimum value of the labels for the y-axis. This value will not be displayed if YVMIN is less than YMIN, but it will be displayed if YVMIN is greater than or equal to YMIN. This virtual minimum is used to determine "nice" numbers for labeling the large tic marks on the y-axis. If YVMIN is not equal to YMIN then the y-axis will not begin on a large, labeled, tic mark.

The value of YVMIN is changed to the value of YMIN when the value of YMIN is changed. So, if you want to make YVMIN different from YMIN, it must be changed *after* YMIN is changed.

If NLYINC is set to zero, then the y-axis will begin at YMIN and end at YMAX, but if these are not

"nice" numbers, the virtual minimum and maximum will be set to values outside the actual minimum and maximum so that the large tic marks can be labeled with "nice" numbers.

YMOD

Default: YMOD = 0

YMOD is another control on the numbering for the y-axis.

- > 1 the numbers displayed on the y-axis will be modulo YMOD and positive. If the number to be displayed on the axis is x then the number that is actually displayed is $|x \text{YMOD} \times (x/\text{YMOD})| + \text{YOFF}$ -1 < YMOD < 1 the numbers will be displayed normally
- $-1 \le \mathtt{YMOD} \le 1$ the numbers will be displayed normally <-1 the numbers will be modulo $|\mathtt{YMOD}|$ and may be negative

YOFF

Default: YOFF = 0

YOFF is another control on the numbering for the y-axis. If |YMOD| > 1, then YOFF is added to the numbers to be displayed on the y-axis. If $|YMOD| \le 1$, then YOFF is ignored.

YLEADZ

Default: YLEADZ = 0

YLEADZ controls whether leading zeros are displayed on the y-axis numbering.

- $\neq 0$ the numbers displayed on the will have leading zeros if they are between 0 and 1
- 0 the numbers displayed on the will *not* have leading zeros if they are between 0 and 1

The default is to not display these leading zeros. Numbers that are between -1 and 0 always have the leading zero displayed.

YPAUTO

Default value: YPAUTO = 1

YPAUTO, along with YPOW, controls the y-axis scale factor that is appended to the y-axis text

label. This scale factor is needed when there are not enough digits allowed for the numbers labeling the y-axis.

- 2 determine YPOW, but do not append the scale factor to the text label even if it is needed
- 1 determine YPOW, and append the scale factor to the text label
- 0 use the present value of YPOW and if YPOW \neq 0 append the scale factor to the text label

If the user wishes to completely specify the appearance of the y-axis, YPAUTO must be set to 0, otherwise the number of digits and decimal places, NYDIG and NYDEC, may be changed.

YPOW

Default: YPOW = 0

YPOW controls the y-axis scale factor that is appended to the y-axis text label. This scale factor is a power of ten, that is, 10^{YPOW} , and the numbers labeling the y-axis should be multiplied by this scale factor to get the correct graph units.

NYDIG

Default: NYDIG = 5

NYDIG controls the total number of digits to be displayed in each of the numbers labeling the y-axis. If NYDIG is smaller than required to display the y-axis numbers, NYDIG will not be increased but a scale factor, $(\times 10^n)$, will be appended to the y-axis text label. If NYDIG is larger than required, NYDIG will be reduced to the minimum value required. The value of NYDIG is updated after each graph is drawn.

NYDEC

Default: NYDEC = -1

NYDEC controls the number of digits to be displayed in the fractional parts of the numbers labeling the y-axis. The value of NYDEC is updated after each graph is drawn.

GPLOT Keywords

- -1 display the numbers labeling the y-axis as integers, with no decimal point
- display the numbers labeling the y-axis with no fractional part, but with a decimal point
- >0 display the numbers labeling the y-axis with NYDEC digits in the fractional part

YNUMSZ

Default: %YNUMSZ = 3

YNUMSZ controls the size of the numbers labeling the y-axis. See Figure A.39 on page 367. %YNUMSZ is a percentage of the height of the window, that is, YNUMSZ = %YNUMSZ \times (YUWIND – YLWIND) \div 100

YNUMA

Default: YNUMA = -YAXISA

YNUMA controls the angle of the base line for the numbers labeling the y-axis. YNUMA is the angle, in degrees, measured counterclockwise, between a line parallel to the y-axis and the base line of a number. See Figure A.39 on page 367. By default, YNUMA is set as a percentage. If YNUMA is set as a percentage, for example, SET %YNUMA 0, then YNUMA is set to -YAXISA. The actual value of %YNUMA will be ignored. This allows the user to change the angle of the y-axis and keep the base line of the y-axis numbers horizontal.

YITICA

Default: YITICA = XAXISA - YAXISA + 180

YITICA, along with YITICL, controls the location of the numbers labeling the y-axis at the large tic marks. YITICA is the angle, in degrees, measured counterclockwise, between the y-axis and a line joining the base of each large tic mark on the y-axis to the center of the number labeling that tic mark. See Figure A.39 on page 367. By default, YITICA is set as a percentage. If YITICA is set as a percentage, then YITICA is set to XAXISA — YAXISA — YAXISA + 180. The actual value of %YITICA will be ignored. This allows the user to change the angle of the axes and keep the relative location of the y-axis numbers the same.

YITICL

Default: %YITICL = 3

YITICL, along with YITICA, controls the location of the numbers labeling the y-axis at the large tic marks. YITICL is the distance from the base of each large tic mark on the y-axis to the lower left hand corner of the number labeling that tic mark. See Figure A.39 on page 367. %YITICL is a percentage of the height of the window, that is, YITICL = %YITICL \times (YUWIND – YLWIND) \div 100

A.6 Axis Box Characteristics

Figure A.41 shows the relation of the world coordinate system to the window and to the graph axis corner locations.

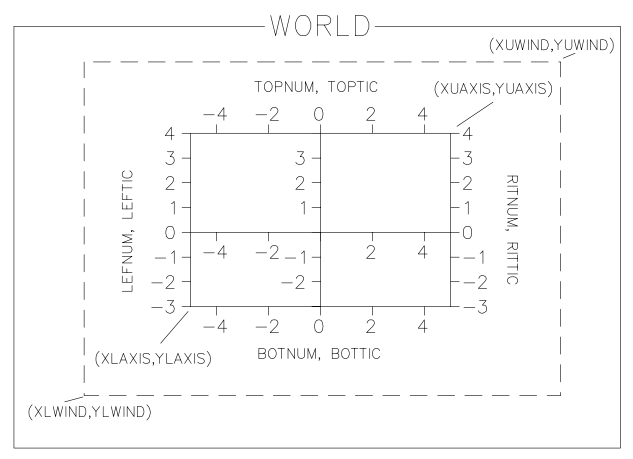


Figure A.41: The window and axis locations

XLWIND

Default: %XLWIND = 0

XLWIND is the left edge of the window box. See Figure A.41 on page 377. %XLWIND is a

GPLOT Keywords

percentage of the entire width of the world coordinate system.

XUWIND

Default: %XUWIND = 0

XUWIND is the right edge of the window box. See Figure A.41 on page 377. %XUWIND is a percentage of the entire width of the world coordinate system.

YLWIND

Default: %YLWIND = 0

YLWIND is the bottom edge of the window box. See Figure A.41 on page 377. %YLWIND is a percentage of the entire height of the world coordinate system.

YUWIND

Default: %YUWIND = 0

YUWIND is the top edge of the window box. See Figure A.41 on page 377. %YUWIND is a percentage of the entire height of the world coordinate system.

BOX

Default: BOX = 1

BOX controls whether or not an axis box is placed around the graph.

0 do not draw an axis box

 $\neq 0$ draw an axis box

XLAXIS

Default: %XLAXIS = 15

XLAXIS controls the position of the left, or lower, end of the x-axis, see Figure A.41 on page 377. This is also the horizontal coordinate of the lower left hand corner of the graph box, if BOX = 1. %XLAXIS is a percentage of the width of the window, that is, XLAXIS = XLWIND + %XLAXIS \times (XUWIND - XLWIND) \div 100

XUAXIS

Default: %XUAXIS = 95

XUAXIS controls the position of the right, or upper, end of the x-axis, see Figure A.41 on page 377. This is also the horizontal coordinate of the upper right hand corner of the graph box, if BOX = 1. %XUAXIS is a percentage of the width of the window, that is, XUAXIS = XLWIND + %XUAXIS \times (XUWIND - XLWIND) \div 100

XAXISA

Default: XAXISA = 0

XAXISA is the angle, in degrees, measured counterclockwise, between a horizontal line and the x-axis. See Figure A.36 on page 356.

YLAXIS

Default: %YLAXIS = 15

YLAXIS is the position of the bottom, or lower, end of the y-axis, see Figure A.41 on page 377. This is also the vertical coordinate of the lower left hand corner of the graph box, if BOX = 1. %YLAXIS is a percentage of the height of the window, that is, YLAXIS = %YLAXIS \times (YUWIND – YLWIND) \div 100

YUAXIS

Default: %YUAXIS = 90

YUAXIS is the position of the upper end of the y-axis, see Figure A.41 on page 377. This is also the vertical coordinate of the upper right hand corner of the graph box, if BOX = 1. %YUAXIS is a percentage of the height of the window, that is, YUAXIS = %YUAXIS \times (YUWIND - YLWIND) \div 100

YAXISA

Default: YAXISA = 90

YAXISA is the angle, in degrees, measured counterclockwise, between a horizontal line and the y-axis. See Figure A.39 on page 367.

BOTNUM

Default: BOTNUM = 0

GPLOT Keywords

BOTNUM controls the height of the numbers on the bottom edge of the box. |BOTNUM| is the ratio of the height of the numbers on the bottom edge of the box to XNUMSZ, the height of the numbers on the x-axis. The height of these numbers will be $|BOTNUM| \times XNUMSZ$.

BOTNUM is ignored if BOX = 0 or if the bottom edge of the axis box overlaps the x-axis.

- 0 no numbers on the bottom edge of the box
- > 0 numbers on bottom edge of the box on same side as x-axis numbers
- < 0 numbers on bottom edge of the box on opposite side as x-axis numbers

BOTTIC

Default: BOTTIC = 1

BOTTIC controls the lengths of the large and short tic marks on the bottom edge of the box. |BOTTIC| is the ratio of the lengths of the tic marks on the bottom edge of the box to the lengths of the tic marks on the x-axis, where XTICL is the length of the large x-axis tic marks and XTICS is the length of the short x-axis tic marks. The large tic marks on the bottom of the box will have a length of $|BOTTIC| \times XTICL$ and the short tic mark length will be $|BOTTIC| \times XTICS$.

BOTTIC is ignored if BOX = 0 or if the bottom edge of the axis box overlaps the x-axis.

- 0 no tic marks on the bottom edge of the box
- > 0 tic marks on bottom edge of the box on the same side as the x-axis tic marks
- < 0 tic marks on bottom edge of the box on the opposite side as the x-axis tic marks

RITNUM

Default: RITNUM = 0

RITNUM controls the height of the numbers on the right edge of the box. |RITNUM| is the ratio of the height of the numbers on the right edge of the box to YNUMSZ, the height of the numbers on the y-axis. This number height will be $|RITNUM| \times YNUMSZ$.

RITNUM is ignored if BOX = 0 or if the right edge of the axis box overlaps the y-axis.

- 0 no numbers on the right edge of the box
- > 0 numbers on right edge of the box on the same side as y-axis numbers
- < 0 numbers on right edge of the box on the opposite side as y-axis numbers

RITTIC

Default: RITTIC = -1

RITTIC controls the lengths of the large and short tic marks on the right edge of the box. |RITTIC| is the ratio of the lengths of the tic marks on the right edge of the box to the lengths of the tic marks on the y-axis. YTICL is the length of the large y-axis tic marks and YTICS is the length of the short y-axis tic marks. The large tic marks on the right edge of the box will have a length of $|RITTIC| \times YTICL$ and the short tic mark length will be $|RITTIC| \times YTICS$.

RITTIC is ignored if BOX = 0 or if the right edge of the axis box overlaps the y-axis.

- 0 no tic marks on the right edge of the box
- > 0 tic marks on right edge of the box on the same side as y-axis tic marks
- < 0 tic marks on right edge of the box on the opposite side as y-axis large tic marks

TOPNUM

Default: TOPNUM = 0

TOPNUM controls the height of the numbers on the top edge of the box. |TOPNUM| is the ratio of the height of the numbers on the top edge of the box to XNUMSZ, the height of the numbers on the x-axis. The height of these numbers will be $|TOPNUM| \times XNUMSZ$.

TOPNUM is ignored if BOX = 0 or if the top edge of the axis box overlaps the x-axis.

- 0 no numbers on the top edge of the box
- > 0 numbers on top edge of the box on the same side as x-axis numbers
- < 0 numbers on top edge of the box on the opposite side as x-axis numbers

TOPTIC

Default: TOPTIC = -1

GPLOT Keywords

TOPTIC controls the lengths of the large and short tic marks on the top edge of the box. |TOPTIC| is the ratio of the lengths of the tic marks on the top edge of the box to the lengths of the tic marks on the x-axis, where XTICL is the length of the large x-axis tic marks and XTICS is the length of the short x-axis tic marks. The large tic marks on the top edge of the box will have a length of $|TOPTIC| \times XTICL$ and the short tic mark length will be $|TOPTIC| \times XTICS$.

