# GAUSS ${ }^{\text {™ }}$ 

## User Guide

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## Introduction

### 1.1 Product Overview

GAUSS $^{\text {TM }}$ is a complete analysis environment suitable for performing quick calculations, complex analysis of millions of data points, or anything in between. Whether you are new to computerized analysis or a seasoned programmer, the GAUSS family of products combine to offer you an easy to learn environment that is powerful and versatile enough for virtually any numerical task.

Since its introduction in 1984, GAUSS has been the standard for serious number crunching and complex modeling of large-scale data. Worldwide acceptance and use in government, industry, and the academic community is a firm testament to its power and versatility.

The GAUSS System can be described several ways: It is an exceptionally efficient number cruncher, a comprehensive programming language, and an interactive analysis environment. GAUSS may be the only numerical tool you will ever need.

### 1.2 Documentation Conventions

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

| Text Style | Use | Example |
| :---: | :---: | :---: |
| regular text | narrative | "... text formatting is used ..." |
| bold text | emphasis | "...not supported under UNIX." |
| italic text | variables | "... If vnames is a string or has fewer elements than $x$ has columns, it will be ..." |
| monospace | code example | ```if scalerr(cm); cm = inv(x); endif;``` |
| monospace | filename, path, etc. | "...is located in the examples subdirectory..." |
| monospace bold | reference to a GAUSS command or other programming element within a narrative paragraph | "...as explained under create..." |
| Small Caps | reference to section of the manual | "...see Operator Precedence, Section 11.7..." |

## Getting Started

### 2.1 Installation Under UNIX/Linux

1. Make a directory to install GAUSS in.
2. cd to that directory.
3. Gunzip the .gz file if there is one.
4. Untar the .tar file.
5. Run the executable script ginstall.
6. Put the installation directory in the executable path.
7. Put the installation directory in the shared library search path.
8. Install the license. (To receive a license and license installation instructions, email license@aptech.com.)

For last-minute information, see README. term.

### 2.2 Installation Under Windows

### 2.2.1 Machine Requirements

- A Pentium or AMD computer or equivalent.
- Operating System and Memory (RAM) requirements:
- Windows NT4.0, SP6 IE4.0, 32 MB minimum 256 MB recommended.
- Windows 2000, 64 MB minimum, 256 MB recommended.
- Windows XP, 128 MB minimum, 256 MB recommended.
- Minimum of 100 MB free hard disk space, more may be needed depending on the size of matrices and the complexity of the program.
- Monthly defragmenting is recommended.


### 2.2.2 Installation from Download

For download instructions, email info@aptech.com.

### 2.2.3 Installation from CD

Insert the GAUSS compact disc into the CD-ROM drive, and setup should start automatically. If setup does not start automatically, click Start, then click Run. Type D: \setup. exe in the dialog box (where $D$ is the drive letter of the CD-ROM drive).

You can use this procedure for the initial installation of GAUSS, and for additions or modifications to GAUSS components.

To receive a license and license installation instructions, email license@aptech.com.

## Using the Command Line Interface

TGAUSS is the command line version of GAUSS. The executable file, tgauss is located in the GAUSS installation directory.

The format for using TGAUSS is:
tgauss flag(s) program program...
-b Execute file in batch mode and then exit. You can execute multiple files by separating file names with spaces.
-1 logfile Set the name of the batch mode log file when using the -b argument. The default is tmp/gauss.log\#\#\#, where \#\#\# is the process ID.
-e expression Execute a GAUSS expression. This command is not logged when GAUSS is in batch mode.
-o Suppress the sign-on banner (output only).
-T Turn the dataloop translator on.
-t Turn the dataloop translator off.

### 3.1 Viewing Graphics

GAUSS generates .tkf files for graphical output. The default output for graphics is graphic.tkf. On Windows, you can use vwr.exe to view the graphics file; on UNIX/Linux/Mac, you can use vwrmp. Two functions are available to convert . tkf files to PostScript for printing and viewing with external viewers: the tkf2ps function will convert .tkf files to PostScript (.ps) files, and the tkf2eps function will convert . tkf files to encapsulated PostScript (.eps) files. For example, to convert the file graphic.tkf to a postscript file named graphic.ps use:

```
ret = tkf2ps('`filename.tkf',, '`filename.ps'`)
```

If the function is successful it returns 0 .

### 3.2 Command Line History and Command Line Editing

When you run a command at the TGAUSS prompt, it is added to your command line history, which is stored in a file called . gauss_prompt_history in your \$(HOME) directory on UNIX/Linux or in your \$ (HOMEDRIVE) <br>\$ (HOMEPATH) directory on Windows. A separate history for commands entered in the command line debugger is stored in a file called .gauss_debug_prompt_history in the same directory. By default, the last 500 commands executed at the TGAUSS and debugger command lines are stored in these files. You can change this number by changing prompt_hist_num in your gauss.cfg file. The following keystrokes are supported for movement and editing at the command line and for retrieving the command line history:

### 3.2.1 Movement

$$
\begin{aligned}
& \text { LEFT ARROW or } \quad \text { Moves cursor left one character } \\
& \text { CTRL+B }
\end{aligned}
$$

RIGHT ARROW or CTRL+F

HOME or CTRL+A

END or CTRL+E

ALT+LEFT ARROW or CTRL+LEFT ARROW

ALT+RIGHT ARROW or CTRL+RIGHT ARROW

Moves cursor right one character

Moves cursor to beginning of line

Moves cursor to end of line

Moves cursor left one word

Moves cursor right one word

Deletes character at cursor

Deletes character left of cursor
CTRL+H

CTRL+U

CTRL+K

CTRL+X

ESC (Win only)
CTRL+V

CTRL+T

Deletes whole line
Pastes text from buffer to left of cursor
Cuts all characters left of cursor

Cuts all characters right of cursor, including cursor

Cuts whole line

Transposes character at cursor and character left of cursor

### 3.2.3 History Retrieval

| UP ARROW or CTRL+P | Retrieves previous line in history |
| :--- | :--- |
| DOWN ARROW or CTRL+P | Retrieves next line in history |
| PAGE UP or CTRL+W | Retrieves previous line in history that matches text <br> to left of cursor |
| PAGE DOWN or CTRL+S | Retrieves next line in history that matches text to left <br> of cursor |
| ALT+H or <br> OPTION+H (MAC only) | Prints prompt history to screen |
| !! | Runs last line in history |
| !num | Runs the line num before current line in history; <br> !-1 is equivalent to !! |
| !-num | Runs last line in history beginning with text |

Note that some of these keystrokes are mapped differently on different computers. For example, on some computers, SHIFT+RIGHT ARROW behaves the same as RIGHT ARROW, while ALT+RIGHT ARROW moves the cursor right one word. Therefore, multiple keystroke mappings have been supported to maximize the availability of these commands on any given machine.

### 3.3 Interactive Commands

### 3.3.1 quit

The quit command will exit TGAUSS.
The format for quit is:
quit

You can also use the system command to exit TGAUSS from either the command line or a program (see system in the GAUSS Language Reference).

The format for system is:

```
system
```


### 3.3.2 ed

The ed command will open an input file in an external text editor (see ed in the GAUSS Language Reference).

The format for ed is:
ed filename

### 3.3.3 browse

The browse command allows you to search for specific symbols in a file and open the file in the default editor. You can use wildcards to extend search capabilities of the browse command.

The format for browse is:
browse symbol

### 3.3.4 config

The config command gives you access to the configuration menu allowing you to change the way GAUSS runs and compiles files.

The format for config is:
config

## Run Menu

| Translator | Toggles on/off the translation of a file using dataloop. The translator |
| :--- | :--- |
| is not necessary for GAUSS program files not using dataloop. |  |

Translator line Toggles on/off execution time line number tracking of the original number tracking file before translation.

Line number Toggles on/off the execution time line number tracking. If the tracking translator is on, the line numbers refer to the translated file.

## Compile Menu

Autoload Toggles on/off the autoloader.
Autodelete Toggles on/off autodelete.
GAUSS Library Toggles on/off the GAUSS library functions.
User Library Toggles on/off the user library functions.
Declare Toggles on/off the declare warning messages during compiling. Warnings

Compiler Trace Includes the following options:

| Off | Turns off the compiler trace function. |
| :--- | :--- |
| File | Traces program file openings and closings. |
| Line | Traces compilation by line. |
| Symbol | Creates a report of procedures and the local and <br> global symbols they reference. |

### 3.4 Debugging

The debug command runs a program under the source level debugger.
The format for debug is:
debug filename

### 3.4.1 General Functions

? Displays a list of available commands.
q/Esc Exits the debugger and returns to the GAUSS command line.
+/- Disables the last command repeat function.

### 3.4.2 Listing Functions

1 number Displays a specified number of lines of source code in the current file.
lc Displays source code in the current file starting with the current line.
11 file line Displays source code in the named file starting with the specified line.
11 file Displays source code in the named file starting with the first line.
11 line Displays source code starting with the specified line. File does not change.
11 Displays the next page of source code.
lp Displays the previous page of source code.

### 3.4.3 Execution Functions

s number Executes the specified number of lines, stepping into procedures.
n number Executes the specified number of lines, stepping over procedures.
$\mathbf{x}$ number Executes code from the beginning of the program to the specified line count, or until a breakpoint is hit.
$\mathbf{g} \llbracket a r g s \rrbracket \quad$ Executes from the current line to the end of the program, stopping at breakpoints. The optional arguments specify other stopping points. The syntax for each optional argument is:
filename line period The debugger will stop every period times it reaches the specified line in the named file.
filename line $\quad$ The debugger will stop when it reaches the specified line in the named file.
filename ,, period The debugger will stop every period times it reaches any line in the named file.
line period The debugger will stop every period times it reaches the specified line in the current file.

| filename | The debugger will stop at every line in the named file. <br> line |
| :--- | :--- |
| The debugger will stop when it reaches the specified line <br> in the current file. |  |
| procedure period | The debugger will stop every period times it reaches the <br> first line in a called procedure. |
| procedure | The debugger will stop every time it reaches the first line <br> in a called procedure. |

$\mathbf{j} \llbracket \operatorname{args} \rrbracket \quad$ Executes code to a specified line, procedure, or period in the file without stopping at breakpoints. The optional arguments are the same as $\mathbf{g}$, listed above.
$\mathbf{j x}$ number Executes code to the execution count specified (number) without stopping at breakpoints.

0 Executes the remainder of the current procedure (or to a breakpoint) and stops at the next line in the calling procedure.

### 3.4.4 View Commands

v 【vars』 Searches for (a local variable, then a global variable) and displays the value of a specified variable.
$\mathbf{v} \$ \llbracket v a r s \rrbracket \quad$ Searches for (a local variable, then a global variable) and displays the specified character matrix.

The display properties of matrices and string arrays can be set using the following commands.
$\mathbf{r} \quad$ Specifies the number of rows to be shown.
c Specifies the number of columns to be shown.
num, num $\quad$ Specifies the indices of the upper left corner of the block to be shown.
w Specifies the width of the columns to be shown.
p Specifies the precision shown.
$\mathbf{f} \quad$ Specifies the format of the numbers as decimal, scientific, or auto format.
q Quits the matrix viewer.