TOPTIC is ignored if BOX = 0 or if the top edge of the axis box overlaps the x-axis.

- 0 no tic marks on the top edge of the box
- >0 tic marks on top edge of the box on the same side as x-axis large tic marks
- < 0 tic marks on top edge of the box on the opposite side as x-axis large tic marks

LEFNUM

Default: LEFNUM = 0

LEFNUM controls the height of the numbers on the left edge of the box. |LEFNUM| is the ratio of the height of the numbers on the left edge of the box to YNUMSZ, the height of the numbers on the y-axis. The height of these numbers will be $|LEFNUM| \times YNUMSZ$.

LEFNUM is ignored if BOX = 0 or if the left edge of the axis box overlaps the y-axis.

- 0 no numbers on the left edge of the box
- > 0 numbers on left edge of the box on the same side as y-axis numbers
- < 0 numbers on left edge of the box on the opposite side as y-axis numbers

LEFTIC

Default: LEFTIC = 1

LEFTIC controls the lengths of the large and short tic marks on the bottom edge of the box. |LEFTIC| is the ratio of the lengths of the tic marks on the left edge of the box to the lengths of the tic marks on the y-axis, where YTICL is the length of the large y-axis tic marks and YTICS is the length of the short y-axis tic marks. The large tic marks on the left edge of the box will have a length of $|\text{LEFTIC}| \times \text{YTICL}$ and the short tic mark length will be $|\text{LEFTIC}| \times \text{YTICS}$.

LEFTIC is ignored if BOX = 0 or if the left edge of the axis box overlaps the y-axis.

GPLOT Keywords

- 0 no tic marks on the left edge of the box
- >0 tic marks on left edge of the box on the same side as y-axis tic marks
- < 0 tic marks on left edge of the box on the opposite side as y-axis tic marks

VAX/VMS COMMAND PROCEDURE

This command procedure must be executed *before* you invoke PHYSICA. This command procedure can be found in:

PHYSICA\$DIR: PHYSICA_USER_FUNCTIONS.COM

If the define command:

```
$ define PHYSICA_USER_FUNCTIONS your_source_file
```

is placed in your LOGIN.COM file, this command procedure need not be executed more than once. The shareable image created with this procedure will automatically be found when you run PHYSICA.

```
$SET VERIFY
$! This command procedure creates a shareable image for the 8 PHYSICA
$! user functions AND subroutines. Normally you need not do this again
$! since the shareable image should still be around from the last use.
$!
$! The FORTRAN sources should be installed in your own directory in a file,
$! named, for example: disk:[directory]TEST.FOR
$! If this is not present then the defaults installed in
$! PHYSICA$DIR:PHYSICA_USER_FUNCTIONS.FOR will be used.
$!
$! The following DEFINE command should have been set by your system manager
$! define PHYSICA_USER_FUNCTIONS PHYSICA$DIR:PHYSICA_USER_FUNCTIONS
$1
$! If PHYSICA_USER_FUNCTIONS is not defined, the image activator cannot find
$! the shareable image. To activate your own set of functions and subroutines,
$! the following DEFINE command should be in effect. If this is not defined
$! the image activator will then find the default shareable image.
$!
$ define PHYSICA_USER_FUNCTIONS disk:[directory]TEST
$ fortran PHYSICA_USER_FUNCTIONS
$! The following scripts are needed since transfer vectors may only be
$! created in VAX MACRO.
$macro/object=xfruser1 sys$input
      .title xfruser1 - Transfer vector for USER1
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
```

```
.transfer user1
      .mask user1
      jmp
               1^user1+2
      .end
$eod
$macro/object=xfruser2 sys$input
      .title xfruser2 - Transfer vector for USER2
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer user2
      .mask
               user2
      jmp
               1^user2+2
      .end
$eod
$macro/object=xfruser3 sys$input
      .title xfruser3 - Transfer vector for USER3
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer user3
      .mask
               user3
              1^user3+2
      jmp
      .end
$eod
$macro/object=xfruser4 sys$input
      .title xfruser4 - Transfer vector for USER4
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer user4
      .mask user4
      qmj
               1^user4+2
      .end
$eod
$macro/object=xfruser5 sys$input
      .title xfruser5 - Transfer vector for USER5
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer user5
      .{\tt mask}
               user5
               l^user5+2
      jmp
      .end
$eod
$macro/object=xfruser6 sys$input
      .title xfruser6 - Transfer vector for USER6
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer user6
```

```
.mask
               user6
      jmp
               1^user6+2
      .end
$eod
$macro/object=xfruser7 sys$input
      .title xfruser7 - Transfer vector for USER7
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer user7
      .mask
              user7
               1^user7+2
      jmp
      .end
$eod
$macro/object=xfruser8 sys$input
      .title xfruser8 - Transfer vector for USER8
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer user8
      .mask
              user8
               1^user8+2
      qmj
      .end
$macro/object=xfrsub1 sys$input
      .title xfrsub1 - Transfer vector for SUB1
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer sub1
      .mask
               sub1
      jmp
               1^sub1+2
      .end
$eod
$macro/object=xfrsub2 sys$input
      .title xfrsub2 - Transfer vector for SUB2
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer sub2
      .mask sub2
               1^sub2+2
      jmp
      .end
$eod
$macro/object=xfrsub3 sys$input
      .title xfrsub3 - Transfer vector for SUB3
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer sub3
      .{	t mask}
               sub3
```

```
1^sub3+2
     jmp
      .end
$eod
$macro/object=xfrsub4 sys$input
      .title xfrsub4 - Transfer vector for SUB4
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer sub4
      .{\tt mask}
               sub4
     jmp
               1^sub4+2
      .end
$eod
$macro/object=xfrsub5 sys$input
      .title xfrsub5 - Transfer vector for SUB5
     .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer sub5
      .mask
               1^sub5+2
     jmp
      .end
$eod
$macro/object=xfrsub6 sys$input
     .title xfrsub6 - Transfer vector for SUB6
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer sub6
      .mask sub6
               1^sub6+2
     qmj
      .end
$macro/object=xfrsub7 sys$input
      .title xfrsub7 - Transfer vector for SUB7
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer sub7
      .mask
               sub7
               1^sub7+2
     jmp
      .end
$eod
$macro/object=xfrsub8 sys$input
      .title xfrsub8 - Transfer vector for SUB8
      .ident /v01-001/
      .psect $$xfrvectors,exe,nowrt
      .transfer sub8
      .mask sub8
          l^sub8+2
      jmp
```

```
.end
$eod
$!
$! Link these 8 user functions and 8 subroutines as a shareable image,
         disk:[directory]test.exe
$! The names usern_vector and subn_vector are dummy but must be unique.
$link/share PHYSICA_USER_FUNCTIONS,sys$input/opt
gsmatch=lequal,1,0
cluster=user1_vector,,,xfruser1
cluster=user2_vector,,,xfruser2
cluster=user3_vector,,,xfruser3
cluster=user4_vector,,,xfruser4
cluster=user5_vector,,,xfruser5
cluster=user6_vector,,,xfruser6
cluster=user7_vector,,,xfruser7
cluster=user8_vector,,,xfruser8
cluster=sub1_vector,,,xfrsub1
cluster=sub2_vector,,,xfrsub2
cluster=sub3_vector,,,xfrsub3
cluster=sub4_vector,,,xfrsub4
cluster=sub5_vector,,,xfrsub5
cluster=sub6_vector,,,xfrsub6
cluster=sub7_vector,,,xfrsub7
cluster=sub8_vector,,,xfrsub8
$eod
$delete/NOconfirm xfruser*.obj;*,xfrsub*.obj;*
$set noverify
```

AlphaVMS COMMAND PROCEDURE

This command procedure must be executed *before* you invoke PHYSICA. This command procedure can be found in:

PHYSICA\$DIR: PHYSICA_USER_FUNCTIONS.COM

If the define command:

```
$ define PHYSICA_USER_FUNCTIONS your_source_file
```

is placed in your LOGIN.COM file, this command procedure need not be executed more than once. The shareable image created with this procedure will automatically be found when you run PHYSICA.

```
$SET VERIFY
$! This command procedure creates a shareable image for the 8 user functions
$! and 8 subroutines. The FORTRAN sources should be installed in your own
$! directory in a file, named, for example:
                                              disk:[directory]TEST.FOR
$! If this is not present then the defaults installed in
$! PHYSICA$DIR:PHYSICA_USER_FUNCTIONS.FOR will be used.
$! Your system manager should have created the default logical name:
$! DEFINE PHYSICA_USER_FUNCTIONS PHYSICA$DIR:PHYSICA_USER_FUNCTIONS
$! If PHYSICA_USER_FUNCTIONS is not defined, the image activator cannot find
$! the shareable image. To activate your own set of functions and subroutines,
$! the following DEFINE command should be in effect. If this is not defined
$! the image activator will find the default shareable image.
$! define PHYSICA_USER_FUNCTIONS disk:[directory]test
$! The logical name PHYSICA_USER_FUNCTIONS will need to be re-defined in each
$! new process where you want to make use of your own routines.
$ fortran PHYSICA_USER_FUNCTIONS.for
$!
$! Link these 8 user functions and 8 subroutines as a sharerable image,
        disk:[directory]test.exe
$! The names usern_vector and subn_vector are dummy but must be unique.
$link/shareable PHYSICA_USER_FUNCTIONS.obj, sys$input/opt
 GSMATCH=lequal,1,1000
 SYMBOL_VECTOR=(user1=PROCEDURE,-
                user2=PROCEDURE,-
```

user3=PROCEDURE, user4=PROCEDURE, user5=PROCEDURE, user6=PROCEDURE, user7=PROCEDURE, user8=PROCEDURE, sub1=PROCEDURE, sub2=PROCEDURE, sub3=PROCEDURE, sub4=PROCEDURE, sub5=PROCEDURE, sub5=PROCEDURE, sub6=PROCEDURE, sub7=PROCEDURE, sub7=PROCEDURE, sub8=PROCEDURE)

\$EOD

\$set noverify

USER ROUTINE SOURCE CODE

VMS: Following is the default source code for the eight functions and the eight subroutines that are linked to PHYSICA via the shareable image. This source code can be found in:

PHYSICA\$DIR: PHYSICA_USER_FUNCTIONS.FOR

UNIX: Following is the source code for the eight user defined functions and the eight user defined subroutines that are linked into PHYSICA by default. This source code can be found in:

phys_user.f

The process to change the user defined routines in physica follows:

- 1) edit the file phys_user.f to put in your versions of user1, ..., user8, sub1, ..., sub8
- 2) compile it, e.g., f77 -c phys_user.f
- 3) archive it with ar -rsv physica.a phys_user.o
- 4) link the program with physica.link

For off-site users: phys_user.f can be found in the physica-link tar file, along with the necessary archives.

```
REAL*8 FUNCTION USER1(A)
IMPLICIT REAL*8 (A-H,0-Z)
USER1=A
RETURN
END

REAL*8 FUNCTION USER2(A,B)
IMPLICIT REAL*8 (A-H,0-Z)
USER2=A+B
RETURN
END

REAL*8 FUNCTION USER3(A,B,C)
IMPLICIT REAL*8 (A-H,0-Z)
USER3=A+B+C
RETURN
END
```

CCC

```
REAL*8 FUNCTION USER4(A,B,C,D)
   IMPLICIT REAL*8 (A-H,0-Z)
   USER4=A+B+C+D
   RETURN
   END
   REAL*8 FUNCTION USER5(A)
   IMPLICIT REAL*8 (A-H,0-Z)
   USER5=5.0
   RETURN
   END
   REAL*8 FUNCTION USER6(A)
   IMPLICIT REAL*8 (A-H,0-Z)
   USER6=6.0
   RETURN
   END
   REAL*8 FUNCTION USER7(A,B)
   IMPLICIT REAL*8 (A-H,0-Z)
   SUM = 0.0
   DO I = 1, INT(A)
     SUM = SUM+B
   END DO
   USER7 = SUM
   RETURN
   END
   REAL*8 FUNCTION USER8(A)
   IMPLICIT REAL*8 (A-H,0-Z)
   USER8=8.0
   RETURN
   END
   SUBROUTINE SUB1(IATYPE, ICODE, IUPDATE, IER, X1, X2, ADIFF)
   INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
             ADIFF, X1(1), X2(1)
   REAL*8
Determine if vector X1 is different than vector X2
 If there is a difference set ADIFF = 1, otherwise set ADIFF = 0
   IF( IATYPE(1) .NE. 1 )GO TO 991
   IF( IATYPE(2) .NE. 1 )GO TO 992
   IF( IATYPE(3) .NE. 0 )GO TO 993
   IF( ICODE(1,1) .NE. ICODE(1,2) )GO TO 994
```

```
! third argument to be updated
      IUPDATE(3) = 1
      ADIFF = 0.0D0
     DO I = 1, ICODE(1,1)
        IF( X1(I) .NE. X2(I) )THEN
         ADIFF = 1.0D0
          RETURN
       END IF
     END DO
     RETURN
 991 WRITE(*,*)' error in SUB1: first argument is not a vector'
     IER = -1
     RETURN
 992 WRITE(*,*)' error in SUB1: second argument is not a vector'
      IER = -1
     RETURN
 993 WRITE(*,*)' error in SUB1: third argument is not a scalar'
     IER = -1
     RETURN
 994 WRITE(*,*)' error in SUB1: input vectors not the same length'
      IER = -1
     RETURN
C Setting IER to -1 indicates to PHYSICA that an error has occured in
C the subroutine. This is like an alternate RETURN. If you call this
C subroutine in a script command file, and the error occurs, the script
C will stop execution
     END
      SUBROUTINE SUB2(IATYPE, ICODE, IUPDATE, IER, XIN, XOUT, A)
      INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
     REAL*8
               A, XIN(1), XOUT(1)
  Divide a vector by a scalar and put the output in another vector
   e.g., CALL SUB2 XIN XOUT A and XOUT will be XIN/A
CCC
      IF( IATYPE(1) .NE. 1 )GO TO 991
      IF( IATYPE(2) .NE. 1 )GO TO 992
      IF( IATYPE(3) .NE. 0 )GO TO 993
                         ! second argument to be updated
      IUPDATE(2) = 1
     NPTS = MIN(ICODE(1,1),ICODE(1,2)) ! use minimum length
     DO I = 1, NPTS
        IF( XIN(I) .GT. 100.0D0 )G0 TO 994
       XOUT(I) = XIN(I)/A
     END DO
      RETURN
```

```
991 WRITE(*,*)' error in SUB2: first argument is not a vector'
     IER = -1
     RETURN
 992 WRITE(*,*)' error in SUB2: second argument is not a vector'
     IER = -1
     RETURN
 993 WRITE(*,*)' error in SUB2: third argument is not a scalar'
     IER = -1
     RETURN
 994 WRITE(*,*)' error in SUB2: XIN(I) > 100'
     IER = -1
     RETURN
     END
     SUBROUTINE SUB3(IATYPE, ICODE, IUPDATE, IER, MATRIX, DIAGONAL)
      INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
     REAL*8 MATRIX(1), DIAGONAL(1)
 Extract the diagonal from a square matrix and put it into a vector
  e.g., CALL SUB3 MATRIX DIAGONAL
С
        and DIAGONAL will be MATRIX[J,J] for J = 1, #
CCC
     IF( IATYPE(1) .NE. 2 )GO TO 991
      IF( IATYPE(2) .NE. 1 )GO TO 992
     IUPDATE(2) = 1
                       ! second argument to be updated
     NROWS = ICODE(1,1)
     NCOLS = ICODE(2,1)
     IF( NCOLS .NE. NROWS )GO TO 993
     DO J = 1, NCOLS
       DIAGONAL(J) = MATRIX(J+(J-1)*NCOLS)
     END DO
 991 WRITE(*,*)' error in SUB3: first argument is not a matrix'
     IER = -1
     RETURN
 992 WRITE(*,*)' error in SUB3: second argument is not a vector'
     IER = -1
 993 WRITE(*,*)' Use a square matrix for SUB3'
     IER = -1
     RETURN
     END
     SUBROUTINE SUB4(IATYPE, ICODE, IUPDATE, IER, X, M, XOUT)
      INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
             M(1), X(1), XOUT(1)
     REAL*8
```

```
Calculate XOUT = X<>M (inner product)
С
   where X is a vector, M is a matrix, XOUT is the output vector
CCC
     IF( IATYPE(1) .NE. 1 )GO TO 991
      IF( IATYPE(2) .NE. 2 )GO TO 992
     IF( IATYPE(3) .NE. 1 )GO TO 993
      IUPDATE(3) = 1
                        ! third argument to be updated
     NROWS = ICODE(1,2)
     NCOLS = ICODE(2,2)
     NUMX = ICODE(1,1)
     NUMXO = ICODE(1,3)
     IF( NUMX .NE. NROWS )GO TO 994
     IF( NUMXO .NE. NCOLS )GO TO 995
     DO I = 1, NCOLS
       XOUT(I) = 0.0D0
       DO J = 1, NROWS
          XOUT(I) = XOUT(I) + X(J)*M(I+(J-1)*NCOLS)
       END DO
     END DO
     RETURN
 991 WRITE(*,*)' *** error in SUB4'
     WRITE(*,*)' first argument is not a vector'
     IER = -1
     RETURN
 992 WRITE(*,*)' *** error in SUB4'
     WRITE(*,*)'
                     second argument is not a matrix'
     IER = -1
     RETURN
 993 WRITE(*,*)' *** error in SUB4'
     WRITE(*,*)' third argument is not a vector'
     IER = -1
     RETURN
 994 WRITE(*,*)' *** error in SUB4'
                    vector length not equal to row dimension of matrix'
     WRITE(*,*)'
     IER = -1
     RETURN
 995 WRITE(*,*)' *** error in SUB4'
     WRITE(*,*)
            output vector length not equal to column dimension of matrix'
     IER = -1
     RETURN
     END
     SUBROUTINE SUB5(IATYPE, ICODE, IUPDATE, IER, LFILE, X, Y)
      INTEGER*4
                  IATYPE(15),ICODE(3,15),IUPDATE(15),IER
```

```
LENF, IUNIT, I, NPTS
      INTEGER*4
     REAL*8
             X(1), Y(1), XDUM
     LOGICAL*4 LUOPEN
     LOGICAL*1 LFILE(1)
     CHARACTER*80 AFILE
 Read a vector X from a file, multiply it by 5 and add it to vector Y
 with the result put into X
C e.g., CALL SUB5 'FILE.DAT' X Y
         and X will be X*5+Y
CCC
     IF( IATYPE(1) .NE.-1 )GO TO 991
     IF( IATYPE(2) .NE. 1 )GO TO 992
      IF( IATYPE(3) .NE. 1 )GO TO 993
     IF( ICODE(1,1) .GT. 80 )GO TO 994
     IUPDATE(2) = 1
                        ! second argument to be updated
     LENF = ICODE(1,1)
     DO I = 1, LENF
       AFILE(I:I) = CHAR(LFILE(I))
     END DO
     D0\ 2\ I = 30,\ 99
       INQUIRE(UNIT=I,OPENED=LUOPEN,ERR=2)
       IF( .NOT.LUOPEN )THEN
         IUNIT = I
         GO TO 4
       END IF
   2 CONTINUE
   4 OPEN(FILE=AFILE(1:LENF), UNIT=IUNIT, STATUS='OLD', READONLY, SHARED, ERR=995)
     I = 1
  10 READ(33,*,ERR=996,END=20)XDUM
     IF( I .GT. ICODE(1,2) )THEN
       WRITE(*,*)' message from SUB5'
       WRITE(*,*)' max. number of elements from the file has been read'
       WRITE(*,*)' but there is more that could be read'
       GO TO 20
     END IF
     X(I) = XDUM
     I = I+1
     GO TO 10
 20 NPTS = MIN(ICODE(1,2),ICODE(1,3))
     DO I = 1, NPTS
       X(I) = X(I)*5.0D0+Y(I)
     END DO
      ICODE(1,2) = NPTS ! update the length of X
     RETURN
 991 WRITE(*,*)' *** error in SUB5'
```

```
WRITE(*,*)'
                     first argument is not a string'
      IER = -1
     RETURN
 992 WRITE(*,*)' *** error in SUB5'
     WRITE(*,*)'
                     second argument is not a vector'
     IER = -1
     RETURN
 993 WRITE(*,*)' *** error in SUB5'
     WRITE(*,*)'
                    third argument is not a vector'
     IER = -1
     RETURN
 994 WRITE(*,*)' *** error in SUB5'
     WRITE(*,*)'
                     string is longer than 80 characters'
     IER = -1
     RETURN
 995 WRITE(*,*)' *** error in SUB5'
     WRITE(*,*)'
                    unable to open file: '//AFILE(1:LENF)
     IER = -1
     RETURN
 996 WRITE(*,*)' *** error in SUB5'
     WRITE(*,9961)I,AFILE(1:LENF)
9961 FORMAT('
                  reading line#', I3,' from file: ',A)
     IER = -1
     RETURN
     END
     SUBROUTINE SUB6(IATYPE, ICODE, IUPDATE, IER, X, Y)
      INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
     REAL*8 X(1), Y(1)
  Multiply a vector by 6 and add it to another vector
  e.g., CALL SUB6 X Y
         and X will be X+6*Y
C
CCC
      IF( IATYPE(1) .NE. 1 )GO TO 991
      IF( IATYPE(2) .NE. 1 )GO TO 992
                         ! second argument to be updated
      IUPDATE(2) = 1
     NPTS = MIN(ICODE(1,1),ICODE(1,2))
     DO I = 1, NPTS
       X(I) = X(I) + Y(I) *6.000
     END DO
     RETURN
 991 WRITE(*,*)' *** error in SUB6'
     WRITE(*,*),
                     first argument is not a vector'
     IER = -1
     RETURN
```

```
992 WRITE(*,*)' *** error in SUB6'
     WRITE(*,*)' second argument is not a vector'
     IER = -1
     RETURN
     F.ND
     SUBROUTINE SUB7(IATYPE, ICODE, IUPDATE, IER, X, Y)
      INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
     REAL*8 X(1), Y(1)
 Multiply a vector by 7 and add it to another vector
 e.g., CALL SUB7 X Y
С
         and X will be X+7*Y
CCC
     IF( IATYPE(1) .NE. 1 )GO TO 991
     IF( IATYPE(2) .NE. 1 )GO TO 992
     IUPDATE(2) = 1 ! second argument to be updated
     NPTS = MIN(ICODE(1,1),ICODE(1,2))
     DO I = 1, NPTS
       X(I) = X(I) + Y(I) * 7.000
     END DO
     RETURN
 991 WRITE(*,*)' *** error in SUB7'
     WRITE(*,*)' first argument is not a vector'
     IER = -1
     RETURN
 992 WRITE(*,*)' *** error in SUB7'
     \mbox{WRITE(*,*)'} second argument is not a vector'
     IER = -1
     RETURN
     END
     SUBROUTINE SUB8(IATYPE, ICODE, IUPDATE, IER, X, Y)
      INTEGER*4 IATYPE(15),ICODE(3,15),IUPDATE(15),IER
     REAL*8 X(1), Y(1)
C Multiply a vector by 8 and add it to another vector
 e.g., CALL SUB8 X Y
C
         and X will be X+8*Y
CCC
     IF( IATYPE(1) .NE. 1 )GO TO 991
      IF( IATYPE(2) .NE. 1 )GO TO 992
     IUPDATE(2) = 1 ! second argument to be updated
     NPTS = MIN(ICODE(1,1),ICODE(1,2))
     DO I = 1, NPTS
       X(I) = X(I) + Y(I) * 8.000
```

<u>User Routine Source Code</u>

```
END DO
RETURN

991 WRITE(*,*)' *** error in SUB8'
WRITE(*,*)' first argument is not a vector'
IER = -1
RETURN

992 WRITE(*,*)' *** error in SUB8'
WRITE(*,*)' second argument is not a vector'
IER = -1
RETURN
END
```

| Hcor | Pouting | e Source | Code |
|------|---------|----------|------|
| O2CI | NOULIII | 5 JUUICE | COUC |

Index

| ! character, 78, 187 | virtual minimum, 362 |
|------------------------------------|---|
| Γ function, 301 | y-axis, 359, 366 |
| incomplete, 302 | angle, 379 |
| natural logarithm, 301 | digits, 375 |
| *M keyword, 45 | digits in fraction, 375 |
| *S keyword, 45 | menu, 64 |
| *T keyword, 45 | numbers, 368, 370, 372-377 |
| *V keyword, 45 | angle, 376 |
| β functions, 294 | drop first/last, 371 |
| complete, 301, 305 | size, 376 |
| β functions complete, 294 | plot characteristics, 366 |
| β functions incomplete, 295 | position, 379 |
| χ^2 minimization, 97 | scale factor, 374, 375 |
| χ^2 of fit, 97, 99 | small tics, 368, 372 |
| χ^2 probability function, 297 | text label, 368, 375 |
| μ SR MUD data sets, 221 | tic marks, 368, 370-372, 376 |
| μ SR data sets, 223 | virtual maximum, 372 |
| ψ function, 295, 296, 298 | virtual minimum, 373 |
| x-axis, 357, 370 | .physicarc file, 29 |
| angle, 379 | // operator, 284, 314 |
| digits, 365 | ? for script parameters, 78 |
| digits in fraction, 365 | @ for EXECUTE, 76 |
| menu, 64 | \$HOME/.physicarc file, 29 |
| numbers, 357, 359, 361-364, 366 | \ NORMAL qualifier, 94 |
| angle, 366 | \ALLWINDOWS qualifier, 209, 210, 212 |
| drop first/last, 360 | \ALPHANUMERIC qualifier, 19 |
| size, 365 | \APPEND qualifier, 28, 135, 187, 190, 248, |
| plot characteristics, 355 | 276 |
| position, 378, 379 | \AREAS qualifier, 23, 25, 40 |
| scale factor, 364, 365 | \ASCII qualifier, 186, 194, 196, 204 |
| small tics, 357, 361 | \AVERAGE qualifier, 9 |
| text label, 357, 364, 365 | \AXES qualifier, 23, 24, 31, 116, 208 |
| tic marks, 359–361, 366 | \AXESONLY qualifier, 116, 118 |
| virtual maximum, 361 | \BORDER qualifier, 27, 32 |