### 3.4.5 Breakpoint Commands

lb Shows all the breakpoints currently defined.
b $\llbracket \arg s \rrbracket \quad$ Sets a breakpoint in the code. The syntax for each optional argument is:
filename line period The debugger will stop every period times it reaches the specified line in the named file.
filename line The debugger will stop when it reaches the specified line in the named file.
filename ,, period The debugger will stop every period times it reaches any line in the named file.

| line period | The debugger will stop every period times it reaches the <br> specified line in the current file. |
| :--- | :--- |
| filename | The debugger will stop at every line in the named file. |
| line | The debugger will stop when it reaches the specified line <br> in the current file. |
| procedure period | The debugger will stop every period times it reaches the <br> first line in a called procedure. |
| procedure | The debugger will stop every time it reaches the first line <br> in a called procedure. |

d $\llbracket a r g s \rrbracket \quad$ Removes a previously specified breakpoint. The optional arguments are the same arguments as $\mathbf{b}$, listed above.

## Introduction to the Windows Interface

The GAUSS graphical user interface is a multiple document interface. The interface consists of the Menu Bar, the Toolbar, edit windows, the Command Input-Output window, and the Status bar (see Figure 4.1).

### 4.1 GAUSS Menus

You can view the commands on a menu by either clicking the menu name or pressing ALT $+n$, where $n$ is the underlined letter in the menu name. For example, to display the File menu, you can either click File or press ALT+F.

### 4.1.1 File Menu

The File menu lets you access the file, printer setup, and exit commands. Some of these actions can also be executed from the toolbar. The File menu contains the following commands:


Figure 4.1: GAUSS Graphical User Interface

| New | Opens a new, untitled document in an Edit window. <br> Note: New, unsaved documents are not automatically backed up until you <br> save them, giving them a file name. |
| :--- | :--- |
| Open | Opens an existing file for viewing or editing. |
| Reload | Updates the active file. |
| Save | Saves your changes to the file in the active window. If the file is untitled, you <br> are prompted for a path and filename. |
| Save As | Saves your changes to the file in the active window using a new or different <br> path or file name. |
| Close | Closes the document in the active window. You are prompted to save the file <br> if it has been modified since you last saved it. |
| Close All | Closes all open files. You are prompted to save any file that has been <br> modified since you last saved it. |
| Run Program | Runs a GAUSS program file. |
| Insert File | Opens an existing text file and copies the contents into the active document. <br> This is similar to pasting text from the Windows clipboard. |
| Print | Prints the active file or selected text from the active window. |
| Print Setup | Specifies the printer you want to use. Other printer options, such as page <br> orientation and paper tray, are also accessed with this command. |
| Properties | Displays information about the active file. |
| Change Working | Changes the directory where GAUSS looks for the files it uses for normal <br> operation. This command does not affect the Open or Save As paths. |
| Directory |  |
| Clear Working | Clears the working directory list. |
| Directory List | Closes all open files and exits GAUSS. You are prompted to save any file <br> that has been modified since it was last saved. |
| Exit | GAUSS maintains a list of the ten most recent files you opened, at the end of <br> the File menu. If the file you want to open is on this list, click it and GAUSS <br> opens it in an Edit window. |
| Recent Files |  |

### 4.1.2 Edit Menu

The Edit menu lets you access the set of editing commands. Some of these actions can also be executed from the toolbar. The Edit menu contains the following commands:

| Undo | Restores your last changes in the active window. |
| :---: | :---: |
| Redo | Restores changes in the active window that you removed using the Undo Edit command. |
| Cut | Removes selected text from the active window and places it on the Windows clipboard. |
| Copy | Copies selected text from the active window to the Windows clipboard. |
| Paste | Copies text from the Windows clipboard to the active window at the cursor position. |
| Select All | Selects all text in the active window. |
| Clear All | Clears all text in the active window |
| Find | Finds the specified text in the active window. The search starts at the cursor position and continues to the end of the text in the active window. The search can be case sensitive or case insensitive. You may also limit the search to regular expressions. |
| Find Again | Resumes the search for the next occurrence of the text you specified in the previous Find action. Subsequent searches for the same text can also be performed by pressing F3. |
| Replace | Locates the specified text in the active window and replaces it with the text you entered in the "Replace with" field in the Search dialog box. The search starts at the cursor position and continues to the end of the text in the active window. The search can be case sensitive or case insensitive, and the replacement can be unique or global. |
| Insert Time/Date | Inserts the current time and date at the cursor position. GAUSS uses the time and date that appears in the Microsoft Windows Date/Time Properties window. |


| Go To Line | Moves the cursor to the specified line number. |
| :---: | :---: |
| Go To Next Bookmark | Moves to the next bookmark in the program. |
| Toggle Bookmark | Sets or clears existing bookmarks from the program. |
| Edit Bookmarks | Opens the Edit Bookmarks window. From the Edit Bookmarks window you can add, remove, or go to any set bookmark in a program. |
| Record Macro | Places a series of keystrokes into memory so that they can be called at a later date. For more information about recording macros see Using Keystroke Macros, Section 5.1.4. |
| Clear Macros | Clears macros from memory. |
| 4.1.3 View Menu |  |
| The View menu lets you toggle the Main Toolbar, the Status Bar, the Working Directory Toolbar, and the Debug Toolbar on or off. |  |
| Main Toolbar | Toggles the Main toolbar on or off. For more information about the Main toolbar, see Main Toolbar, Section 4.1.11. |
| Status Bar | The Status Bar is located along the bottom of the GAUSS window. For more information about the status bar, see Status Bar, Section 4.1.15. |
| Working Directory | Toggles the Working Directory toolbar on or off. For more information about the working directory toolbar, see Working Directory Toolbar, Section 4.1.12. |
| Debug Toolbar | Toggles the Debug toolbar on or off. For more information about the Debug toolbar, see Debug Toolbar, Section 4.1.13. |
| Window Toolbar | Toggles the Window toolbar on or off. For more information about the Window toolbar, see Window Toolbar, Section 4.1.14. |
| Error Output | Opens or closes the Error Output window. |
| Source View | Displays or undisplays the Source/Symbols window. |

Formation about the Main The Status Bar is located along the bottom of the GAUSS window. For more information about the status bar, see Status Bar, Section 4.1.15.

Toggles the Working Directory toolbar on or off. For more information about the working directory toolbar, see Working Directory Toolbar, Section 4.1.12.

Toggles the Debug toolbar on or off. For more information about the Debug toolbar, see Debug Toolbar, Section 4.1.13.

Toggles the Window toolbar on or off. For more information about the Window toolbar, see Window Toolbar, Section 4.1.14.

Opens or closes the Error Output window.
Displays or undisplays the Source/Symbols window.

### 4.1.4 Configure Menu

The Configure menu lets you customize the GAUSS environment.

| Preferences | Opens the General Preferences window. From the General Preferences <br> window you can define Run options, Compile options, DOS window <br> options, and Autosave options. For more information on configuring <br> GAUSS General Preferences, see Preferences Dialog Box, Section 5.6.1. |
| :--- | :--- |
| Editor | Opens the Editor Properties window. From the Editor Properties window <br> Properties |
| you can define colors and fonts, the language syntax, tabs, or general editor |  |
| properties. For more information on configuring editor properties, see |  |
| Editor Properties, Section 5.6.2. |  |

### 4.1.5 Run Menu

The Run menu lets you run the code you have entered, a block of code you selected, or the active file, depending on the operating mode.

| Insert GAUSS | Manually adds the GAUSS prompt ( $\gg$ ) at the cursor position. The GAUSS |
| :--- | :--- |
| prompt | prompt is automatically displayed following the execution of GAUSS code. |

Insert Last Cmd Re-enters the last command written to the Input buffer.

## Run Selected Text

Run Active File
Test Compile
Active File

Run Main File
Test Compile Test compiles the main file. During compilation, any errors are displayed in

| Main File | the Output window. <br> Note: This command is different than the GAUSS compile command, which <br> compiles a program and saves the pseudocode as a file. |
| :--- | :--- |
| Edit Main File | Opens the specified main file in an edit window. |
| Stop Program | Stops the program currently running and returns control to the editor. |
| Build GCG File | Creates GAUSS pseudocode file that can be run over and over with no <br> compile time. |
| from Main | Makes the active file the main file. |

### 4.1.6 Debug Menu

The Debug menu lets you access the commands used to debug your active file or main file.
The Debug menu contains the following Commands:

Debug Main File Runs the main file in the debugger.
Debug Active Runs the active file in the debugger.
File
Set/Clear Enables or disables a breakpoint at the cursor in the active file. Breakpoint

Edit Opens a list of all breakpoints in your program. The breakpoints are listed Breakpoints by line number. Any procedure breakpoints are also listed.

Clear All Removes all line and procedure breakpoints from the active file. Breakpoints

| Go | Starts the debugger. |
| :--- | :--- |
| Stop | Stops the debugger. |
| Run to Cursor | Runs the program until it reaches the cursor position. <br> Step Into |
| Runs the next executable line of code in the application and steps into |  |
| procedures. |  |

### 4.1.7 Tools Menu

The Tools menu lets you open GAUSS tools windows. The following commands can be used:

| Matrix Editor | Lets you create or edit data in a matrix (or grid). A cell can be edited by <br> typing in a new value and pressing ENTER. For more information see <br> Matrix Editor, Chapter 6. |
| :--- | :--- |
| Graphics Editor | Opens the Graphics Editor, which is an interactive TKF file editor. This <br> menu item will be inactive if you have not purchased the Graphics Editor. |
| Source Browser | Searches source files for string patterns. For more information see GAUSS <br> Source Browser, Chapter 8. |

Lib Tool Lets you manage the contents of libraries. For more information see Library Tool, Chapter 7.

DOS Opens a DOS Compatibility window.
Compatibility
Window

### 4.1.8 Window Menu

The Window menu commands let you manage your workspace. You can toggle the focus between all open windows using CTRL+TAB, or clicking in the window you want active. All open windows are listed at the end of the Window menu. The following commands can be used:

Cmd Window Makes the Command Input - Output window the active window.
Output Window Splits the output from the Command Input - Output window.
Debug Window Starts the debugger on the current file.
Re-use Window If checked, the next file browsed in the Source Browser will be displayed in the same window.

Command Log Loads the command log window into an editor.
Close All Closes all TKF File Viewer windows.
Graphics
\(\left.$$
\begin{array}{ll}\text { Dual Horizontal } & \begin{array}{l}\text { Horizontally tiles the program source and execution windows within the } \\
\text { main window, and minimizes all other windows. }\end{array} \\
\text { Dual Vertical } & \begin{array}{l}\text { Vertically tiles the program source and execution windows within the main } \\
\text { window, and minimizes all other windows. }\end{array} \\
\text { Cascade } & \begin{array}{l}\text { Arranges all open windows on the screen, overlapping each, with the active } \\
\text { window on top. }\end{array}
$$ <br>

Tile Horizontal \& Arranges all open windows horizontally on the screen without any overlap.\end{array}\right\}\)| Arranges all open windows vertically on the screen without any overlap. |
| :--- |
| Tile Vertical |
| Arrange Icons |
| Arranges all minimized windows across the bottom of the main GAUSS |
| window. |
| Horizontally | | Splits the active window into two horizontal panes. This allows you to view |
| :--- |
| two different areas of the same document to facilitate split-window editing. |
| Note: You can move the splitter bar by dragging it with the mouse. You can |
| remove the splitter bar from the window by dragging it to the end of the |
| window. |


| Split Vertically | Splits the active window into two vertical panes. This allows you to view <br> two different areas of the same document to facilitate split-window editing. <br> Note: You can move the splitter bar by dragging it with the mouse. You can <br> remove the splitter bar from the window by dragging it to the end of the <br> window. |
| :--- | :--- |
| Open Window | GAUSS maintains a list of all the windows you have opened at the end of <br> Lhe Window menu. If the window you want to view is on this list, click it and <br> List becomes the active window. |

### 4.1.9 Help Menu

The Help menu lets you access information in the GAUSS Help system. The GAUSS Help menu contains the following Commands:

User's Guide Accesses the online GAUSS User's Guide.
Keyboard Accesses the list of keystrokes you can use for cursor movement, editing, and text selection.