<u>INDEX</u>

| \BOUNDS qualifier, 125 | \EXPAND qualifier, 45 |
|--|--|
| BOXES qualifier, 40, 110, 233 | EXTEND qualifier, 187, 190 |
| \CHAOS qualifier, 224 | FIOWA qualifier, 115, 154, 212 |
| CHECKDUP qualifier, 124 | FIOWABIG qualifier, 115, 154 |
| CHISQ qualifier, 97, 99 | FIXED qualifier, 244 |
| \CIT467 qualifier, 19 | FLIP qualifier, 50 |
| \CL qualifier, 100 | \FLIPPED qualifier, 196 |
| \CLEAR qualifier, 183 | FORMAT qualifier, 146, 186, 194, 196, |
| \CLOSE qualifier, 185 | 276 |
| COLOUR qualifier, 34, 35, 50, 255 | \FRACTION qualifier, 71, 72 |
| \COLOURS qualifier, 22 | \FREE qualifier, 98 |
| COMMENSURATE qualifier, 226 | \GRAPH qualifier, 83, 139, 143, 163, 209, |
| CONFIRM qualifier, 56, 74, 84, 257, 262 | 261 |
| CONTINUE qualifier, 23, 185 | \GREY qualifier, 50 |
| CONTOURS qualifier, 39 | \HBOOK qualifier, 153, 214 |
| COORDINATES qualifier, 23 | \HIST qualifier, 218 |
| CORRMAT qualifier, 99 | \HISTOGRAM qualifier, 34, 117, 240 |
| COUNTER qualifier, 146 | \HORIZONTAL qualifier, 279 |
| COUNTS qualifier, 162 | \IMSR qualifier, 223 |
| COVMAT qualifier, 99 | \INDEX qualifier, 28 |
| CYCLES qualifier, 7 | \INDICES qualifier, 126, 268 |
| DERIV qualifier, 32 | \INITIALIZE qualifier, 29 |
| DIFFUSION qualifier, 36, 214 | \INSIDE qualifier, 182 |
| DIR qualifier, 215 | INTERPOLATE qualifier, 124 |
| \DISCARD qualifier, 7, 10 | INTERPSIZE qualifier, 26 |
| \DISPLAY qualifier, 159 | \ITMAX qualifier, 100 |
| \DITHER qualifier, 38 | \KEYPAD qualifier, 12 |
| \DOTPLOT qualifier, 219 | \LAGRANGE qualifier, 8 |
| DOWN qualifier, 244, 245, 283, 284 | \LEGEND qualifier, 22, 23, 34, 39 |
| DUMMY qualifier, 218, 225, 244, 342- | \LEVELS qualifier, 39 |
| 345 | \LIBRARY qualifier, 130 |
| \E1 qualifier, 99 | \LINES qualifier, 39 |
| \E2 qualifier, 99 | \LIST qualifier, 215 |
| \EDGES qualifier, 9 | \LOG qualifier, 31 |
| \EMPTY qualifier, 9, 10 | \MACRO qualifier, 57, 75, 135 |
| \EQUALLY_SPACED qualifier, 40 | \MATRICES qualifier, 243 |
| \ERASE qualifier, 262 | \MATRIX qualifier, 11, 146, 161, 162, 195, |
| ERRFILL qualifier, 110, 187, 194, 233 | 216, 271, 278 |
| \ERRSKIP qualifier, 187 | \MAX qualifier, 161 |
| ERRSTOP qualifier, 187 | \MEAN qualifier, 87 |
| EXECUTE qualifier, 248 | \MEDIAN qualifier, 87 |

| \MESSAGES qualifier, 71, 72, 101, 104, | \UPDATE qualifier, 101 |
|---|---|
| 185, 248, 268, 272 | \VARNAMES qualifier, 94 |
| \MIN qualifier, 161 | VARY qualifier, 94, 225, 244 |
| \MOMENT qualifier, 251 | VECTORS qualifier, 186, 243 |
| \MSR qualifier, 223 | VERTICAL qualifier, 279 |
| \MUD qualifier, 221 | VIRTUAL qualifier, 109, 232 |
| \NBINS qualifier, 8 | VOLUMES qualifier, 25, 40 |
| \NONRECURSIVE qualifier, 88 | \WEIGHTS qualifier, 6, 7, 10, 97, 250 |
| \NOREPLOT qualifier, 19 | WORLD qualifier, 83, 139, 143, 163 |
| \NPTS qualifier, 71, 159 | WRITE qualifier, 12 |
| \OVERLAY qualifier, 187, 190 | XAXIS qualifier, 136, 357 |
| \PAGE qualifier, 130, 146 | \XDISCARD qualifier, 10 |
| \PARAMETERS qualifier, 71, 73 | XFIOWA qualifier, 214 |
| \PARTIAL qualifier, 22, 32 | XPROFILE qualifier, 32 |
| \PEARSON qualifier, 252 | XYOUT qualifier, 70, 71, 125-127 |
| PERCENT qualifier, 83, 139, 143, 163 | \YAXIS qualifier, 136, 368 |
| \PHYSICA qualifier, 11, 211 | YBOS qualifier, 154, 218 |
| \PNUM qualifier, 157 | \YDISCARD qualifier, 10 |
| \POINTS qualifier, 37 | YPROFILE qualifier, 32, 34, 39 |
| POLAR qualifier, 24, 33, 119, 124, 271 | \ZEROS qualifier, 98 |
| \POLYGON qualifier, 161, 183 | 3-d figures, 254 |
| PROFILE qualifier, 32, 34, 39, 214 | 3DPLOT command, 4 |
| RANDOM qualifier, 34, 106, 112, 237, 291 | hardcopy, 5 |
| \READ qualifier, 12 | . 404 |
| \RECURSIVE qualifier, 90 | acute, 181 |
| REPLOT qualifier, 57, 70, 71, 75, 117, | adjacent duplicate points, 268 |
| 209 | AERF function, 300 |
| REPLOTONLY qualifier, 19, 210 | AERFC function, 300 |
| \RESET qualifier, 23, 24, 32, 34, 39, 94 | AIRY function, 294 |
| \RWN qualifier, 215 | AIX, 211 |
| \SCALAR qualifier, 20, 193, 243, 277 | ALIAS command, 5 , 29, 225 |
| \SIZE qualifier, 125 | ALL keyword, 29 |
| \SPECIFIC qualifier, 21 | Alpha, 3 |
| \STATIC qualifier, 12 | alphanumeric clear, 19 |
| \TEXT qualifier, 203, 209, 243, 278 | alphanumeric monitor width, 112, 238 |
| \TILE qualifier, 274 | alphanumeric terminal window, 5 |
| \TOLERANCE qualifier, 96 | alphanumeric window, 4 |
| \TTBUFFERS qualifier, 11, 212 | AlphaVMS, 14 |
| \UNFORMATTED qualifier, 189, 195, 201, | altering vector content, 44 |
| 204 | AMP&PHASE keyword, 324 |
| \UP qualifier, 245, 283, 284 | angular momentum, 306 |

<u>INDEX</u>

| append operator, 284, 314 | AUTOHEIGHT keyword, 138 |
|--|--------------------------------|
| example, 284 | automatic plotting, 208 |
| append scalar, 314 | AUTOSCALE keyword, 109, 231 |
| append string, 284 | autoscaling, 225 |
| append to string variable, 314 | autoscaling axes, 29, 116, 231 |
| append vectors, 284 | average deviation, 249, 251 |
| arc drawing, 83, 84 | average filter, 87 |
| ARC keyword, 84 | axis autoscaling, 231 |
| arc length, 332, 334 | axis box, 357, 368 |
| arc tangent, 290 | axis box characteristics, 377 |
| area calculation, 316 | axis commensurate, 226 |
| area fill, 111, 234 | axis corner locations, 377 |
| AREA function, 316 | axis drawing, 116 |
| area inside polygon, 316 | axis length, 109, 232 |
| arithmetic mean, 249 | axis scales, 115, 226 |
| array function | axis text labels, 136 |
| LOOP, 345 | AXP OSF/1, 153 |
| PROD, 343 | 1 000 |
| RPROD, 345 | bar graphs, 263 |
| RSUM, 344 | BEI function, 303 |
| SUM, 342 | BELL command, 271 |
| array string variable, 108, 204 | BER function, 303 |
| length, 266 | BESI0 function, 296 |
| ARROLEN keyword, 86, 108, 230 | BESI1 function, 296 |
| ARROTYP keyword, 86, 108, 231 | BESJ0 function, 295 |
| arrow drawing, 83, 85, 230 | BESJ1 function, 295 |
| arrow head length, 86, 108 | BESK0 function, 296 |
| arrow head width, 86, 108 | BESK1 function, 296 |
| ARROW keyword, 85, 110, 233 | Bessel functions, 295, 303 |
| arrow styles, 86 | first kind, 295 |
| arrow type, 108 | modified, 296, 303 |
| ARROWID keyword, 86, 108, 231 | second kind, 295 |
| ASCII code, 239 | BESTFIT command, 6 |
| ASCII decimal representation, 311, 315 | cycles, 7 |
| ASSIGN command, 5 | iterations, 7 |
| associated Legendre functions, 304 | weights, 6 |
| asymmetric error bars, 119 | BESY0 function, 295 |
| asynchronous trapping, 16, 78, 149 | BESY1 function, 295 |
| ATAN2 function, 290 | BETA function, 294 |
| ATAN2D function, 290 | beta functions, 294 |
| attaching to a subprocess, 28 | complete, 294 |

| incomplete, 295 | , 379–382 |
|---|---|
| BETAIN function, 295 | box number size, 380-382 |
| BIN command, 7 | breve, 181 |
| averages, 9 | BROADCAST keyword, 55, 73 |
| counts, 7 | BUFFER command, 11, 58, 76 |
| data, 7 | |
| discard extremes, 7 | CALL command, 14 , 147, 152 |
| edge defined bins, 9 | CAREA vector, 25 |
| increment only if empty, 9 | causal recursive filter, 87 |
| Lagrange type, 8 | CCONT vector, 23 |
| number of bins, 8 | centimeters, 30, 83, 111, 112, 140, 163, |
| weight, 7 | 234, 236, 261 |
| BIN2D command, 9, 161 | central measures, 248 |
| bins defined by box corners, 11 | CERN library, 153, 215 |
| dimensions, 10 | CHAOS data sets, 224 |
| discard extremes, 10 | CHAR function, 311 , 315 |
| extremes, 10 | CHARA keyword, 354 |
| increment only if empty, 10 | CHARSZ keyword, 4, 354 |
| weights, 10 | CHEBY function, 296 |
| binary file read, 189 | Chebyshev polynomials, 296, 303 |
| BINARY keyword, 276 | chi-square minimization, 97 |
| binary ordered Walsh function, 308 | chi-square probability function, 297 |
| BINOM function, 296 | inverse, 297 |
| binomial coefficient, 296 | CHISQ function, 297 |
| BIRY function, 294 | CHISQINV function, 297 |
| bitmap | CHLOGU function, 302 |
| compressed format, 128 | choose window, 56, 74 |
| device, 6 | choosing data points, 159 |
| erase, 262 | circle drawing, 83, 84 |
| hardcopy, 140, 352 | CIRCLE keyword, 84, 110, 233 |
| bivariate normal probability function, 297 | circumflex, 181 |
| BIVARNOR function, 297 | CIT467 |
| blurring vector, 328 | toggle graphics, 19 |
| bolding, 172 | CIT467 keyword, 155 |
| Boolean, 80 | clear alphanumerics, 19 |
| Boolean operator, 281 | CLEAR command, 19 , 113, 209, 210, |
| BORDER keyword, 55, 73 | 238, 274 |
| BOTNUM keyword, 379 | clear graphics, 54, 155, 156, 183, 209 |
| BOTTIC keyword, 380 | Clebsch-Gordan coefficients, 306 |
| box drawing, 83, 85 | Clebsch-Gordan coefficients function, 307 |
| BOX keyword, 85, 110, 233, 357, 368, 378 | CLEBSG function, 307 |