Reference Accesses the online GAUSS Language Reference, which contains the syntax for each GAUSS command.

Tip of the Day Displays a tip to help you make better use of the features available in the GAUSS Windows Interface.

About GAUSS... Provides information about your version of GAUSS, your license type and ID, as well as copyright information.

### 4.1.10 GAUSS Toolbars

The toolbar buttons let you have fast access to the most commonly used commands. Place the mouse pointer over the button to display a description of the command.


Figure 4.2: Main Toolbar

### 4.1.11 Main Toolbar

| New | Opens a new, untitled document in an Edit window. <br> Note: New, unsaved documents are not automatically backed up until you <br> save them, giving them a file name. |
| :--- | :--- |
| Open | Opens an existing file for viewing or editing. <br> Saves your changes to the file in the active window. If the file is untitled, you <br> are prompted for a path and filename. |
| Save | Removes selected text from the active window and places it on the Windows <br> clipboard. |
| Cut | Copies selected text from the active window to the Windows clipboard. |
| Copy | Copies text from the Windows clipboard to the active window at the cursor <br> position. |
| Prints the active file or selected text from the active window. |  |

Text window.
Run Active $\quad$ Runs the active file. The file then becomes the main file.
File
Main File Displays the name of the main file and lets you quickly change the main file List to one of the files listed.

Run Main Runs the file specified in the Main File list.
File
Stop Program Stops the program currently running and returns control to the editor.
Test Compile Compiles the main file. During compilation, any errors are displayed in the
Main File
Output window.
Note: This command is different than the GAUSS compile command, which compiles a program and saves the pseudocode as a file.

Edit Main Opens the specified main file in an edit window.
File
Debug Main Runs the main file in the debugger.
File

### 4.1.12 Working Directory Toolbar

You can use the Working Directory toolbar to quickly change your working directory.


Figure 4.3: Working Directory Toolbar

Current $\quad$\begin{tabular}{l}
Displays the name of the current working directory and lets you quickly <br>
Working

$\quad$

change the working directory to one of the directories listed.
\end{tabular}

## Directory List

$\begin{array}{ll}\text { Change } \quad \text { Browses to a new directory. } \\ \text { Working } \\ \text { Directory } & \end{array}$

### 4.1.13 Debug Toolbar

You can use the Debug toolbar for quick access to commands while debugging a file.


Figure 4.4: Debug Toolbar
Go Starts the debugger.

Stop Stops the debugger.
Toggle Enables or disables a breakpoint at the cursor in the active file.
Breakpoint
Clear All $\quad$ Removes all line and procedure breakpoints from the active file.
Breakpoints

Set Watch Opens the Matrix Editor for watching changing variable data. For more information about viewing variables see Viewing Variables, Section 5.5.5.

Step Into Runs the next executable line of code in the application and steps into procedures.

Step Over Runs the next executable line of code in the application but does not step into procedures.
Step Out Runs the remainder of the current procedure and stops at the next line in the

Run to Cursor Runs the program until it reaches the cursor position.

### 4.1.14 Window Toolbar

You can use the Window toolbar for quick access to window commands.


Figure 4.5: Window Toolbar
Activate Cmd Makes the Command window the active window.
Window

| Activate/ | Splits the output from the Command Input-Output window, or deactivates |
| :--- | :--- |
| Deactivate | Output window. |
| Output Window |  |

Activate Debug Makes the Debug window the active window. Window

Source/Symbol Displays or undisplays the Source/Symbol window.
View
Error Output Opens or closes the Error Output window.
Window
Tile windows Tiles the active window and the Output or Command Input - Output horizontally window horizontally.

Tile windows Tiles the active window and the Output or Command Input - Output

## vertically window vertically.

```
Src Browser
replaces window
with new file
contents
If selected, the next file browsed in the Source Browser will be displayed in the same window.
```


### 4.1.15 Status Bar

The status bar is located along the bottom of the GAUSS window. The status of the windows and processes are shown on the status bar.

### 4.1.16 GAUSS Status

The first section of the status bar shows the current GAUSS status. From time to time you are alerted to the task GAUSS is performing by new messages appearing in the status bar.


Figure 4.6: Status Bar

Cursor Location The line number and column number where the cursor is located appear on the status bar for the active window. When a block of text is selected, the values indicate the first position of the selected text.

DATALOOP DATALOOP appears on the status bar to indicate the Dataloop Tranlator is turned on.

OVR OVR appears on the status bar when typing replaces the existing text with text you enter. When OVR does not appear on the status bar, typing inserts
text without deleting the existing text. Press the INSERT key to toggle between the two conditions.

CAP

NUM

CAP appears on the status bar to indicate the Caps Lock key has been pressed and all text you enter will appear in upper case.

NUM appears on the status bar to indicate the Num Lock key has been pressed and the keypad numbers are active.

## Using the Windows Interface

## 5

The GAUSS graphical user interface is a multiple document interface. The interface consists of edit windows and the Command Input - Output window. Integrated into GAUSS is a full debugger with breakpoints and watch variables. The GAUSS graphical user interface also incorporates the Matrix Editor (see Chapter 6), Library Tool (see Chapter 7), and GAUSS Source Browser (see Chapter 8), as well as a context-sensitive HTML Help system (see Chapter 9).

### 5.1 Using the GAUSS Edit Windows

The GAUSS edit windows provide syntax color coding and auto-formatting as well as easy access to the Matrix Editor and Library Tool, and include an integrated context-sensitive help system accessible through the F1 key.

The edit windows provide standard text editing features like drag and drop text editing, and find and replace. The editor also lets you set bookmarks, define keystroke macros, find and replace using regular expressions, and run selected text from the editor.

### 5.1.1 Editing Programs

To begin editing, open an edit window by browsing to the source file, or by typing edit and the filename in the Command Input - Output window. If more than one file is open, the last file opened or run becomes the active window.

### 5.1.2 Using Bookmarks

Bookmarks are efficient placeholders used to identify particular sections or lines of code. To add or remove bookmarks, place the cursor in the line you want to bookmark and then press CTRL+F2, or click Toggle Bookmark on the Edit menu. You can jump to the next bookmark by pressing F2, or go to the previous bookmark by pressing SHIFT+F2.

To edit a list of all currently defined bookmarks, click Edit Bookmarks on the Edit menu. The Edit Bookmarks window allows you to add, remove, name or select the bookmark to which you wish to jump.

### 5.1.3 Changing the Editor Properties

You can customize the formatting of your code and text by changing font colors, fonts, adding line indentations, and adding line numbering to your programs. To access these properties, on the Configure menu click Editor Properties, or right-click on an edit window and click Properties on the context menu.For more information about the Editor Properties see Editor Properties, Section 5.6.2.

### 5.1.4 Using Keystroke Macros

GAUSS will save up to 10 separate keystroke macros.
To record a keystroke macro, press CTRL + SHIFT + R, or click Record Macro on the Edit menu. When you start recording the macro, a stop button will appear in the GAUSS window.

You create a macro by clicking Record Macro and pressing the keystrokes you want recorded. Once you have completed recording the macro, you can stop recording with the stop button. Once you have finished recording the macro, you can select one of ten macro names for it.

Use the following guidelines when creating and using your macro:

- Only keystrokes in the active window are recorded, not keystrokes in a dialog box.
- Only keystrokes are recorded, not mouse movements.

Macros are not saved when you close GAUSS.
If your macro is lengthy, consider creating a separate file and copying the information from the file into the active window, rather than using a macro to enter the information.

### 5.1.5 Using Margin Functions

The margin of the edit window can be used to show currently set bookmarks, currently set breakpoints, and line numbers. You can also select an entire line of text with a single click in the Selection Margin.

You can turn on or off the margin in the Misc tab of the Editor Properties dialog box.

### 5.1.6 Editing with Split Views

Using split views, you can edit two parts of the same program in the same buffer. To open split views, click Split Horizontally or Split Vertically on the Window menu.

### 5.1.7 Finding and Replacing Text

Along with a standard find and replace function, you can use the edit window to find and replace regular expressions. To find regular expressions, open the Find dialog box and select the checkbox for regular expressions.

### 5.1.8 Running Selected Text

There are three ways you can run selected text. First, highlight the text you want to run, then either press CTRL+R, drag and drop the selected text into the Command Input - Output window, or click "Run Selected Text" on the Run menu.

### 5.2 Using The Command Input - Output Window

The Command Input - Output window lets you input interactive commands and view the results. The Command Input - Output window can be split into two separate windows, one for input and one for output, by clicking Output Window on the Window menu.

Output will be written at the insertion point in the Command Input - Output window or the Output window, when it is a separate window. GAUSS commands cannot be executed from this window.

From the Command Input - Output window, you can run saved programs. You can view or edit the data of any variable in the active workspace with the Matrix Editor. You can also open files for editing or to debug.

The GAUSS Command Input - Output window has many of the same features that the GAUSS text editor has. You can cut and paste text. You can search the buffer of the Command Input - Output window. You can also save the contents of the Command Input - Output window to a text file.

### 5.2.1 Running Commands

The GAUSS interface allows you to run programs that consist of single commands or blocks of commands executed interactively, as well as large-scale programs that may consist of commands in one or more files. The file that is run to execute the command is the main file (the file name displayed in the Main File list).

When you run commands interactively, the actual code being processed is called the "active block." The active block is all code between the GAUSS prompt ( $\gg$ ) and the end of the current line. Thus, the active block can be one or more lines of code.

Interactive commands can be entered at the " $\gg$ " prompt in the Command Input - Output window or selected using the mouse and clicking the Run Selected Text button on the Main toolbar.

A block of code can be executed by selecting the block with the mouse and then running that block using the Run Selected Text function.

Note: The GAUSS prompt (>>) at the beginning of the selected text is ignored.
You can enter multi-line commands into the Command Input - Output window by pressing CTRL+ENTER at the end of each line. At the end of the final line in a multi-line command, press ENTER. The Command Input - Output window will automatically place a semicolon at the end of a single-line command before it is interpreted. For multi-line commands, you must enter a semicolon at the end of each line.

You can also run multi-line commands by pasting the text of a file at the GAUSS prompt, or selecting multiple lines of code from the Command Input - Output window and pressing CTRL+R.

You can repeat any of the last 20 lines entered into the command buffer by pressing CTRL+L to cycle through the last command buffer.

### 5.2.2 Running Programs in Files

You can execute the active file by clicking Run Active File on the Run menu, or by clicking the Run Currently Active File button on the Main toolbar.

You can execute the file displayed in the Main File list (the main file) by clicking Run Main file on the Run menu, or by clicking the Run Main File button on the Main toolbar.

### 5.3 Using Source View

Source View is a dockable dialog bar with two tabs that provide easy access to source files and symbols associated with your current GAUSS workspace.

### 5.3.1 Source Tab

The Source tab is a tree view that displays a list of active libraries and the source files they contain. Under each source file is a list of the symbols and procedures which they define. By using the right mouse button, you can search for symbols, open source files or view source file properties.

## Opening a Source File

To open a source file, double click the file name or right click the file and click Edit.

## Finding Commands in Source Files

To search the source files right click any file name in the source tab and click Find. In the Find dialog enter a keyword and click OK.

### 5.3.2 Symbols Tab

The Symbols tab contains a tree view of the GAUSS workspace global symbols organized by symbol type: Matrices, Arrays, Strings, String Arrays, and Structures.

## Editing or Viewing a Symbol

To edit or view a symbol, double-clicking on it or right-clicking and selecting Edit from the menu.

## Finding Symbols in Source Files

To search the source files right click any file name in the source tab and click Find. In the Find dialog enter a keyword and click OK.

### 5.4 Using the Error Output Window

The Error Output window allows errors messages to be output to a separate window, instead of the GAUSS Input - Output window. When an error occurrs, you can open to program of source file directly from the Error Output window.

To open the program or source file, press F4 or double click the error message. The file will open at the line the error occurred.

### 5.5 Using The Debugger

The debugger greatly simplifies program development. With all of the features of a dedicated debugging system, the debugger can help you to quickly identify and solve logic errors at run-time.

The debugger is integrated into the multiple document interface of GAUSS; it uses the interface tools, such as the edit windows, the Matrix Editor, and the Command Input - Output window for debugging. So while using the debugger, you still have all the features of the edit windows and Matrix Editor, along with GAUSS's suite of debugging tools.

You use the debugger to watch the program code as it runs. Prior to running the debugger, breakpoints and watch variables can be set to stop the program at points you set and provide additional data as the code is run.

### 5.5.1 Starting and Stopping the Debugger

You can start the debugger by clicking Go on the Debug menu or the Debug toolbar.
When starting the debugger, you can choose to debug the active file or to debug the main file of a program. If you are debugging a single file and already have the file open, you can use the menu or toolbar to start the debugger on the file, or simply type debug and the filename in the Command Input - Output window.

When you start the debugger, the debugger automatically highlights the first line of code to be run. Any breakpoints are shown in the left margin of the window.

You can stop the debugger at any time by clicking Stop on the Debug menu or the Debug toolbar.

### 5.5.2 Using Breakpoints

Breakpoints stop code execution where you have inserted them. Breakpoints are normally set prior to running the debugger, but can also be set or cleared during debugging by clicking the Set/Clear Breakpoint command on the Debug menu.

The debugger supports two types of breakpoints: procedure breakpoints and line number breakpoints. Procedure breakpoints pause execution when the specified procedure or function is reached. Line number breakpoints pause execution when the specified line is reached. In either case, the break occurs before any of the GAUSS code for the procedure or line is executed. The debugger also allows you to specify a certain cycle of execution for a line number or procedure where you want the execution to be paused. The cycle count is for the occurrence of the line number or procedure, not the number of times a line is to be skipped.

### 5.5.3 Setting and Clearing Breakpoints

You can set or clear a line breakpoint in the highlighted line of code by clicking Set/Clear Breakpoint on the Debug menu or by pressing the F9 key.

To set breakpoints in any part of the file not currently being executed, just click the line where you want the breakpoint to be, then click Toggle Breakpoint.

To clear breakpoints in the file, click a line of code that has a breakpoint set and then click Set/Clear Breakpoint. You can also clear all breakpoints from the active file by clicking Clear All Breakpoints.

## Using the Breakpoint Editor to Set and Clear Breakpoints

The Breakpoint Editor allows you to set or clear both line and procedure breakpoints. It also lets you specify cycles of execution for breakpoints. With the Breakpoint Editor, you can set or clear breakpoints in any program currently in your working directory.

### 5.5.4 Stepping Through a Program

GAUSS's debugger includes the ability to step into, step out of, and step over code during debugging.

Use Step Into to execute the line of code currently highlighted by the debugger.
Use Step Out to execute to the end of the current function without pause and return to the calling function.

Use Step Over to execute the line of code currently highlighted by the debugger without entering the functions that are called.

### 5.5.5 Viewing and Editing Variables

GAUSS allows you to view and edit the values of variables during debugging.

## Viewing Variable Values During Debugging

Once the debugger is started, the editor window uses floatover variable windows for viewing variable data. Floatover variable windows give a quick view of the value a variable currently holds by simply moving your mouse over the variable name in the edit window.

The floatover variable window is only intended to give a quick view of the data, so it may not show all data held by the variable. If the variable data is incomplete, the floatover variable window will display an arrow to show that there is more data. If you need to view more data, open the Matrix Editor by highlighting the variable name and pressing CTRL+E.

## Editing Variable Values During Debugging

The debugger integrates the Matrix Editor to edit values of loaded variables, or to use as a watch window to view the changing values of variables as you step through a program.

To edit a variable value, highlight the variable in the edit window, or the Command Input - Output window and then open the Matrix Editor. You can use the menu or toolbar to start the Matrix Editor, or simply type CTRL+E.

## Making a Watch Window

You can make the Matrix Editor a Watch window, allowing you to watch the changing value of a variable as the lines of the program are executed. You can activate the Watch window by clicking Set Watch on the Debug menu, or by highlighting a variable name in the debugger window and pressing CTRL+E.

You use a Watch window to see how variables change in value during debugging. Watch variables can be specified prior to running the debugger or during a debugging session.

The debugger searches for a watch variable using the following order:

1. A local variable within a currently active procedure.
2. A global variable.

A watch variable can be the name of a matrix, a scalar, a string array, or a string. For a matrix or a string array, the first element is displayed. If a matrix element is clicked, the Matrix Editor is loaded with the matrix. The matrix elements can be changed during the debugging session.

### 5.6 Customizing GAUSS

### 5.6.1 Preferences Dialog Box

The Preferences dialog box lets you specify how GAUSS operates. To open the Preferences dialog box, click Preferences... on the Configure menu. The changes you make in the Preferences dialog box remain set between sessions.

## Run Options

| Dataloop | Specifies whether or not GAUSS will translate data loops into |
| :--- | :--- |
| Translator | procedures. |

Translate Line Specifies whether or not GAUSS will preserve the line numbers of data Number
Tracking
Line Number Specifies whether or not GAUSS will preserve line numbers of a file
Tracking being compiled for the interpreter.

Sound at End Determines whether or not a sound is played at the end of the execution of Job of GAUSS code. The sound can be selected using the Select button and played using the Test button.

The default is OFF.

## Compile Options

The Compile tab contains options that let you control how GAUSS compiles a program before it is run.

| Autoload | Specifies whether the autoloader will automatically resolve references <br> in your code. If Autoload is off, you must define all symbols used in <br> your program. |
| :--- | :--- |
| Autodelete | Use Autodelete in conjunction with Autoload to control the handling of <br> references to unknown symbols. |
| GAUSS Library | Specifies whether the autoloader will use the standard GAUSS library <br> in compiling your code. |
| User Library | Specifies whether the autoloader will use the User Libraries in <br> compiling your code. |
| Declare | Specifies whether the GAUSS compiler will display declare warnings |

Warnings in the Command Input - Output window. For more information on declare warnings see Using .dec Files, Section 19.3.1.

Compiler Trace Specifies whether you would like to trace the file compilation by file opening and closing, specific lines, or whether you would like to trace by local and global symbols.

## Cmd Window

The Cmd Window tab contains options that let you control how the GAUSS Command Window operates.

| Action on | Specifies whether pressing ENTER executes the current whole line <br> Entways or only when the cursor is at the end of a line. Also, specifies <br> whether placing a semi-colon at the end of a line causes GAUSS to <br> enter multi-line mode. |
| :--- | :--- |
| Performance | Specifies whether or not output is buffered, and sets the buffer size in <br> kilobytes. |
| Cmd Prompt | Specifies whether new GAUSS prompts are inserted at the current <br> cursor location, appended to the text in the Command window, or <br> relocated on the line following the cursor. |
| Output | Specifies whether output from a GAUSS program is inserted at the <br> current cursor location, appended to the text in the Command window, <br> or written over the text following the cursor. |

## DOS Compatibility

The DOS Compatibility tab lets you control the appearance of the DOS Compatibility window.

$$
\text { Change Font } \quad \text { Specifies what font the DOS Compatibility window will use. }
$$

Tab size $\quad$ Specifies the tab size in a DOS Compatibility window

## Stay on top of Specifies whether the DOS Compatibility window will stay on top of GAUSS GAUSS.

## File

The File tab contains options that let you control how GAUSS auto-saves your work.

| Save file on <br> Execute | Specifies whether open files will <br> automatically be saved when a file is run. If the file you are running is <br> loaded, it will be saved prior to execution, regardless of how it is <br> executed (Run file, command line, main file, or active file). All open <br> editor files, including the active file, are saved before execution. |
| :--- | :--- |
|  | Note: New, unsaved documents are not automatically backed up until <br> you save them, giving them a file name. After you save the new file, it <br> will be automatically backed up with all other open files. |
| Autosave | Specifies whether you want GAUSS to automatically save your files at <br> a set interval of time. |
| Edit Window | Specifies the initial window size of opened <br> files. |

## Misc

The Misc tab contains several general options to control GAUSS.
Show Tip of the Turns on/off the Tip of the Day at startup. Day at startup

Specifies whether the Help window always stays on top of GAUSS
Keep Help when opened. of GAUSS

| Set Initial | Specifies initial position of the Help |
| :--- | :--- |
| Window Position | window. |
| Graphics Editor If you have purchased the Graphics Editor, <br> this is where you will enter or change the Graphics Editor license key.  | Licens |

### 5.6.2 Editor Properties

You can customize the formatting of your code and text by changing font colors, fonts, adding line indentations, and adding line numbering to your programs. To access these properties, on the Configure menu click Editor Properties.

## Color/Font

Color Specifies the way syntax coloring works in the editor.
Font Specifies what font the edit window will use.

## Language/Tabs

Auto Indentation Specifies how the autoindenter will indent your code. Style

Tabs Specifies how many spaces a tab has.
Language Specifies what syntax the GAUSS editor will recognize for syntax coloring.

Fixup Text Case Specifies whether the editor will automatically change the case of While Typing Language Keywords

## Misc

Smooth Enables or disables smooth scrolling when the window is scrolled Scrolling up/down by one line or left/right by one character.

Show Left Enables or disables the editor's margin. The margin is used for
Margin showing breakpoints, bookmarks, or line numbers.