INDEX

| CLEN function, 314 | size, 23 |
|---|------------------------------------|
| CLIP keyword, 353 | contour level |
| clipping, 139 | colour, 22 |
| close drawing file, 69 | exact, 21 |
| close file, 185 | label, 22 |
| CLOSE keyword, 69 | separation, 22 |
| closest value, 337 | size, 22 |
| CNTSEP keyword, 109, 232 | saving, 23 |
| colour, 209, 225, 240 | selection, 21 |
| colour bitmap, 48 | zooming in, 22 |
| COLOUR command, 20, 174, 261 | specific, 21 |
| using a scalar, 20 | contour matrix |
| COLOUR keyword, 352 | boundary, 27 |
| commensurate and windows, 272 | contour matrix data, 24 |
| commensurate axes, 109, 226, 232 | contour plot |
| commensurate graph and windows, 226 | legend entry size, 110, 232 |
| comment, 78 | legend format, 110, 233 |
| in formatted text, 166 | contour polar coordinates, 24 |
| comment line, 263 | contour scattered points, 24 |
| common scale, 117 | contour volume, 23, 25 |
| complementary error function, 300 | calculation, 25 |
| inverse, 300 | control keys, 11 |
| complete β function, 294, 301, 305 | control-c trapping, 78, 94 |
| compressed format, 128 | control-I, 136 |
| conditional statements, 80 | control-z, 28 |
| confidence level of fit, 100 | conventions used in this manual, 3 |
| CONFIRM keyword, 56, 74, 84 | CONVOL function, 328 |
| confirmation request, 56, 74 | convolution, 328 |
| contour, 212 | even number of points, 329 |
| contour area, 23, 25 | formula, 335 |
| calculation, 25 | noise effects, 329 |
| interpolation size, 26 | odd number of points, 328 |
| contour axes, 24 | convolution formula, 86 |
| CONTOUR command, 21 , 110, 136, 231, | convolution integral, 309 |
| 233 | coordinate conversion, 275 |
| contour coordinates, 23, 25 | COPY command, 27 |
| contour label, 23 | append, 28 |
| separation, 109, 232 | conditional, 27 |
| size, 110, 232 | unconditional, 28 |
| contour legend, 23 | correlation matrix, 99 |
| axis relocation, 23 | COS&SIN keyword, 325 |

| cosine integral, 298 | filled boxes, 41 |
|--|-------------------------------|
| COSINT function, 298 | input variables, 40 |
| covariance matrix, 99 | diffusion, 36 |
| crosshair, 157, 354 | grey scales, 36 |
| cubic spline, 112, 237, 317, 331-334 | dithering pattern, 37, 38 |
| CUNITS keyword, 83, 113, 143, 157, 159, | areas, 40 |
| 242 | contours, 39 |
| CURSOR keyword, 258, 354 , 355 | data range, 38 |
| CVOLM vector, 25 | input variables, 38 |
| CXMAX vector, 25 | legend, 39 |
| CXMIN vector, 25 | user defined, 38 |
| CYMAX vector, 25 | volumes, 40 |
| CYMIN vector, 25 | grey scales, 37 |
| | legend, 32 |
| data 'spikes', 87 | legend entry size, 110, 232 |
| data interpolation, 330 | legend format, 110, 233 |
| DATE function, 262, 310 | log of data, 31 |
| example, 310 | matrix data boundary, 32 |
| DAWSON function, 298 | of derivative, 32 |
| Dawson's integral, 298 | polar coordinates, 33 |
| DCL command, 28 | profiles, 32 |
| DEALIAS command, 5, 28 | random points, 34 |
| deconvolution, 328 | input variables, 35, 36 |
| even number of points, 329 | random points colour, 35 |
| DEFAULTS command, 29 | solid filled regions, 33 |
| initialization file, 29 | input variables, 33 |
| reset windows, 29 | legend, 34 |
| degrees of freedom, 98 | types, 31 |
| DENS\$AREA vector, 40 | zooming in, 32 |
| DENS\$CONT vector, 39 | DERIV function, 112, 237, 317 |
| DENS\$VOLM vector, 40 | example, 318 |
| DENSITY command, 30 , 50, 110, 112, | Fritsch and Carlson, 318 |
| 136, 214, 231, 233, 237 | interpolating splines, 317 |
| \DIFFUSION qualifier, 218 | Lagrange polynomials, 317 |
| \PROFILES qualifier, 218 | smoothing splines, 317 |
| density plot, 31, 212 | DESTROY command, 44, 183 |
| axes, 31 | all matrices, 45 |
| boxes, 40 | all scalars, 45 |
| accentuating values, 41 | all string variables, 45 |
| box size scale factor, 41 | all vectors, 45 |
| delimiting values, 41 | conditional, 45 |

<u>INDEX</u>

| examples, 46 | PROMPTING keyword, 57, 79 |
|--|---|
| expand names, 45 | REPLAY keyword, 57 |
| expression, 45 | REPLOT keyword, 57 |
| index ranges, 45 | SHELL keyword, 58 |
| keywords, 45 | STACK keyword, 58, 247 |
| unconditional, 44 | discrete Fourier series, 325 |
| DET function, 322 | dispersion, 248 |
| determinant of a matrix, 322 | DISPLAY command, 4, 30, 58, 111, 112, |
| DEVICE command, 47, 69, 84, 114, 128, | 119, 166, 234, 236, 239, 241, 242, |
| 140, 157, 163, 212, 225 | 261, 271 |
| keywords, 47 | FILL keyword, 120, 164, 173, 264 |
| DEVICE keyword, 114 | FONT keyword, 59, 113, 180 |
| dieresis, 181 | HATCH keyword, 59, 110, 234 |
| differentiating nonrecursive filters, 88 | LINES keyword, 59, 82 |
| diffraction theory, 301 | MENU keyword, 62, 108 |
| DIGAMMA function, 295, 296, 298 | message, 58 |
| Digi-Pad, 54 | PCHAR keyword, 62 |
| digital filter, 86 | SPECIAL keyword, 59 |
| digital halftoning, 36 | display file, 53 |
| digital smoothing polynomial filter, 334 | plotting units, 53 |
| Digital Unix, 211 | display variable, 243 |
| DIGITIZE command, 53 | dithering pattern, 41, 83, 110, 233 |
| keyboard key codes, 55 | DO loop, 57, 74, 77, 80, 269 |
| mouse button codes, 55 | nested, 80 |
| optional output variables, 54 | problems, 114, 237 |
| preparation, 54 | dot fill pattern, 112, 120, 121, 164, 172, |
| digitizing crosshair, 54 | 173, 237, 264, 353 |
| digitizing data, 53 | PostScript, 50 |
| digitizing pad types, 54 | dot fill pattern definition, 164, 173 |
| digitizing points, 160 | dotplot, 219 |
| DILOG function, 299 | dots per inch, 33, 35, 36, 38, 48, 49, 121, |
| Dilogarithm function, 299 | 164, 173, 241, 264 |
| DIM function, 292 | draw arc, 84 |
| DISABLE command, 55 , 73, 209 | draw arrow, 85, 108, 230 |
| BORDER keyword, 55, 272 | draw box, 85 |
| BROADCAST keyword, 55 | draw circle, 84 |
| CONFIRM keyword, 56, 84, 257 | draw circle sector, 84 |
| ECHO keyword, 56 | draw ellipse, 85 |
| local effect, 56 | draw figures, 56, 74, 82 |
| HISTORY keyword, 56 | draw polygon, 85 |
| JOURNAL keyword, 57, 75, 135 | draw rectangle, 85 |

| draw string, 56, 74, 263 | ELLIPSE command, 70 , 111, 208, 235 |
|--|--|
| draw string characteristics, 257 | explicitly defined, 71 |
| draw wedge, 84 | fit an ellipse, 71 |
| drawing file close, 69 | fraction, 72 |
| drawing file edit, 69 | messages, 72 |
| drawing file frame, 70 | method, 72 |
| drawing file open, 69 | number of points, 71 |
| dummy variable, 225, 342 | output variables, 70, 73 |
| duplicate points eliminate, 268 | parameter order, 71, 72 |
| dynamic buffer, 11, 12, 212 | replotting, 70 |
| maximum size, 11 | ellipse drawing, 83, 85 |
| dynamic load, 14, 147 | ELLIPSE keyword, 85, 110, 233 |
| | elliptic integrals, 299 |
| ECHO keyword, 56, 59, 74 | first kind, 299 |
| EDGR, 47, 76, 84, 145, 155, 180, 182, | second kind, 299 |
| 257, 262, 272 | ELTIME function, 291 |
| EDGR command, 69 , 156 | ENABLE command, 55, 73, 209, 257 |
| CLOSE keyword, 69 | BORDER keyword, 73 |
| EDIT keyword, 69 | BROADCAST keyword, 73 |
| FRAME keyword, 70 | CONFIRM keyword, 74 |
| OPEN keyword, 69 | ECHO keyword, 74, 78 |
| edit a drawing file, 69 | local effect, 74 |
| EDIT keyword, 69 | HISTORY keyword, 74 |
| EI function, 300 | JOURNAL keyword, 57, 74, 135 |
| EIGEN function, 323 | PROMPTING keyword, 75 |
| eigenvalue, 323 | REPLAY keyword, 75 |
| eigenvector, 323 | REPLOT keyword, 75 |
| EINELLIC function, 299 | SHELL keyword, 75 |
| elapsed time, 291 | STACK keyword, 76, 247 |
| electrical engineering, 303 | ENDDO statement, 80 |
| elements of array string variable, 266 | ENDIF statement, 80 |
| eliminate | environment variable, 156, 270 |
| matrices, 44 | file name, 77, 135, 166, 185, 211, 247 |
| scalars, 44 | 270, 276 |
| string variables, 44 | environment variable translation, 314 |
| eliminate vectors, 44 | environmnet variable, 77 |
| ELLICE function, 299 | EQS function, 315 |
| ELLICK function, 299 | equation solve, 320 |
| ellipse | erase, 140 |
| drawing, 70 | erase alphanumeric, 13 |
| populate, 70 | erase dots, 120, 164, 173, 234, 241, 264 |

<u>INDEX</u>

| erase fill, 233 | example, 311 |
|--|---|
| erase graphics, 209, 352 | expand variable name, 45 |
| erase legend background, 140 | EXPINT function, 300 |
| erase line segment, 145 | EXPN function, 300 |
| erase text, 262 | exponential integrals, 300 |
| ERASEWINDOW command, 76, 209, 262 | exponential integrals of order n , 300 |
| ERF function, 299 , 306 | expression, 281 |
| ERFC function, 300 | with DESTROY, 45 |
| ERRBAR keyword, 63, 108, 230 | expression function, 14, 152 |
| ERRFILL keyword, 110, 188, 194, 233 | expression in fit, 94, 101 |
| error bars, 119, 209 | expression variable, 311, 316 |
| asymmetric, 119 | expansion, 311 |
| clipping, 120 | expression with COPY, 27 |
| shape, 119 | EXTENSION command, 77, 81 |
| symmetric, 119 | extrema, 248 |
| error function, 299, 301 | A . 7 |
| complementary, 300 | fast Fourier transform, 324, 329, 330 |
| inverse, 300 | coefficients, 325 |
| error trapping, 152 | prime factors, 326 |
| Euler's constant, 295, 296, 298, 299 | restrictions, 326 |
| EVAL function, 316 | FC keyword, 318, 331 |
| example, 316 | FERDIRAC function, 301 |
| evaluation forced, 316 | Fermi-Dirac function, 301 |
| executable files, 225 | Feynman diagram, 299 |
| EXECUTE command, 29, 76 , 81, 84, | FFT function, 267, 324 , 328 |
| 247, 257, 271 | example, 326 |
| abort, 78 | prime factors, 326 |
| comments, 78 | restrictions, 326 |
| conditional statements, 80 | FIGURE command, 56, 74, 82 , 108, 110, |
| DO loops, 80 | 111, 113, 230, 233–235 |
| echoing, 78 | ARROW keyword, 108 |
| filename extension, 77 | fillable figures, 82 |
| generalized parameters, 78 | units, 83 |
| parameter passing, 78 | X Windows, 83 |
| prompting, 79 | figure types, 82 |
| returning from, 78 | file stack, 247 |
| script library, 77 | filename extension, 77, 81, 225 |
| transferring control, 78 | fill area under a curve, 120 |
| EXIST function, 336 | fill histogram, 118 |
| example, 336 | FILL keyword, 83, 110, 233 |
| EXPAND function, 311 | fill pattern, 41, 83, 110, 233, 240 |

| fill with dot pattern, 120, 164, 173, 264, | weights, 97 |
|--|--------------------------------------|
| 353 | zeros, 98 |
| fill with hatch pattern, 120, 164, 173, 264, | FIT command, 94 |
| 353 | hint for physicists, 98 |
| fillable figures, 82 | parameters, 225 |
| filter | FIT\$CHISQ variable, 98, 99 |
| causal recursive, 87 | FIT\$CL variable, 100 |
| coefficients, 86 | FIT\$CORR variable, 99 |
| digital, 86 | FIT\$COVM variable, 99 |
| noise amplification, 87 | FIT\$E1 variable, 99 |
| nonrecursive, 86, 88, 335 | FIT\$E2 variable, 99 |
| recursive, 90 | FIT\$FREE variable, 98 |
| time-invariant, 87 | FIT\$VAR array string variable, 95 |
| FILTER command, 86 | FMIN command, 101 |
| examples, 91 | example, 102 |
| FINELLIC function, 299 | FOLD function, 218, 338 |
| FIOWA data sets, 212 | example, 338 |
| FIOWA histograms, 212 | folding a matrix, 338 |
| FIOWA scatterplots, 213 | font, 174 |
| first derivative, 317 | font default, 113, 242 |
| FIRST function, 337 | FONT keyword, 59, 108, 113, 180, 242 |
| FISHER function, 301 | font names, 113, 242 |
| Fisher's F-distribution function, 301 | font table, 113, 242 |
| fit, 6 | formatted text, 166 |
| confidence level, 100 | accents, 181 |
| degrees of freedom, 98, 100 | blank lines, 169 |
| expression, 94, 101 | bold, 172 |
| informational messages, 101 | character height, 169 |
| iterations, 100 | colour, 173 |
| method, 95 | command delimiters, 166 |
| normal, 96 | comments, 166 |
| chi-square, 97 | continuation lines, 168 |
| correlation matrix, 99 | emphasis mode, 179 |
| covariance matrix, 99 | filled, 172 |
| errors, 99 | font, 174 |
| weights, 97 | hexadecimal mode, 180 |
| parameters, 94 | horizontal space, 177 |
| Poisson, 98 | italics mode, 179 |
| chi-square, 99 | justification centre, 174 |
| tolerance, 96 | justification left, 175 |
| update, 101 | justification right, 176 |