Line Tooltips Shows the first line number on screen as a tooltip as you scroll up and on Scroll down the file.
Allow Drag Enables or disables drag and drop functionality. and Drop
Allow Column Lets you select and manipulate columns of text.Selection
Confine Caret to Text
Tells the GAUSS editor to interpret carets as text only rather than as ..... Text substitution symbols or text.
Color Syntax Toggles on or off color syntax highlighting.
Highlighting
Show Horizontal Toggles on or off the horizontal scrollbar. Scrollbar
Show Vertical Toggles on or off the vertical scrollbar.Scrollbar
Allow Vertical Toggles on or off the ability to split editor panes vertically. Splitting
Allow Toggles on or off the ability to split editor panes horizontally.HorizontalSplitting
Line Numbering Specifies the style and starting digit for line numbering.
Max Undoable Sets the number of actions that you can undo.
Actions
5.7 Using GAUSS Keyboard Assignments

### 5.7.1 Cursor Movement Keys

UP ARROWDOWN ARROWLEFT ARROWUp one lineDown one lineLeft one character

| RIGHT ARROW | Right one character |
| :--- | :--- |
| CTRL+LEFT ARROW | Left one word |
| CTRL+RIGHT ARROW | Right one word |
| HOME | Beginning of line |
| END | End of line |
| PAGE UP | Next screen up |
| PAGE DOWN | Next screen down |
| CTRL+PAGE UP | Scroll window right |
| CTRL+PAGE DOWN | Scroll window left |
| CTRL+HOME | Beginning of document |
| CTRL+END | End of document |

### 5.7.2 Edit Keys

BACKSPACE Delete character to left of cursor, or delete selected text
DEL Delete character to right of cursor, or delete selected text
CTRL+INS or CTRL+C Copy selected text to Windows clipboard
SHIFT + DEL or CTRL+X Delete selected text and place it onto Windows clipboard
SHIFT + INS or CTRL+V Paste text from Windows clipboard at the cursor position
CTRL+Z
Undo last editing action

### 5.7.3 Text Selection Keys

| SHIFT+UP ARROW | Select one line of text up |
| :--- | :--- |
| SHIFT+DOWN ARROW | Select one line of text down |
| SHIFT+LEFT ARROW | Select one character to the left |
| SHIFT+RIGHT ARROW | Select one character to the right |
| SHIFT+CTRL+LEFT ARROW | Select one word to the left |
| SHIFT+CTRL+RIGHT ARROW | Select one word to the right |
| SHIFT+HOME | Select to beginning of the line |
| SHIFT+END | Select to end of the line |
| SHIFT+PAGE UP | Select up one screen |
| SHIFT+PAGE DOWN | Select down one screen |
| SHIFT+CTRL+HOME | Select text to end of document |

### 5.7.4 Command Keys

| CTRL+A | Redo |
| :--- | :--- |
| CTRL+C | Copy selection to Windows clipboard |
| CTRL+D | Open Debug window |
| CTRL+E | Open Matrix Editor |
| CTRL+F | Find/Replace text |
| CTRL+G | Go to specified line number |
| CTRL+I | Insert GAUSS prompt |


| CTRL+L | Insert last |
| :--- | :--- |
| CTRL+N | Make next window active |
| CTRL+O | Open Output window |
| CTRL+P | Print current window, or selected text |
| CTRL+Q | Exit GAUSS |
| CTRL+R | Run selected text |
| CTRL+S | Save window to file |
| CTRL+W | Open Command window |
| CTRL+V | Paste contents of Windows clipboard |
| CTRL+X | Cut selection to Windows clipboard |
| CTRL+Z | Undo |
| 5.7.5 Function Keys |  |
| F1 | Open GAUSS Help system or context-sensitive Help |
| F2 | Go to next bookmark |
| F3 | Find again |
| F4 | Go to next search item in Source Browser |
| F5 | Run Main File |
| F6 | Run Active File |
| F7 | Edit Main File |
| F8 | Step Into |
| F9 | Set/Clear breakpoint |




## Matrix Editor

## 6

### 6.1 Using the Matrix Editor

The Matrix Editor lets you view and edit matrix data in your current workspace. You can open the Matrix Editor from either the Command Input - Output window or a GAUSS edit window by highlighting a matrix variable name and typing CTRL+E. You can view multiple matrices at the same time by opening more than one Matrix Editor.

### 6.1.1 Editing Matrices

The Matrix Editor will allow you to format matrices in decimal, scientific, Hexadecimal, or as text characters.

Just like a spreadsheet, when using the Matrix Editor, you can use your keyboard's arrow keys to quickly move between matrix positions. To edit a scalar value, select a cell and press Enter. You can use the Home and End keys to move to the beginning or end of a scalar. When finished editing, press Enter again.

| \#x-matrix (50x10) |  |  |  |  |  |  | - $\square$ | $\times$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Matrix Format Edit View |  |  |  |  |  |  |  |  |
| 123 e ff a | [..] 国 (\%) 宿 |  |  |  |  |  |  | ^ |
|  | 1 | 2 | 3 | 4 | 5 | 6 |  |  |
| 1 | -0.148443 | -0.553305 | -0.508804 | -0.552587 | 0.526161 | -0.251634 |  |  |
| 2 | -0.437972 | -0.938730 | 0.981923 | 0.472761 | 0.860515 | -0.063889 |  |  |
| 3 | 1.353409 | 0.627728 | 1.189957 | 0.508362 | -1.498672 | -0.710302 |  |  |
| 4 | 0.967591 | -0.113163 | -0.097851 | 0.521970 | -1.132649 | 0.023005 |  |  |
| 5 | 0.973790 | -0.699632 | -0.543146 | 1.287314 | 1.441660 | -0.167420 |  |  |
| 6 | -0.545715 | 1.273839 | -0.697932 | -0.220956 | 0.810306 | 2.081371 |  |  |
| 7 | 1.086625 | 0.367896 | -0.468893 | 1.380549 | -0.587850 | 0.663834 |  |  |
| 8 | 0.501997 | -1.161872 | -0.790847 | -0.963507 | -1.433578 | 1.624267 |  |  |
| 9 | 0.621958 | -0.922859 | 0.333411 | 0.421978 | 1.621670 | 0.900518 |  |  |
| 10 | 1.764253 | -0.938587 | 0.177002 | 0.756723 | -0.193889 | -0.541515 |  |  |
| 11 | -0.080564 | -1.906603 | 0.327855 | -1.023495 | 0.085362 | 0.001252 |  |  |
| 12 | 0.059547 | 1.075533 | -0.424191 | 1.648687 | -1.893974 | -1.204153 |  |  |
| 13 | -0.206388 | 1.271439 | 0.527988 | 0.643007 | -1.307583 | 1.207785 |  |  |
| 14 | -1.319452 | 1.172313 | 1.576035 | -0.288744 | -0.485703 | 1.478021 |  |  |
| 15 | 2.458355 | -0.663845 | 1.214584 | -1.427925 | 0.159592 | -1.876784 |  |  |
| 16 | 0.140758 | -0.773888 | -0.323972 | 0.668909 | 1.281200 | -0.783832 |  |  |
| 17 | -0.086262 | 0.764467 | 0.839743 | $-0.448715$ | 0.908994 | -0.430262 |  |  |
| 18 | -0.688029 | -0.661764 | -0.982795 | 0.024450 | -0.805820 | 0.647185 |  |  |
| 19 | -0.871597 | -0.607500 | -0.278373 | 1.029881 | 0.603642 | 0.242880 |  |  |
| 20 | 0.221217 | -0.639253 | -0.700970 | -1.922023 | -0.148534 | 0.638206 |  |  |
|  |  |  |  |  |  | R 1, C 1 | Real | $\checkmark$ |
| $\leqslant$ |  |  |  | ] |  |  | , |  |

Figure 6.1: Matrix Editor

### 6.1.2 Viewing Variables

All variables are treated as matrices in GAUSS. A scalar is simply a $1 \times 1$ matrix. A vector is a $(\mathrm{N} \times 1)$ or $(1 \times \mathrm{N})$ matrix. So you can use the Matrix Editor to view and monitor the value of any variable. You can update the value of a variable at any time by using the Reload function. When using the Matrix Editor to view, edit or monitor smaller matrices, you can minimize space it occupies on the screen by selecting Minimal View from the View menu.

By using the Auto-reload function, GAUSS will automatically update the values of variables in the Matrix Editor. Using Auto-reload you can create a watch window.

## Setting Watch Variables

Watch Variables allow you to see how variables change in value while debugging a program. A watch variable can be the name of a matrix, a scalar, an array, a string array, or a string.

The debugger searches for a watch variable in the following order:

- a local variable within a currently active procedure
- a global variable


### 6.1.3 Matrix Editor Menu Bar

## Matrix Menu

The Matrix menu lets you control the data of the Matrix in the Matrix Editor as an entire set.

Load $\begin{aligned} & \text { Clears any existing grid and loads any named matrix from the GAUSS } \\ & \text { workspace to the grid. }\end{aligned}$ l
Reload Reloads the existing matrix with the name shown on the Title bar.

Auto-Reload Automatically updates the data shown in the Matrix Editor, creating a watch window.

Save Saves the grid as a matrix in the GAUSS workspace. If a matrix of the same name already exists in the workspace, it is overwritten.

## Format Menu

The Format menu lets you control the way the data is presented in the Matrix Editor.

| Decimal | Display selected elements as decimal numbers. |
| :--- | :--- |
| Scientific | Display selected elements using scientific notation. |
| Hexadecimal | Display selected elements as hexadecimal numbers. |
| Character | Display selected elements as character data. |
| Precision... | Specify precision of selected elements. |

## Edit Menu

The Edit menu gives you tools to control the data in the Matrix Editor.

| Clear All | Clears the grid of all values but keep the row and column order. |
| :--- | :--- |
| Preferences | Sets several matrix options, including the number of digits to the right <br> of the decimal point, cell height and width, and whether pressing the <br> Enter key moves the cursor down or over one cell. These options, along <br> with screen position and window state, are saved between sessions. |

## View Menu

The View menu lets you control the Matrix Editor window. The View menu also lets you control your view of imaginary numbers.

| Dimension | Specifies which two-dimensional subarray of an N-dimensional array <br> to display by indexing the N-2 leading dimensions of the array. |
| :--- | :--- |
| Real Parts | Specifies that you want the real parts of imaginary numbers to be <br> displayed in the Matrix Editor. |
| Imaginary Parts | Specifies that you want the imaginary parts of numbers to be displayed <br> in the Matrix Editor. |
| Minimal View | Minimizes the amount of screen space occupied by the Matrix Editor. <br> This is especially useful for creating watch windows for single |
| variables. |  |$\quad$| Forces the Matrix Editor window to remain visible on the screen even |
| :--- |
| when the interface focus has shifted to another window. |



## Library Tool

## 7

### 7.1 Using the Library Tool

The Library Tool lets you quickly manage your libraries. You can add and remove libraries and you can add and remove files within the libraries.

### 7.1.1 Managing Libraries

Using the New Library button, you can create a new library for organizing your code. You can remove a library by selecting the Delete Library button.

### 7.1.2 Managing the Library Index

To add absolute path names to the library index, use the Add Paths button. To only use file names for searching libraries, use the Strip Paths button. Use Rebuild to recompile all the files used in the library, and rebuild the library index file. Use the Revert to Original button to revert to the configuration the library was in when the Library Tool was opened.


Figure 7.1: Library Tool

### 7.1.3 Managing Library Files

You can add files to a library with the Add button. You can remove files from a library with the Remove button. After changing source files referred to in a library, select the files in the file list and update the library index with the Update button. To remove multiple files from a library, select the files in the file selection window, and use the Clear Selection button.

For more information about libraries, see Libraries, Chapter 19.


## GAUSS Source Browser

The GAUSS Source Browser lets users quickly find, view, and if necessary, modify source code. Both the TGAUSS and GAUSS Source Browsers can be used to search for external symbols in active libraries. The GAUSS Source Browser can also be used to search for symbols in any directory or source file.

### 8.1 Using the Source Browser in TGAUSS

To start the Source Browser in TGAUSS, type BROWSE followed by a symbol name. When the Source Browser is active, the prompt displays Browse: GAUSS searches through all active libraries for the file in which the symbol is defined. If found, the file containing the source code is opened in the default editor.

Wildcard (*) searches can also be used. When using wildcard searches, each symbol that the string matches will be displayed on-screen in a numbered list. To select a specific command to view in the default editor, select the number from the list.

The Source Browser will remain active until you type CTRL-C to return to the GAUSS prompt.

### 8.2 Using the Source Browser in GAUSS

To open the Source Browser in GAUSS, from the Tools menu select Source Browser.
Using the Source Browser you can search a file for a specified symbol or search across all the source files in a directory. Using a comma separated list of files and directories in the Look in: list, you can search multiple locations in a single search. When searching for symbols, you can use wildcards $(*)$ to further modify the scope of the search.

Note: The Source Browser does not search recursively through sub-folders. To search sub-folders during a search, add the sub-folder names to the Look in: list.

Once the search is complete, the Source Browser lists where the specified symbol was found. The Filename column of the Results List shows the file in which the symbol was found. The Line column shows the line number where symbol was found and the Line Description column shows the text of the line where the symbol was found.

Search locations typed into the Look in: text box will persist between Source Browser sessions.

## Pattern: Defines search pattern.

Look in: Limits the scope of the search to specific files or directories. Using a comma separated list, searches multiple files and directories in a single search.

Match Case Makes search case-sensitive.
Match whole Limits search to entire words. word

Stay on top Keeps the Source Browser on top even when another window is active.
F4 Opens New Sets F4 to open new window.
Window
Browse Lets you limit the scope of the search to specific files or directories.
Search Initiates search.
Results List Lists occurrences of selected symbol in specified files or directories.


Figure 8.1: Source Browser


#### Abstract

Status Lists how many occurrences there were, and how many files the symbol occurred in.

Close Closes the Source Browser.


### 8.2.1 Opening Files From the Source Browser

Double-click the file name to open a file in its own editor window. When opened, the cursor is placed at the beginning of the line selected in the Results List. By double-clicking different files in the Source Browser, you can open each file in its own separate editor window.

Use the F4 key to quickly view or edit the next file in the Results List using the active editor window. Using the F4 key opens the file in the active editor window and places the cursor at the beginning of the line in which the symbol was found. The F4 key uses the active editor window to display the source file; it will not open an editor window to display files. You can use the F4 key from either the Source Browser or from the active editor window to move to the next occurrence of the symbol shown in the Results List.

Use SHIFT+F4 to quickly view or edit the previous file in the Results List using the active editor window. Using the F4 key opens the file in the active editor window and places the cursor at the beginning of the line in which the symbol was found.

### 8.2.2 Source Browser Keyboard Controls

UP ARROW Moves to the previous occurrence in the Results List.
DOWN ARROW Moves to the next occurrence in the Results List.
HOME Moves to the first occurrence in the Results List.
END Moves to the last occurrence in the Results List.
F4 Shows the next occurrence in the active editor window.
SHIFT+F4 Shows the previous occurrence in the active editor window.
TAB Moves to next field.
ENTER Starts Search.

## GAUSS Help

### 9.1 Help Menu

From the Help menu, you can directly access the online User Guide, Keyboard Assignments list, and Language Reference. Pressing F1 also accesses the Help system, displaying either the Introduction to the User Guide or, if an object has focus and help can be directly accessed, help for that object.

### 9.2 Context-Sensitive Help

GAUSS integrates a context-sensitive Help system to help you use the GAUSS environment and the GAUSS language. Context-sensitive means that Help for the object with focus is displayed without navigating through the Help system. For example, to display Help on a keyword in the GAUSS language in a GAUSS edit window or the Command Input - Output window, place the insertion point on the keyword and press F1.

Several areas of the GAUSS interface are context-sensitive, including:

- GAUSS windows
- Toolbar buttons
- GAUSS menus
- The GAUSS language

For intrinsic commands and functions, the GAUSS Language Reference for the command is displayed. For other external procedures in active libraries, a window displays a source code file, allowing you to scroll to the desired symbol.

### 9.3 SHIFT+F1 Support

If you press SHIFT + F1 or click on the Help toolbar button (an arrow with a question mark), the pointer changes to a Help pointer (arrow + ?). Click on an object to display the Help system or, if available, context-sensitive Help for that object.

### 9.4 CTRL+F1 Support

You can search through all active libraries for any global symbol by placing the cursor on the symbol name and pressing CTRL+F1.

GAUSS searches through all active libraries for the file that the symbol is defined in. If found, the file containing the source code is opened in an edit window. If the file contains the code string "**> symbol_name" (without quotes) at the beginning of a line of commented code, the cursor will be placed at the beginning of that line. If the string is not found in the file, the cursor will be placed at the beginning of the file.

To properly implement this functionality in your own source code, place
**> symbol_name
at the beginning of a line in a comment block.

### 9.5 ToolTips

A ToolTip is a small label that is displayed when the mouse pointer is held over a GAUSS button. The ToolTip will give a brief description of the button's function.

### 9.6 Other Help

GAUSS includes full online versions of the GAUSS Language Reference and GAUSS User Guide in PDF format. These manuals are available for download; contact Aptech Systems for download instructions.

The Gaussians mail list is an e-mail list providing users of GAUSS an easy way to reach other GAUSS users. Gaussians provides a forum for information exchange, tips and experiences using GAUSS. For more information about the Gaussians mail list, see the Resource Library page at http://www. aptech. com. You can also e-mail support@aptech. com when you have current Premier Support.


## Language Fundamentals

## 10

GAUSS is a compiled language. GAUSS is also an interpreter. A compiled language, because GAUSS scans the entire program once and translates it into a binary code before it starts to execute the program. An interpreter, because the binary code is not the native code of the CPU. When GAUSS executes the binary pseudocode it must "interpret" each instruction for the computer.

How can GAUSS be so fast if it is an interpreter? Two reasons. First, GAUSS has a fast interpreter, and the binary compiled code is compact and efficient. Second, and most significantly, GAUSS is a matrix language. It is designed to tackle problems that can be solved in terms of matrix or vector equations. Much of the time lost in interpreting the pseudocode is made up in the matrix or vector operations.

This chapter will enable you to understand the distinction between "compile time" and "execution time", two very different stages in the life of a GAUSS program.

### 10.1 Expressions

An expression is a matrix, string, constant, function reference, procedure reference, or any combination of these joined by operators. An expression returns a result that can be assigned to a

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variable with the assignment operator ' $=$ '.

### 10.2 Statements

A statement is a complete expression or command. Statements end with a semicolon.

$$
\mathrm{y}=\mathrm{x} * 3 ;
$$

If an expression has no assignment operator (=), it will be assumed to be an implicit print statement:

```
print x*3;
```

or

$$
x * 3
$$

Here is an example of a statement that is a command rather than an expression:
output on;

Commands cannot be used as a part of an expression.
There can be multiple statements on the same line as long as each statement is terminated with a semicolon.

### 10.2.1 Executable Statements

Executable statements are statements that can be "executed" over and over during the execution phase of a GAUSS program (execution time). As an executable statement is compiled, binary code is added to the program being compiled at the current location of the instruction pointer. This binary code will be executed whenever the interpreter passes through this section of the program. If this code is in a loop, it will be executed each iteration of the loop.

Here are some examples of executable statements:

```
y = 34.25;
print y;
x = 1372940 3;
```


### 10.2.2 Nonexecutable Statements

Nonexecutable statements are statements that have an effect only when the program is compiled (compile time). They generate no executable code at the current location of the instruction pointer.

Here are two examples:

```
declare matrix x = 1 2 3 4 ;
external matrix ybar;
```

Procedure definitions are nonexecutable. They do not generate executable code at the current location of the instruction pointer.

Here is an example:

```
zed = rndn(3,3);
proc sqrtinv(x);
    local y;
    y = sqrt(x);
    retp(y+inv(x));
endp;
zsi = sqrtinv(zed);
```

There are two executable statements in the example above: the first line and the last line. In the binary code that is generated, the last line will follow immediately after the first line. The last line is the call to the procedure. This generates executable code. The procedure definition generates no code at the current location of the instruction pointer.

There is code generated in the procedure definition, but it is isolated from the rest of the program. It is executable only within the scope of the procedure and can be reached only by calling the procedure.

### 10.3 Programs

A program is any set of statements that are run together at one time. There are two sections within a program.

### 10.3.1 Main Section

The main section of the program is all of the code that is compiled together WITHOUT relying on the autoloader. This means code that is in the main file or is included in the compilation of the main file with an \#include statement. ALL executable code should be in the main section.

There must always be a main section even if it consists only of a call to the one and only procedure called in the program.

### 10.3.2 Secondary Sections

Secondary sections of the program are files that are neither run directly nor included in the main section with \#include statements.

The secondary sections of the program can be left to the autoloader to locate and compile when they are needed. Secondary sections must have only procedure definitions and other nonexecutable statements.
\#include statements are allowed in secondary sections as long as the file being included does not violate the above criteria.

Here is an example of a secondary section:

```
declare matrix tol = 1.0e-15;
proc feq(a,b);
    retp(abs(a-b) <= tol);
endp;
```


### 10.4 Compiler Directives

Compiler directives are commands that tell GAUSS how to process a program during compilation. Directives determine what the final compiled form of a program will be. They can affect part or all of the source code for a program. Directives are not executable statements and have no effect at run-time. They do not take a semicolon at the end of the line.

The \#include statement mentioned earlier is actually a compiler directive. It tells GAUSS to compile code from a separate file as though it were actually part of the file being compiled. This code is compiled in at the position of the \#include statement.

Here are the compiler directives available in GAUSS:
\#define Define a case-insensitive text-replacement or flag variable.

| \#definecs | Define a case-sensitive text-replacement or flag variable. |
| :--- | :--- |
| \#undef | Undefine a text-replacement or flag variable. |
| \#ifdef | Compile code block if a variable has been \#define'd. |
| \#ifndef | Compile code block if a variable has not been \#define'd. |
| \#iflight | Compile code block if running GAUSS Light. |
| \#ifdos | Compile code block if running DOS. |
| \#ifos2win | Compile code block if running OS/2 or Windows. |
| \#ifunix | Else clause for \#if-\#else-\#endif code block. <br> \#else |
| \#endif | Include code from another file in program. |
| \#include | Compile program with line number and file name records. |
| \#lineson | Insert source file name record at this point (currently used when <br> \#loing data loop translation). |
| \#linesoff | Insert source file line number record at this point (currently used <br> \#hen doing data loop translation). |
| \#srcfile |  |

The \#define statement can be used to define abstract constants. For example, you could define the default graphics page size as:

```
#define hpage
    9.0
#define vpage 6.855
```

and then write your program using hpage and vpage. GAUSS will replace them with 9.0 and 6.855 when it compiles the program. This makes a program much more readable.

The \#ifdef-\#else-\#endif directives allow you to conditionally compile sections of a program, depending on whether a particular flag variable has been \#define'd. For example:

```
#ifdef log_10
    y = log(x);
#else
    y = ln(x);
#endif
```

This allows the same program to calculate answers using different base logarithms, depending on whether or not the program has a \#define $\log _{\mathbf{l}} \mathbf{1 0}$ statement at the top.
\#undef allows you to undefine text-replacement or flag variables so they no longer affect a program, or so you can \#define them again with a different value for a different section of the program. If you use \#definecs to define a case-sensitive variable, you must use the right case when \#undef'ing it.

With \#lineson, \#linesoff, \#srcline, and \#srcfile you can include line number and file name records in your compiled code, so that run-time errors will be easier to track down. \#srcline and \#srcfile are currently used by GAUSS when doing data loop translation.

For more information on line number tracking, see Debugging, Section 20.3 and see Debugging Data Loops, Section 23.3. See also \#lineson in the GAUSS Language Reference.