| left margin, 171 | example, 104 |
|--|---|
| line spacing, 170 | method, 102 |
| slanted mode, 179 | |
| subscript mode, 178 | GAMMA function, 301 |
| superscript mode, 179 | gamma function, 301 |
| formatted text special characters, 166 | incomplete, 302 |
| Fourier coefficients, 325 | gamma functions |
| Fourier series, 325 | natural logarithm, 301 |
| Fourier transform, 267, 324 | GAMMACIN function, 302 |
| inverse, 325, 328 | GAMMATIN function, 302 |
| FRAME keyword, 70, 139 | GAMMLN function, 301 |
| frame within a drawing file, 70 | GAMMQ function, 302 |
| FREC1 function, 301 | GAUSS function, 297 |
| FREC2 function, 301 | Gauss series, 302 |
| free format, 186 | Gauss-Jordan method, 320 |
| FREQ function, 297 | Gauss-Newton method, 95 |
| FRES1 function, 301 | Gaussian, 329 |
| FRES2 function, 301 | Gaussian distribution |
| Fresnel integrals, 301 | integral, 299 |
| Fritsch and Carlson, 318 | normalized, 297 |
| interpolation, 331 | Gaussian function, 309 |
| FULL keyword, 63 | Gaussian probability function, 297 |
| full width half maximum, 334 | inverse, 298 |
| function, 288 | GAUSSIN function, 298 |
| array, 288 | GAUSSJ function, 320 |
| looping, 288 | examples, 320 |
| basic, 290 | general characteristics, 352 |
| define dummy variable, 225 | GENERAL keyword, 64 |
| element by element, 288 | generalized parameter, 312 |
| looping, 342 | GENERATE command, 105, 112, 237, |
| numeric, 288 | 291 |
| numeric analysis, 316 | random number, 106 |
| numeric with string argument, 314 | random number seed, 107 |
| return variable's characteristics, 336 | GENERIC keyword, 155 |
| shape changing, 338 | generic terminal driver, 155 |
| special mathematical, 294 | geometric figures, 82 |
| string, 288 | geometric mean, 249, 250 |
| that return a string, 309 | GET command, 63, 107 , 228, 261, 348 |
| trigonometric, 289 | and script files, 107 |
| FWHM, 334 | ERRFILL keyword, 188, 194 |
| F7FRO command 102 | FONT keyword, 59 |

| GPLOT keywords, 108 | graphics display device, 155 |
|--|--|
| PHYSICA keywords, 108 | colour, 20 |
| GKS, 52 | graphics editor, 69 |
| plotting units, 53 | graphics erase, 76 |
| global section, 115, 153 | graphics font, 113, 242 |
| GLOBALS command, 115, 154 | graphics hardcopy, 19, 84, 128, 140, 163 |
| GOTO statement, 77, 79, 269 | bitmap, 76 |
| GPLOT defaults, 29 | HPLaserJet, 48 |
| GPLOT keywords, 108, 228, 230 | InkJet, 48 |
| GPLOT menu, 63, 230 | LA100, 49 |
| x-axis, 64 | PostScript, 49 |
| y-axis, 64 | Printronix, 49 |
| full, 63 | ThinkJet, 49 |
| general, 64 | graphics orientation, 156 |
| short, 64 | graphics output disable/enable, 155 |
| GPLOT window, 273 | graphics page, 272 |
| GR1105 keyword, 155 | graphics pixels, 352 |
| graph | graphics window |
| autoscaling, 109 | borders, 55, 73 |
| graph axes, 208 | grave, 181 |
| GRAPH command, 109, 111, 113, 115 , | greek letters, 261 |
| 136-138, 208, 218, 231, 235, 238- | grey scale, 240, 241 |
| 240 | GRID command, 124 |
| axes only, 116 | duplicate points, 124 |
| data and axes, 116 | non-interpolated grid example, 126 |
| data only, 116 | output matrix size, 125 |
| error bars, 119 | output vectors, 125–127 |
| examples, 121 | polar coordinates, 124 |
| filling, 120 | range of interpolation, 125 |
| histogram, 117 | sparse data, 126 |
| legend entry, 116 | sparse data example, 127 |
| polar coordinates, 119 | grid lines, 359, 368 |
| replot data, 117 | и . В и од |
| graph commensurate axes, 226 | Hamming, R.W., 87 |
| graph coordinates, 275 | Hankel transform, 267 |
| graph legend, 117, 137 | hardcopy |
| graph scales, 226 | device type, 114 |
| graph units, 83, 140, 163, 261 | page, 272 |
| graphics clear, 19, 47, 156, 209 | plot file, 257 |
| graphics cursor, 84, 113, 139, 157-159, | HARDCOPY command, 4, 128 |
| 161 162 257 273 354 | examples, 130 |

| print and save codes, 128 harmonic oscillator, 302 HATCH keyword, 59, 111, 234, 241 hatch pattern, 41, 83, 110, 120, 172, 173, 233, 240, 241, 264 defaults, 30 hatch pattern definition, 164, 173 hatch pattern fill, 59, 353 hatch pattern redefining, 111, 234 HBOOK data set, 214 histograms, 216 listing, 215 | hardcopy device, 48 plotting units, 48 HPPlotter plotting units, 52 hypergeometric function, 302, 304 logarithmic confluent, 302 HYPGEO function, 302 IµSR data sets, 223 IATYPE array, 15, 148 ICHAR function, 311, 315 ICLOSE function, 337 ICODE array, 15, 148, 152 |
|---|--|
| Ntuples, 215 | IDENTITY function, 323 |
| scatterplots, 217 | identity matrix, 323 |
| HELP command, 130 library user defined, 130 | IEQUAL function, 337 |
| paging the output, 130 | IER variable, 16, 149 |
| HERMITE function, 302 | IF block, 77, 80, 134, 269 |
| Hermite polynomials, 302 | IF statement, 269 |
| histogram, 115, 240, 353 | IFF keyword, 27, 46 |
| bar | IFFT function, 325, 328 |
| appearance, 108, 238 | IMAGEN, 52 |
| colour, 242 | plotting units, 52 |
| size, 115, 118 | IMSR data sets, 223 |
| width, 241 | inches, 83, 111, 112, 140, 163, 234, 236, |
| colour, 115, 118 | 261 |
| fill, 111, 118, 234 | incomplete β function, 252, 295 |
| fill pattern, 240 | incomplete Γ function, 302 |
| grey scale, 241 | incomplete gamma function, 302 |
| hatch fill, 115, 241 | index |
| plotting, 117 | range, 45 |
| type, 117, 209 | INDEX function, 315 |
| HISTORY keyword, 56, 74 | example, 316 |
| HISTYP keyword, 117, 239, 353 | influence function, 6 |
| HLDIR routine, 215 | initialization file, 29 InkJet |
| Houston | |
| plotting units, 52 | hardcopy device, 48 plotting units, 49 |
| HP PaintJet | inner product operator, 285 , 321 |
| bitmap device, 48 | example, 285, 286 |
| HPLaserJet | INPUT command, 131 |
| graphics resolution, 48 | na or communa, 101 |

| and script files, 131 | JAHNUF function, 308 |
|--|--------------------------------------|
| input line | JOIN function, 336 |
| recall buffers, 11, 58, 75, 212, 253 | JOURNAL command, 57, 75, 135 |
| control keys, 11 | initial defaults, 135 |
| INQUIRE command, 133 | journal file, 59 |
| examples, 134 | JOURNAL keyword, 57, 74 |
| inside a polygon, 182 | justification of text, 258 |
| INTEGRAL function, 112, 237, 318 | |
| example, 319 | KEI function, 303 |
| smoothing splines, 319 | Kelvin functions, 303 |
| integral transform, 267 | first kind, 303 |
| integrating recursive filters, 90 | second kind, 303 |
| integration, 318 | KER function, 303 |
| interactive input, 131 | keyboard focus, 4 |
| INTERP function, 112, 237, 330, 332 | keypad buffer, 11, 12, 212 |
| Fritsch and Carlson, 331 | KEYWORD command, 135 |
| Lagrange interpolation, 331 | wildcard, 136 |
| linear interpolation, 331 | keywords, 3 |
| spline interpolation, 331 | Kummer's function, 303 |
| INTERP keyword, 317 | kurtosis, 249, 251 |
| interpolating nonrecursive filters, 89 | LA100 |
| interpolating polynomial, 318 | bitmap device, 49 |
| interpolation 2-dimensional, 124 | plotting units, 49 |
| interpolation using FFT, 325 | label, 77, 79 |
| intersection operator, 244, 283 | LABEL command, 136 , 357, 368 |
| invalid field on read, 110, 233 | \XAXIS qualifier, 218 |
| inverse χ^2 probability function, 297 | example, 137 |
| inverse complementary error function, 300 | turn off label, 136 |
| inverse error function, 300 | where to draw, 136 |
| inverse Fourier transform, 325 | labeled tic marks, 226 |
| INVERSE function, 321 | LABSIZ keyword, 110, 232 |
| inverse Gaussian probability function, 298 | Lagrange interpolation, 318, 331 |
| inverse normal probability function, 298 | LAGRANGE keyword, 317, 331 |
| inverse of a matrix, 321 | LAGUERRE function, 303 |
| invert matrix, 346 | Laguerre polynomials, 303 |
| IRIX, 211 | LANDSCAPE keyword, 156 |
| IUPDATE array, 16, 149 | Laplace transform, 267 |
| IACODI Carattara 202 | LaserJet III, 128 |
| JACOBI function, 303 | LaserJet IIP, 128 |
| Jacobi polynomials, 303 | LAST function, 337 |
| Jahn's U function, 306, 308 | ETEX output, 130 |
| | |

| LCASE function, 310 | likelihood function, 95 |
|--|--|
| example, 310 | LINE command, 111, 113, 142 , 235 |
| leading zeros, 364, 374 | X Windows, 143 |
| least-squares fit, 6, 72, 335 | line graph |
| least-squares line, 333 | plotting symbols, 239 |
| least-squares residual, 225 | LINE keyword, 111, 235 |
| LEFNUM keyword, 382 | line thickness, 209, 352 |
| LEFTIC keyword, 382 | line type, 59, 82, 138, 209, 352 |
| LEGEND command, 116, 137 | defaults, 30 |
| example, 141 | redefining, 111, 235 |
| FRAME keyword, 56, 74 | styles, 235 |
| legend entry, 137 | linear correlation coefficient, 252 |
| clipping, 139 | linear interpolation, 331 |
| frame box, 139 | LINEAR keyword, 331 |
| coordinates, 139 | LINES keyword, 59 |
| coordinates units, 139 | lines through the origin, 279 |
| move, 140 | link, 147 |
| outline, 139 | LINTHK keyword, 352 |
| resize, 140 | LINTYP keyword, 352 |
| line segment, 138 | LINUX, 211 |
| plotting symbols, 138 | Linux, 3 |
| status, 141 | LIST command, 146 |
| string portion, 138 | literal string |
| text | write, 278 |
| height, 138 | LJ250 |
| title, 140 | bitmap device, 48 |
| height, 141 | LN03+, 52 |
| transparency, 140 | plotting units, 52 |
| legend format, 34, 39, 110, 233 | LOAD command, 14, 147 , 212 |
| LEGENDRE function, 304 | function, 152 |
| Legendre functions, 304 | function arguments, 152 |
| associated, 304 | function example, 152 |
| Legendre polynomials, 303, 304 | subroutine, 147 |
| LEGFRMT keyword, 34, 110, 233 | example, 150 |
| LEGSIZ keyword, 39, 110, 232 | local minimum, 101 |
| LEN function, 336 , 337 | LOGAM function, 301 |
| length of a vector, 270, 336 | logarithmic <i>x</i> -axis, 357, 361, 362 |
| Leo Tick formula, 90 | logarithmic y -axis, 368, 372, 373 |
| leptokurtic, 251 | logarithmic confluent hypergeometric func- |
| Lexmark printer, 33, 35, 36, 38, 121, 164, | tion, 302 |
| 173, 241, 264 | logical name, 29, 77, 156, 270 |