The syntax for \#srcfile and \#srcline is different than for the other directives that take arguments. Typically, directives do not take arguments in parentheses; that is, they look like keywords:

```
#define red 4
```

\#srcfile and \#srcline, however, do take their arguments in parentheses (like procedures):

```
#srcline(12)
```

This allows you to place \#srcline statements in the middle of GAUSS commands, so that line numbers are reported precisely as you want them. For example:

```
#srcline(1) print "Here is a multi-line "
#srcline(2) "sentence--if it contains a run-time error, "
#srcline(3) "you will know exactly "
#srcline(4) "which part of the sentence has the problem.";
```

The argument supplied to \#srcfile does not need quotes:

```
#srcfile(/gauss/test.e)
```


### 10.5 Procedures

A procedure allows you to define a new function which you can then use as if it were an intrinsic function. It is called in the same way as an intrinsic function.

$$
y=\operatorname{myproc}(a, b, c) ;
$$

Procedures are isolated from the rest of your program and cannot be entered except by calling them. Some or all of the variables inside a procedure can be local variables. local variables exist only when the procedure is actually executing and then disappear. Local variables cannot get mixed up with other variables of the same name in your main program or in other procedures.

For details on defining and calling procedures, see Procedures and Keywords, chapter 12.

### 10.6 Data Types

There are four basic data types in GAUSS, matrices, N -dimensional arrays, strings and string arrays. It is not necessary to declare the type of a variable, but it is good programming practice to respect the types of variables whenever possible. The data type and size can change in the course of a program.

The declare statement, used for compile-time initialization, enforces type checking.
Short strings of up to 8 bytes can be entered into elements of matrices, to form character matrices (For details, see Character Matrices, Section 10.6.7).

### 10.6.1 Constants

The following constant types are supported:

## Decimal

Decimal constants can be either integer or floating point values:
$1.34 \mathrm{e}-10$
1.34 e 123
$-1.34 \mathrm{e}+10$
$-1.34 d-10$
1.34 d 10
$1.34 \mathrm{~d}+10$
123.456789345

Up to 18 consecutive digits before and after the decimal point(depending on the platform) are significant, but the final result will be rounded to double precision if necessary. The range is the same as for matrices (For details, see Matrices, Section 10.6.2.

## String

String constants are enclosed in quotation marks:
"This is a string."

## Hexadecimal Integer

Hexadecimal integer constants are prefixed with $\mathbf{0} \mathbf{x}$ :

0x0ab53def2

## Hexadecimal Floating Point

Hexadecimal floating point constants are prefixed with $\mathbf{Q} \mathbf{v}$. This allows you to input a double precision value exactly as you want using 16 hexadecimal digits. The highest order byte is to the left:

0vfff8000000000000

### 10.6.2 Matrices

Matrices are 2-dimensional arrays of double precision numbers. All matrices are implicitly complex, although if it consists only of zeros, the imaginary part may take up no space. Matrices are stored in row major order. A $2 \times 3$ real matrix will be stored in the following way from the lowest addressed element to the highest addressed element:

$$
[1,1] \quad[1,2] \quad[1,3] \quad[2,1] \quad[2,2] \quad[2,3]
$$

A $2 \times 3$ complex matrix will be stored in the following way from the lowest addressed element to the highest addressed element:

| (real part) | $[1,1]$ | $[1,2]$ | $[1,3]$ | $[2,1]$ | $[2,2]$ | $[2,3]$ |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| (imaginary part) | $[1,1]$ | $[1,2]$ | $[1,3]$ | $[2,1]$ | $[2,2]$ | $[2,3]$ |

Conversion between complex and real matrices occurs automatically and is transparent to the user in most cases. Functions are provided to provide explicit control when necessary.

All elements of a GAUSS matrix are stored in double precision floating point format, and each takes up 8 bytes of memory. This is the IEEE 754 format:

| Bytes | Data Type | Significant <br> Digits | Range |
| :---: | :---: | :---: | :---: |
| 8 | floating point | $15-16$ | $4.19 \times 10^{-307} \leq\|X\| \leq 1.67 \times 10^{+308}$ |

Matrices with only one element ( $1 \times 1$ matrices) are referred to as scalars, and matrices with only one row or column $(1 \times \mathrm{N}$ or $\mathrm{N} \times 1$ matrices) are referred to as vectors.

Any matrix or vector can be indexed with two indices. Vectors can be indexed with one index. Scalars can be indexed with one or two indices also, because scalars, vectors, and matrices are the same data type to GAUSS

The majority of functions and operators in GAUSS take matrices as arguments. The following functions and operators are used for defining, saving, and loading matrices:

| [ ] | Indexing matrices. |
| :--- | :--- |
| = | Assignment operator. |
| $\sim$ | Vertical concatenation. |
| con | Horizontal concatenation. |
| cons | Numeric input from keyboard. |
| declare | Character input from keyboard. |
| let | Compile-time matrix or string initialization. |
| load | Load matrix (same as loadm). |
| readr | Read from a GAUSS matrix or data set file. |
| save | Save matrices, procedures and strings to disk. |
| saved | Convert a matrix to a GAUSS data set. |
| stof | Convert string to matrix. |
| submat | Extract a submatrix. |
| writer | Write data to a GAUSS data set. |

Following are some examples of matrix definition statements.
An assignment statement followed by data enclosed in braces is an implicit let statement. Only constants are allowed in let statements; operators are illegal. When braces are used in let statements, commas are used to separate rows. The statement

$$
\text { let } x=123,456,789 \text {; }
$$

or

$$
x=123,456,789 \text {; }
$$

will result in

$$
x=\begin{array}{lll}
1 & 2 & 3 \\
4 & 5 & 6 \\
7 & 8 & 9
\end{array}
$$

The statement

$$
\text { let } \mathrm{x}[3,3]=123456789 \text {; }
$$

will result in

$$
x=\begin{array}{lll}
1 & 2 & 3 \\
4 & 5 & 6 \\
7 & 8 & 9
\end{array}
$$

The statement
let $\mathrm{x}[3,3]=1$;
will result in

$$
x=\begin{array}{lll}
1 & 1 & 1 \\
1 & 1 & 1 \\
1 & 1 & 1
\end{array}
$$

The statement
let $\mathrm{x}[3,3]$;
will result in

$$
x=\begin{array}{lll}
0 & 0 & 0 \\
0 & 0 & 0 \\
0 & 0 & 0
\end{array}
$$

The statement

```
let x = 1 2 3 4 5 6 7 8 9;
```

will result in

$$
\begin{array}{r}
1 \\
2 \\
3 \\
4 \\
5 \\
5 \\
6 \\
7 \\
8 \\
9
\end{array}
$$

Complex constants can be entered in a let statement. In the following example, the + or - is not a mathematical operator, but connects the two parts of a complex number. There should be no spaces between the + or - and the parts of the number. If a number has both real and imaginary parts, the trailing ' $i$ ' is not necessary. If a number has no real part, you can indicate that it is imaginary by appending the ' i '. The statement

$$
\text { let } x[2,2]=1+2 i \quad 3-456 i ;
$$

will result in

$$
x=\begin{array}{cl}
1+2 i & 3-4 i \\
5 & 0+6 i
\end{array}
$$

Complex constants can also be used with the declare, con and stof statements.
An "empty matrix" is a matrix that contains no data. Empty matrices are created with the let statement and braces:

$$
\mathrm{x}=\{ \} ;
$$

Empty matrices are supported by several functions, including rows and cols and the concatenation ( $\sim, \mid$ ) operators.

```
x = {};
hsec0 = hsec;
do until hsec-hsec0 > 6000;
    x = x ~ data_in(hsec-hsec0);
endo;
```

You can test whether a matrix is empty by entering rows $(x), \operatorname{cols}(x)$ and $\operatorname{scalerr}(x)$. If the matrix is empty rows and cols will return a 0 , and scalerr will return 65535.

The $\sim$ is the horizontal concatenation operator and the $\mid$ is the vertical concatenation operator. The statement

$$
y=1 \sim 2 \mid 3 \sim 4 ;
$$

will be evaluated as

$$
y=(1 \sim 2) \mid(3 \sim 4)
$$

and will result in a $2 \times 2$ matrix because horizontal concatenation has precedence over vertical concatenation:

12
34

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The statement

$$
y=1+1 \sim 2 * 2 \mid 3-2 \sim 6 / 2 ;
$$

will be evaluated as

$$
y=((1+1) \sim(2 * 2)) \mid((3-2) \sim(6 / 2))
$$

and will result in a $2 \times 2$ matrix because the arithmetic operators have precedence over concatenation:

24
13

For more information, see Operator Precedence, Section 11.7.
The let command is used to initialize matrices with constant values:

$$
\text { let } x[2,2]=1234 \text {; }
$$

Unlike the concatenation operators, it cannot be used to define matrices in terms of expressions such as

$$
y=x 1-x 2 \sim x 2 \mid x 3 * 3 \sim x 4 ;
$$

The statement

$$
y=x[1: 3,5: 8] ;
$$

will put the intersection of the first three rows and the fifth through eighth columns of $\mathbf{x}$ into the matrix $y$.

The statement

$$
\mathrm{y}=\mathrm{x}\left[\begin{array}{lllll}
1 & 3 & 1,5 & 5 & 9
\end{array}\right] ;
$$

will create a $3 \times 3$ matrix $\mathbf{y}$ with the intersection of the specified rows and columns pulled from $\mathbf{x}$ (in the indicated order).

The following code

```
let r = 1 3 1; let c= 5 5 9; y = x[r,c];
```

will have the same effect as the previous example, but is more general.
The statement

$$
y[2,4]=3 ;
$$

will set the 2,4 element of the existing matrix $\mathbf{y}$ to 3 . This statement is illegal if $\mathbf{y}$ does not have at least 2 rows and 4 columns.

The statement

$$
x=\operatorname{con}(3,2) ;
$$

will cause the following prompt to be printed in the window:

$$
\text { - }(1,1)
$$

indicating that the user should enter the [1,1] element of the matrix. Entering a number and then pressing ENTER will cause a prompt for the next element of the matrix to appear. Pressing ? will display a help screen, and pressing $\mathbf{x}$ will exit.

The statement

```
load x[] = b:mydata.asc
```

will load data contained in an ASCII file into an $N \times 1$ vector $\mathbf{x}$. (Use rows ( $\mathbf{x}$ ) to find out how many numbers were loaded, and use reshape ( $x, N, K$ ) to reshape it to an $N \times K$ matrix).

The statement
load x;
will load the matrix x . fmt from disk (using the current load path) into the matrix $\mathbf{x}$ in memory.
The statement

```
open d1 = dat1;
x = readr(d1,100);
```

will read the first 100 rows of the GAUSS data set dat1. dat.

### 10.6.3 Sparse Matrices

Many GAUSS operators and commands support the sparse matrix data type. You may use any of the following commands to create a sparse matrix:

| denseToSp | Converts a dense matrix to a sparse matrix. |
| :--- | :--- |
| denseToSpRE | Converts a dense matrix to a sparse matrix, using a relative epsilon. |


| packedToSp | Creates a sparse matrix from a packed matrix of non-zero values and <br> row and column indices. |
| :--- | :--- |
| spCreate | Creates a sparse matrix from vectors of non-zero values, row <br> indices, and column indices. |
| spEye | Creates a sparse identity matrix. |
| spOnes | Generates a sparse matrix containing only ones and zeros |
| spZeros | Creates a sparse matrix containing no non-zero values. |

See Sparse Matrices, Chapter 13, for more information.