| logical name assignment, 5 | maximum matrix row index, 249 |
|-------------------------------------|---|
| logical name translation, 314 | maximum vector index, 249 |
| logical search list, 77 | mean, 249 |
| login command file, 29 | mean deviation, 251 |
| LOGIN.COM, 384, 389 | mean filter, 87 |
| LOOP function, 218, 225, 345 | mean value, 250 |
| examples, 346 | median, 249 |
| looping function, 288, 342-346 | median filter, 87 |
| loops, 80 | median value, 250 |
| Lorentzian function, 309 | MENU command, 230 |
| lowercase conversion, 310 | MENU keyword, 62 |
| LU decomposition, 321, 322 | mesokurtic, 251 |
| 404 | MIN function, 293 |
| macron, 181 | minima, 157 |
| MAP command, 115, 153 | minimum, 249 |
| MASK keyword, 63, 108, 230 | minimum local, 101 |
| math symbols, 261 | minimum matrix column index, 249 |
| matrix | minimum matrix row index, 249 |
| change size, 154 | minimum vector index, 249 |
| column interchange, 282 | MOD function, 292 |
| creation, 154 | modified Bessel functions, 296, 303 |
| determinant, 322 | modulus function, 292 |
| folding, 338 | monitor colour, 352 |
| input, 132 | MONITOR command, 58, 75, 155 |
| inverse, 321 | monotone piecewise cubic interpolation, 318 |
| invert, 346 | MSR data sets, 223 |
| listing, 146 | MUD data sets, 221 |
| LU decomposition, 321, 322 | Muller's method, 102 |
| maxima, 161 | Muller, D.E., 103 |
| minima, 161 | multinomial distribution, 98 |
| reflect, 282 | multiple graphs, 272 |
| slices, 245 | multiple pages, 48 |
| transpose, 282 | NOVEN 1 |
| unfolding, 338 | NCURVES keyword, 113, 238 |
| MATRIX command, 154 | NES function, 315 |
| MAX function, 293 | nested IF blocks, 80 |
| MAXHISTORY keyword, 114, 236 | network read, 58, 76 |
| maxima, 157 | new features, 156 |
| maximum, 249 | NEWS command, 156 |
| maximum likelihood function, 95 | NLXINC keyword, 29, 360 |
| maximum matrix column index, 249 | NLYINC keyword, 29, 371 |

| noise amplification by filter, 87 | outer product operator, 284 |
|--|---|
| nonrecursive filter, 86, 88, 335 | example, 285 |
| nonrecursive filter differentiating, 88 | output device, 6, 155 |
| nonrecursive filter interpolating, 89 | outside a polygon, 182 |
| nonrecursive filter smoothing, 89 | D. 4 7 |
| normal distribution, 94 | PaintJet |
| normal distribution of errors, 96 | bitmap device, 48 |
| NORMAL function, 297 | parameter in fit, 94 |
| normal probability function, 297 | parameter passing, 312 |
| normal probability function inverse, 298 | partial derivative, 24 |
| normalized gaussian distribution, 297 | particle physics, 299 |
| normalized tina resolution, 306 | pause during script execution, 271 |
| NSXINC keyword, 29, 357, 360 | PCHAR keyword, 4, 62, 108, 238 |
| NSYINC keyword, 29, 368, 371 | pcm extension, 77, 81 |
| NSYMBOLS keyword, 138 | PEAK command, 113, 157 |
| Ntuples, 215 | default code keys, 158 |
| NUMBLD keyword, 353 | X Windows, 157 |
| numeric evaluation, 316 | Pearson's r, 252 |
| NXDEC keyword, 364, 365 | pen plotter, 50 |
| NXDIG keyword, 364, 365 | pen plotter speed, 112, 238 |
| NXGRID keyword, 359 | penalty function, 6 |
| NYDEC keyword, 375, 375 | perspective projection, 4 |
| NYDIG keyword, 375, 375 | PFACTORS function, 324, 326 |
| NYGRID keyword, 368 | phys_user.f, 391 |
| 111 d2022 1165 1161 d, CCC | PHYSICA defaults, 29, 64 |
| object module, 14, 147 | PHYSICA keywords, 63, 108, 228, 230 |
| OFF keyword, 138, 139, 155, 232 | PHYSICA menu, 63 |
| ON keyword, 138, 139, 155, 232 | PHYSICA version number, 253 |
| on-line help, 130, 136 | PHYSICA.JOURNAL, 135 |
| open a drawing file, 69 | PHYSICA\$DIR, 18 |
| OPEN keyword, 69 | PHYSICA\$INIT file, 29 |
| opening an EDGR file, 156 | PHYSICA\$LIB, 77 |
| OpenVMS, 3 | PHYSICA\$LIB, 270 |
| operator | PHYSICA_INIT file, 29 |
| Boolean, 281 | PHYSICALIB, 270 |
| operator append, 314 | PHYSICA_USER_FUNCTIONS, 14, 18 |
| order property, 283 | PHYSICA_USER_FUNCTIONS.COM, 384, 389 |
| ORIENTATION command, 69, 156, 212, | PHYSICA_USER_FUNCTIONS.FOR, 15, 148, |
| 225 | 391 |
| initial default, 156 | PICK command, 11, 111, 113, 159 , 183, |
| OSF/1, 153 | 235 |

| automatic digitizing, 160 | 209, 238, 353, 354 |
|---|---|
| examples, 160 | angle, 115 |
| choose matrix, 161 | centred, 239 |
| choose matrix maxima, 161 | colour, 115, 239 |
| choose matrix minima, 161 | connecting, 239 |
| choosing a polygon, 161 | maximum value, 239 |
| default code keys, 159 | rotation angle, 240 |
| regional counts, 162 | size, 115, 119, 239, 354 |
| regional counts matrix, 162 | special codes, 239 |
| X Windows, 159 | plotting units, 69, 84, 140, 143, 157, 163, |
| piecewise linear interpolants, 271 | 272 |
| piechart, 162 | plotting units type, 113 |
| coordinates, 162 | PMODE keyword, 63, 108, 230 |
| coordinates units, 163 | POICA function, 304 |
| wedge, 162 | Poisson distribution, 94 |
| wedge filling, 163 | Poisson distribution of errors, 98 |
| piechart wedge drawing, 84 | Poisson-Charlier polynomial, 304 |
| PIEGRAPH command, 111, 162, 234 | polar coordinates, 119, 124, 271 |
| example, 165 | POLYGON command, 182 |
| pixels, 352 | polygon drawing, 83, 85 |
| platykurtic, 251 | POLYGON keyword, 85, 110, 233 |
| PLM function, 304 | polygon vertices, 85 |
| PLMN function, 304 | PORTRAIT keyword, 156 |
| PLMU function, 304 | POSTRES keyword, 112, 237 |
| plot a histogram, 117 | PostScript |
| plot file, 84 | colour, 50 |
| plot keyword | erase, 209, 262 |
| axis box, 351 | grey scale, 50 |
| general, 348 | hardcopy, 352 |
| summary, 348 | hardcopy devices, 49 |
| text, 348 | plotting units, 49 |
| x-axis, 349 | resolution, 33, 35, 36, 38, 49, 112, |
| y-axis, 350 | 121, 164, 173, 237, 241, 264 |
| plotter, 6 | pre-defined windows, 273 |
| file, 262 | prime factors, 324, 326 |
| pen, 20 | print graphics, 128 |
| PLOTTEXT command, 138, 165 | Printronix |
| plotting axes only, 116 | bitmap device, 49 |
| plotting data and axes, 116 | plotting units, 49 |
| plotting data only, 116 | PROB function, 297 |
| plotting symbol. 29, 62, 108, 115, 138. | probability functions, 297 |

| bivariate normal, 297 | ASCII file examples, 196 |
|--|--------------------------------------|
| Gaussian, 297 | formatted read, 196 |
| inverse, 298 | unformatted file, 201 |
| integral of χ^2 distribution, 297 | unformatted file examples, 202 |
| normal, 297 | unformatted file read by record, 202 |
| inverse, 298 | unformatted file stream read, 202 |
| probability functions χ^2 , 297 | maximum record length, 184 |
| inverse, 297 | scalar, 193 |
| probability integral of χ^2 distribution, 297 | scalars |
| PROD function, 225, 343 | ASCII file, 194 |
| example, 343 | ASCII file examples, 194 |
| program input, 269 | formatted read, 194 |
| program version, 113 | invalid field, 194 |
| program version date, 114 | number field, 194 |
| projection, 4 | unformatted file, 195 |
| PROMPTING keyword, 57, 75 | unformatted file examples, 195 |
| PT100G keyword, 155 | text, 203 |
| PTYPE keyword, 63, 108, 230, 352 | ASCII file, 204 |
| | examples, 204 |
| quantum mechanics, 306 | unformatted file, 204 |
| quintic polynomial equations, 24 | vectors, 186 |
| QUIT command, 183 | ASCII file, 186 |
| Racah coefficients, 306, 307 | ASCII file examples, 188 |
| RACAHC function, 307 | binary files, 189 |
| Rademacher function, 305, 309 | column numbers, 186 |
| RADMAC function, 305 | comment lines, 187 |
| RAN function, 107, 112, 237, 291 | field counts, 187 |
| seed, 291 | formats, 186 |
| random number, 291 | invalid field, 187 |
| generation, 106 | line numbers, 186 |
| seed, 107, 112, 237, 291 | output variables, 187 |
| RCHAR function, 134, 313 , 314 | unformatted file, 189 |
| example, 314 | unformatted file examples, 191 |
| format, 313 | unformatted file field counts, 190 |
| read across a network, 58, 76 | unformatted file read by record, 189 |
| READ command, 184 | unformatted file stream read, 190 |
| close the file, 185 | REBIN command, 205 |
| informational messages, 185 | matrix, 206 |
| invalid field, 110, 233 | example, 207 |
| matrix, 195 | vector, 205 |
| ASCII file, 196 | examples, 206 |
| • | |

| rectangle drawing, 83, 85 | Roland |
|--|--|
| recursive filter, 90 | plotting units, 52 |
| recursive filter integrating, 90 | ROLL function, 339 |
| redraw on single graph, 208 | example, 339 |
| reflect operator, 282 | root-mean-square, 249, 250 |
| examples, 282 | root-mean-square error, 99 |
| REFRESH command, 208, 211 | rotation group, 303 |
| regular matrix, 124 | RPROD function, 225, 345 |
| RENAME command, 208 | example, 345 |
| REPLAY keyword, 57, 75 | RSUM function, 225, 344 |
| replot buffers, 19, 71, 209 | example, 344 |
| clear, 19 | run-time link, 147 |
| what is saved, 209 | run-time load, 14, 147 |
| REPLOT command, 19, 57, 70, 75, 117, | arguments, 147 |
| 136, 140, 208 , 212, 224, 231 | function, 152 |
| \TEXT qualifier, 209 | arguments, 152 |
| disable, 209 | example, 152 |
| enable, 209 | restrictions, 147 |
| examples, 210 | subroutine, 147 |
| replot buffers, 19 | example, 150 |
| text, 209, 261 | running mean filter, 87 |
| REPLOT keyword, 57, 75 | running median filter, 87 |
| reserved character names, 166 | running products, 345 |
| RESIZE command, 210 | running sums, 344 |
| RESTORE command, 11, 154, 211 , 224 | |
| μ SR MUD data set, 221 | SAVE command, 211, 224 |
| μ SR data set, 223 | save graphics in a file, 128 |
| CHAOS data set, 224 | SAVGOL function, 332, 334 |
| FIOWA data set, 212 | Savitzky-Golay filter, 332, 334 |
| examples, 214 | Savitzky-Golay smoothing |
| HBOOK data set, 214 | method, 335 |
| examples, 218 | scalar |
| $I\mu SR$ data set, 223 | append, 314 |
| PHYSICA session, 211 | dummy variable, 288, 342–345 |
| XFIOWA data set, 214 | SCALAR command, 94, 218, 225 , 342- |
| YBOS data set, 218 | 345 |
| examples, 220 | dummy variable, 225 |
| RETURN command, 78, 224 | fit parameter, 225 |
| RETURN from DCL, 28 | SCALES command, 22, 32, 115, 116, 226 |
| RITNUM keyword, 380 | , 232 |
| RITTIC keyword, 381 | example, 227 |

| CHARSZ keyword, 119, 239 |
|--|
| CNTSEP keyword, 22 |
| ERRFILL keyword, 188, 194 |
| examples, 229 |
| FILL keyword, 41, 83 |
| FONT keyword, 136, 174, 257, 258, |
| 261 |
| HATCH keyword, 30, 59, 110, 118, 120, |
| 164, 173, 234, 264 |
| how it works, 229 |
| LABSIZ keyword, 22 |
| LEGFRMT keyword, 39 |
| LEGSIZ keyword, 23, 34 |
| LINE keyword, 145, 352 |
| LINES keyword, 30, 62 |
| LINTYP keyword, 82, 111, 120, 235, |
| 240, 279 |
| PCHAR keyword, 62, 115, 117, 118, |
| 120, 139, 353, 354 |
| POSTRES keyword, 33, 35, 36, 38, 50, |
| 121, 164, 173, 241, 264 |
| SEED keyword, 107, 291 |
| TENSION keyword, 317, 319, 331-334 |
| TXTHIT keyword, 138, 141, 258, 261 |
| UNITS keyword, 69, 83, 110-112, 140, |
| 143, 157-159, 163, 169-171, 175- |
| 177, 232, 234, 236, 275 |
| WIDTH keyword, 64 |
| XUAXIS keyword, 23, 34, 39 |
| shareable image, 14 |
| command procedure, 384, 389 |
| creating, 18 |
| function, 152 |
| source code, 18, 391 |
| shared memory, 153, 154 |
| SHELL keyword, 58, 75 |
| shift elements, 339-341 |
| SHORT keyword, 64 |
| SHOW command, 57, 74, 243 , 283 |
| examples, 244 |
| history lines |
| |

| display, 114, 236 | SPLSMOOTH function, 112, 237, 332, 333 |
|---|---|
| max number, 114, 236 | |
| wrap, 114, 237 | with weights, 334 |
| SHOWHISTORY keyword, 114, 236 | STACK command, 58, 76, 84, 247, 257 |
| SIGN function, 292 | append to file, 248 |
| sine integral, 298 | simultaneously execute commands, 248 |
| SININT function, 298 | stack file, 225 |
| skewness, 248, 249, 251 | STACK keyword, 58, 76 |
| SLICES command, 231, 245 | standard deviation, 98, 99, 249, 251, 333 |
| example, 245 | static buffer, 11, 12, 212 |
| SMOOTH function, 112, 237, 330, 332, | STATISTICS command, 248 |
| 334 | central measures, 248 |
| method, 333 | dispersion, 248 |
| tension, 333 | examples, 252 |
| with weights, 332 | extrema, 248 |
| SMOOTH keyword, 317, 319 | linear correlation coefficient, 252 |
| smoothing | messages, 248 |
| Savitzky-Golay filter, 334 | moments, 251 |
| smoothing filter, 329, 330 | parameter definitions, 250 |
| smoothing nonrecursive filters, 89 | skewness, 248 |
| SOLARIS, 211 | weights, 250 |
| solve system of equations, 320 | STATUS command, 135, 253 |
| SORT command, 245 , 283, 284 | STATUS keyword, 141 |
| \UP qualifier, 244 | STEP function, 340 |
| associated vectors, 245 | example, 340 |
| examples, 246 | stop, 183 |
| spawning a subprocess, 28 | string |
| SPECIAL keyword, 59 | formatting, 138 |
| special names, 166 | writing, 278 |
| SPEED keyword, 112, 238 | string font, 136 |
| Spence's integral, 299 | string format, 141 |
| Spencer's formulae, 89 | STRING function, 313 |
| spline interpolation, 331 | string function |
| SPLINE keyword, 331 | CHAR, 311 |
| spline smooth | CLEN, 314 |
| weight, 333 | DATE, 310 |
| spline smooth method, 333 | EQS, 315 |
| spline smooth weight, 332 | EVAL, 316 |
| spline tension, 112, 237, 317, 319, 331, | EXPAND, 311 |
| 333 | ICHAR, 315 |
| SPLINTERP function, 112, 237, 330, 331 | INDEX, 315 |

| LCASE, 310 | tab, 136 |
|---------------------------------------|--|
| NES, 315 | Taylor expansion, 96, 97, 99 |
| RCHAR, 313 | TCASE function, 311 |
| STRING, 313 | example, 311 |
| SUB, 315 | TENSION keyword, 112, 237, 317 |
| SUP, 315 | terminal broadcast messages, 56 |
| TCASE, 311 | TERMINAL command, 78, 224, 256 , 269 |
| TIME, 310 | terminal interface, 12, 58, 75 |
| TRANSLATE, 314 | terminal width, 64, 112, 238 |
| UCASE, 310 | T _E X output, 130 |
| VARNAME, 312 | text angle, 258, 355 |
| VARTYPE, 312 | text append, 284 |
| string height, 138 | text bolding, 111, 234, 258 |
| string length, 315 | text colour, 258 |
| string variable, 45, 204 | TEXT command, 56, 74, 209, 257 , 354, |
| append, 314 | 355 |
| in expression, 311, 316 | confirm, 257 |
| Struve function, 305 | erase, 262 |
| second order, 305 | example, 262 |
| Struve function first order, 305 | script file, 257 |
| STRUVEO function, 305 | stack files, 257 |
| STRUVE1 function, 305 | text draw, 263 |
| STUDENT function, 305 | text emphasis, 258 |
| Student's t-distribution, 305 | text font, 29, 113, 242, 258 |
| Student's t-distribution inverse, 306 | text format, 59, 138, 258 |
| STUDENTI function, 306 | text height, 258, 261, 355 |
| SUB function, 315 | text hexadecimal codes, 258 |
| sub-window boundary, 272 | text horizontal space, 258 |
| SUBn keywords, 14 | text justification, 258, 354, 355 |
| sum, 249, 250 | text location, 258, 261 |
| SUM function, 225, 342 | text plot characteristics, 354 |
| example, 343 | text position, 355 |
| SUP function, 315 | text replot, 261 |
| SURFACE command, 254 | text special characters, 261 |
| colour figure, 255 | text sub-scripts, 258 |
| examples, 256 | text vertical spacing, 258 |
| symbol size, 4 | THEN keyword, 80 |
| symmetric error bars, 119 | Thiessen triangulation, 33, 35, 36, 38, 124, |
| symmetric matrix, 323 | 271 |
| syntax check, 311 | ThinkJet |
| system of equations, 320 | bitmap device, 49 |

| Thinkjet | example, 310 |
|---|---|
| plotting units, 49 | umlaut, 181 |
| tic marks, 109, 226, 227, 231, 232, 357, | unary operator, 282 |
| 359, 361, 362, 368, 370, 372, 373, | UNFOLD function, 338 |
| 380-382 | a matrix, 338 |
| labeled, 226 | example, 339 |
| tilde, 181 | unformatted binary file, 189, 195, 201, 204 |
| TILE command, 111, 234, 263 | union operator, 244, 283 |
| bar definition, 263 | UNIQUE command, 268 |
| example, 265 | examples, 268 |
| string definition, 264 | units, 175-177 |
| time elapsed, 291 | UNITS keyword, 113, 143, 242 |
| TIME function, 262, 310 | unity matrix, 323 |
| example, 310 | UNIX, 3, 14, 28, 29, 77, 131, 135, 156, |
| time-invariant filters, 87 | 166, 185, 211, 247, 270, 276, 314, |
| TINA function, 306 | 391 |
| TITLE keyword, 140 | update after fit, 101 |
| TK4010 keyword, 155 | uppercase conversion, 310 |
| TK4107 keyword, 155 | USE command, 269 |
| TLEN command, 266 | user defined library, 130 |
| example, 266 | user writte |
| toggle case conversion, 311 | subroutine, 212 |
| toggle graphics, 19 | user written |
| TOPNUM keyword, 381 | function, 152, 212, 225 |
| TOPTIC keyword, 381 | arguments, 147, 152 |
| TRANSFORM command, 267 | example, 152 |
| example, 267 | subroutine, 14, 147, 225 |
| TRANSLATE function, 314 | arguments, 147 |
| example, 314 | example, 15, 17, 148, 150 |
| TRANSPARENCY keyword, 140 | IATYPE array, 15, 148 |
| transpose operator, 196, 214, 218, 282 , | ICODE array, 15, 148 |
| 346 | IER variable, 16, 149 |
| example, 282 | IUPDATE array, 16, 149 |
| trapezoidal rule filter, 90 | matrix data, 17, 150 |
| trapping, 78 | name, 14 |
| triangle coefficient, 307 | numeric argument, 16, 149 |
| trigonometric functions, 289 | sharable image, 18 |
| TXTANG keyword, 258, 355 | string argument, 17, 149 |
| TXTHIT keyword, 169, 170, 355 | USERN function, 147, 152 |
| UCASE function, 134, 310 | variable, 224 |

| display, 243 | under a matrix, 271 |
|--|---|
| history, 56, 74 | VT241 keyword, 155 |
| variable name | VT640 keyword, 155 |
| expand, 45 | |
| variable parameters, 225 | WAIT command, 271 |
| variance, 87, 98, 249, 251 | WALSH function, 308 |
| variance-ratio distribution, 301 | wave function, 302 |
| VARNAME function, 312 | wedge drawing, 84 |
| example, 312 | WEDGE keyword, 84, 110, 233 |
| VARTYPE function, 312 | what is in this manual, 1–2 |
| VAX, 3 | WHERE function, 244, 283, 284, 337 |
| VAX/VMS, 14, 115, 147, 154 | example, 338 |
| vector | WIDTH keyword, 112, 238 |
| append, 284 | width of alphanumeric monitor, 112, 238 |
| copies, 27 | WIGN3J function, 307 |
| final index, 337 | WIGN6J function, 308 |
| first index, 337 | WIGN9J function, 308 |
| generation, 105 | Wigner $3 - j$ function, 307 |
| input, 131 | Wigner $6 - j$ function, 308 |
| intersection, 283 | Wigner $9 - j$ function, 308 |
| length, 270, 336, 337 | Wigner symbols, 306 |
| list, 146 | wildcard, 243 |
| paged, 146 | window, 83, 140, 163, 175-177, 209 |
| order property, 283 | commensurate graphs, 226 |
| ordered, 244 | definitions saved, 224 |
| reading, 186 | erase, 76 |
| sorting, 245 | physica, 211 |
| union, 283 | reset, 29 |
| VECTOR command, 270 | WINDOW command, 56, 74, 139, 163, 226, |
| vector coupling coefficients, 306 | 272 |
| VERSION keyword, 113 | boundaries, 272 |
| VERSIONDATE keyword, 114 | define new window, 272 |
| vertex coordinates, 317 | display window coordinates, 272 |
| virtual maximum, 109, 227, 232 | multiple window creation, 274 |
| virtual memory space, 58, 75 | plotting units, 272 |
| virtual minimum, 109, 227, 232 | pre-defined windows, 273 |
| VLEN function, 337 | relation to GPLOT, 273 |
| VMS, 3, 5, 14, 28, 29, 56, 73, 77, 130, 131, | what are windows, 272 |
| 153, 156, 211, 270, 314, 391 | world boundary, 272 |
| Voigt profile function, 309 | WORLD command, 275 |
| VOLUME command. 271 | world coordinate system, 156, 157, 261, |

| 275, 377 | XLWIND keyword, 171, 175-177, 274, 355, |
|--|--|
| world units type, 113, 242 | 377 , 378, 379 |
| WRAP function, 341 | XMAX keyword, 279, 357, 359, 361, 361 |
| example, 341 | XMIN keyword, 279, 357, 359, 362, 362 |
| WRAP keyword, 114, 237 | XMOD keyword, 363 , 364 |
| WRITE command, 275 | XNUMA keyword, 365 |
| appending to a file, 276 | XNUMSZ keyword, 4, 29, 365 , 380, 381 |
| format, 276 | XOFF keyword, 364, 364 |
| matrix, 278 | XPAUTO keyword, 364 |
| scalars, 277 | XPOW keyword, 364, 365 |
| examples, 278 | XPREV keyword, 112, 238 |
| maximum number, 278 | XTICA keyword, 360, 360 |
| string, 278 | XTICL keyword, 360 , 380, 382 |
| text | XTICS keyword, 361 , 380, 382 |
| examples, 278 | XTICTP keyword, 359 |
| unformatted file, 276 | XUAXIS keyword, 29, 34, 39, 378 |
| vectors, 276 | XUWIND keyword, 171, 175-177, 274, 355, |
| examples, 277 | 378, 378 , 379 |
| maximum number, 276 | XVMAX keyword, 361 |
| | XVMIN keyword, 362 |
| X keyword, 155 | XZERO keyword, 359 |
| X Window System, 4, 5, 143, 157, 159 | • |
| cursor readout, 242 | YAXIS keyword, 64, 366 |
| units type, 113 | YAXISA keyword, 360, 366, 371, 376, 379 |
| FIGURE command, 83 | |
| graphics replay, 57, 75 | YBOS data sets, 218 |
| refresh, 208 | YBOS dotplots, 219 |
| zoom window, 208 | YBOS histograms, 219 |
| XAXIS keyword, 64, 355 | YCNT matrix, 23 |
| XAXISA keyword, 354, 360, 366, 371, 376, | YCROSS keyword, 370 |
| 379 | YITICA keyword, 376 , 377 |
| XCNT matrix, 23 | YITICL keyword, 376, 376 |
| XCROSS keyword, 359 | YLABSZ keyword, 136, 368 |
| XFIOWA data sets, 214 | YLAXIS keyword, 29, 379 |
| XITICA keyword, 366 | YLEADZ keyword, 374 |
| XITICL keyword, 366 | YLOC keyword, 258, 354, 355 |
| XLABSZ keyword, 136, 357 | YLOG keyword, 227, 368, 371, 372 |
| XLAXIS keyword, 29, 378 | YLWIND keyword, 22, 110, 169, 170, 232, |
| XLEADZ keyword, 364 | 233, 274, 354, 355, 357, 360, 361, |
| XLOC keyword, 258, 354, 355 | 365, 366, 368, 371, 372, 376, 377, |
| XLOG keyword, 227, 357 , 360 | 378 , 379 |

```
YMAX keyword, 279, 368, 370, 372, 372
YMIN keyword, 279, 368, 370, 373, 373
YMOD keyword, 374, 374
YNUMA keyword, 376
YNUMSZ keyword, 29, 376, 380, 382
YOFF keyword, 374, 374
YPAUTO keyword, 374
YPOW keyword, 374, 375
YPREV keyword, 113, 238
YTICA keyword, 370, 370
YTICL keyword, 371, 381, 382
YTICS keyword, 372, 381, 382
YTICTP keyword, 370
YUAXIS keyword, 29, 32, 379
YUWIND keyword, 22, 110, 169, 170, 232,
      233, 274, 354, 355, 357, 360, 361,
      365, 366, 368, 371, 372, 376, 377,
      378, 379
YVMAX keyword, 372
YVMIN keyword, 373
YZERO keyword, 370
ZEBRA, 215
ZEBRA RZ, 214
zero filled vector, 270
ZEROLINES command, 111, 235, 278
   example, 279
   line type, 279
```