### 10.6.4 N-dimensional Arrays

Many GAUSS commands support arrays of N dimensions. The following commands may be used to create and manipulate an N -dimensional array:

| aconcat | Concatenate conformable matrices and arrays in a user-specified <br> dimension. |
| :--- | :--- |
| aeye | Create an N-dimensional array in which the planes described by the <br> two trailing dimensions of the array are equal to the identity. |
| areshape | Reshape a scalar, matrix, or array into an array of user-specified size. |
| arrayalloc | Create an N-dimensional array with unspecified contents. |
| arrayinit | Create an N-dimensional array with a specified fill value. |
| mattoarray | Convert a matrix to a type array. |

### 10.6.5 Strings

Strings can be used to store the names of files to be opened, messages to be printed, entire files, or whatever else you might need. Any byte value is legal in a string from 0-255. The buffer where a string is stored always contains a terminating byte of ASCII 0 . This allows passing strings as arguments to C functions through the Foreign Language Interface.

Here is a partial list of the functions for manipulating strings:

| \$+ | Combine two strings into one long string. |
| :---: | :---: |
| - | Interpret following name as a variable, not a literal. |
| chrs | Convert vector of ASCII codes to character string. |
| dttostr | Convert a matrix containing dates in DT scalar format to a string array. |
| ftocv | Character representation of numbers in $\mathrm{N} \times \mathrm{K}$ matrix. |
| ftos | Character representation of numbers in $1 \times 1$ matrix. |
| ftostrC | Convert a matrix to a string array using a C language format specification. |
| getf | Load ASCII or binary file into string. |
| indcv | Find index of element in character vector. |
| lower | Convert to lowercase. |
| stof | Convert string to floating point. |
| strindx | Find index of a string within a second string. |
| strlen | Length of a string. |
| strsect | Extract substring of string. |
| strsplit | Split an $\mathrm{N} \times 1$ string vector into an $\mathrm{N} \times \mathrm{K}$ string array of the individual tokens. |


| strsplitPad | Split a string vector into a string array of the individual tokens. Pads <br> on the right with null strings. |
| :--- | :--- |
| strtodt | Convert a string array of dates to a matrix in DT scalar format. |
| strtof | Convert a string array to a numeric matrix. |
| strtofcplx | Convert a string array to a complex numeric matrix. |
| upper | Convert to uppercase. |
| vals | Convert from string to numeric vector of ASCII codes. |

Strings can be created like this:

```
x = "example string";
```

```
x = cons; /* keyboard input */
```

or

```
x = getf("myfile",0); /* read a file into a string */
```

They can be printed like this:

```
print x;
```

A character matrix must have a '\$' prefixed to it in a print statement:

```
print $x;
```

A string can be saved to disk with the save command in a file with a .fst extension and then loaded with the load command:

```
save x;
```

loads x;
or
loads $\mathrm{x}=\mathrm{x} . \mathrm{fst}$;

The backslash is used as the escape character inside double quotes to enter special characters:

| "\b" | backspace (ASCII 8) |
| :--- | :--- |
| "\e" | escape (ASCII 27) |
| "\f" | formfeed (ASCII 12) |
| "\g" | beep (ASCII 7) |
| "\l" | line feed (ASCII 10) |
| "\r" | carriage return (ASCII 13) |
| "\t" | tab (ASCII 9) |
| " |  |
| " | a backslash |
| " |  |
| #\#\#" | the ASCII character whose decimal value is "\#\#\#". |

When entering DOS pathnames in double quotes, two backslashes must be used to insert one backslash:

```
st = "c:\\gauss\\myprog.prg";
```

An important use of strings and character elements of matrices is with the substitution operator ( ${ }^{\wedge}$ ).
In the command
create $f 1=$ olsdat with $\mathrm{x}, 4,2$;
by default, GAUSS will interpret the olsdat as a literal; that is, the literal name of the GAUSS data file you want to create. It will also interpret the $\mathbf{x}$ as the literal prefix string for the variable names: x1 x2 x3 x4.

If you want to get the data set name from a string variable, the substitution operator ( ${ }^{\wedge}$ ) could be used as:

```
dataset="olsdat";
create f1=^dataset with x,4,2;
```

If you want to get the data set name from a string variable and the variable names from a character vector, use

```
dataset="olsdat";
let vnames=age pay sex;
create f1=^dataset with `vnames,0,2;
```

The substitution operator ( ${ }^{\wedge}$ ) works with load and save also:

```
lpath="/gauss/procs";
name="mydata";
load path=^lpath x=^name;
command="dir *.fmt";
```

The general syntax is:

## ^variable name

Expressions are not allowed. The following commands are supported with the substitution operator ( ${ }^{\wedge}$ ):

```
create f1=^dataset with ^vnames,0,2;
create f1=^dataset using `cmdfile;
open f1=^dataset;
output file=^outfile;
load x=^datafile;
load path=`lpath x,y,z,t,w;
loadexe buf=^exefile;
save "name=x;
save path=^spath;
dos ^cmdstr;
run ^prog;
msym ^mstring;
```


### 10.6.6 String Arrays

String arrays are $\mathrm{N} \times \mathrm{K}$ matrices of strings. Here is a partial list of the functions for manipulating string arrays:

| $\$ \mid$ | Vertical string array concatenation operator. |
| :--- | :--- |
| $\$ \sim$ | Horizontal string array concatenation operator. |
| [] | Extract subarrays or individual strings from their corresponding <br> array, |
|  | or assign their values. |
| , | Transpose operator. |
| , | Bookkeeping transpose operator. |
| declare | Initialize variables at compile time. |
| delete | Delete specified global symbols. |
| fgetsa | Read multiple lines of text from a file. |
| fgetsat | Reads multiple lines of text from a file, discarding newlines. |


| format | Define output format for matrices, string arrays, and strings. |
| :---: | :---: |
| fputs | Write strings to a file. |
| fputst | Write strings to a file, appending newlines. |
| let | Initialize matrices, strings, and string arrays. |
| loads | Load a string or string array file (.fst file). |
| lprint | Print expressions to the printer. |
| lshow | Print global symbol table to the printer. |
| print | Print expressions on window and/or auxiliary output. |
| reshape | Reshape a matrix or string array to new dimensions. |
| save | Save matrix, string array, string, procedure, function or keyword to disk and gives the disk file either a .fmt, .fst or .fcg extension. |
| show | Display global symbol table. |
| sortcc | Quick-sort rows of matrix or string array based on character column. |
| type | Indicate whether variable passed as argument is matrix, string, or string array. |
| typecv | Indicate whether variables named in argument are strings, string arrays, matrices, procedures, functions or keywords. |
| varget | Access the global variable named by a string array. |
| varput | Assign the global variable named by a string array. |
| vec | Stack columns of a matrix or string array to form a column vector. |
| vecr | Stack rows of a matrix or string array to form a column vector. |

String arrays are created through the use of the string array concatenation operators. Below is a contrast of the horizontal string and horizontal string array concatenation operators.

```
x = "age";
y = "pay";
n = "sex";
s = x $+ y $+ n;
sa = x $~ y $~ n;
s=agepaysex
sa= age pay sex
```


### 10.6.7 Character Matrices

Matrices can have either numeric or character elements. For convenience, a matrix containing character elements is referred to as a character matrix.

A character matrix is not a separate data type, but gives you the ability to store and manipulate data elements that are composed of ASCII characters as well as floating point numbers. For example, you may want to concatenate a column vector containing the names of the variables in an analysis onto a matrix containing the coefficients, standard errors, t-statistic, and p-value. You can then print out the entire matrix with a separate format for each column with one call to the function printfm.

The logic of the programs will dictate the type of data assigned to a matrix, and the increased flexibility allowed by being able to bundle both types of data together in a single matrix can be very powerful. You could, for instance, create a moment matrix from your data, concatenate a new row onto it containing the names of the variables and save it to disk with the save command.

Numeric matrices are double precision, which means that each element is stored in 8 bytes. A character matrix can thus have elements of up to 8 characters.

GAUSS does not automatically keep track of whether a matrix contains character or numeric information. The ASCII to GAUSS conversion program ATOG will record the types of variables in a data set when it creates it. The create command will, also. The function vartypef gets a vector of variable type information from a data set. This vector of ones and zeros can be used by printfm when printing your data. Since GAUSS does not know whether a matrix has character or
numeric information, it is up to you to specify which type of data it contains when printing the contents of the matrix. (For details, see print and printfm in the GAUSS Language Reference.)

Most functions that take a string argument will take an element of a character matrix also, interpreting it as a string of up to 8 characters.

### 10.6.8 Date and Time Formats

## DT Scalar Format

The DT scalar format is a double precision representation of the date and time. In the DT scalar format, the number

$$
20010421183207
$$

represents 18:32:07 or 6:32:07 PM on April 21, 2001.

## DTV Vector Format

The DTV vector is a $1 \times 8$ vector. The format for the DTV vector is:
[1] Year
[2] Month, 1-12
[3] Day of month, 1-31
[4] Hour of day, 0-23
[5] Minute of hour, 0-59
[6] Second of minute, 0-59
[7] Day of week, 0-6 where 0 is Sunday
[8] Day since beginning of year, 0-365

## UTC Scalar Format

The UTC scalar format is the number of seconds since January 1, 1970, Greenwich Mean Time.

### 10.6.9 Special Data Types

The IEEE floating point format has many encodings that have special meaning. The print command will print them accurately so that you can tell if your calculation is producing meaningful results.

## NaN

There are many floating point encodings which do not correspond to a real number. These encodings are referred to as NaN's. NaN stands for Not A Number.

Certain numerical errors will cause the math coprocessor to create a NaN called an "indefinite". This will be printed as a -NaN when using the print command. These values are created by the following operations:

- $+\infty$ plus $-\infty$
- $+\infty$ minus $+\infty$
- $-\infty$ minus $-\infty$
- $0 * \infty$
- $\infty / \infty$
- $0 / 0$
- Operations where one or both operands is a NaN
- Trigonometric functions involving $\infty$


## INF

When the math coprocessor overflows, the result will be a properly signed infinity. Subsequent calculations will not deal well with an infinity; it usually signals an error in your program. The result of an operation involving an infinity is most often a NaN .

## DEN, UNN

When some math coprocessors underflow, they may do so gradually by shifting the significand of the number as necessary to keep the exponent in range. The result of this is a denormal (DEN). When denormals are used in calculations, they are usually handled automatically in an appropriate way. The result will either be an unnormal (UNN), which like the denormal represents a number very close to zero, or a normal, depending on how significant the effect of the denormal was in the calculation. In some cases the result will be a NaN .

Following are some procedures for dealing with these values. These procedures are not defined in the Run-Time Library. If you want to use them, you will need to define them yourself.

The procedure isindef will return 1 (true) if the matrix passed to it contains any NaN's that are the indefinite mentioned earlier. The GAUSS missing value code as well as GAUSS scalar error codes are NaN's, but this procedure tests only for indefinite:

```
proc isindef(x);
    retp(not x $/= __INDEFn);
endp;
```

Be sure to call gausset before calling isindef. gausset will initialize the value of the global _INDEFn to a platform-specific encoding.

The procedure normal will return a matrix with all denormals and unnormals set to zero.

```
proc normal(x);
    retp(x .* (abs(x) .> 4.19e-307));
endp;
```

The procedure isinf, will return 1 (true) if the matrix passed to it contains any infinities:

```
proc isinf(x);
    local plus,minus;
    plus = __INFp;
```

```
    minus = __INFn;
    retp(not x /= plus or not x /= minus);
endp;
```

Be sure to call gausset before calling isinf. gausset will initialize the values of the globals __INFn and __INFp to platform specific encodings.

### 10.7 Operator Precedence

The order in which an expression is evaluated is determined by the precedence of the operators involved and the order in which they are used. For example, the * and / operators have a higher precedence than the + and - operators. In expressions that contain these operators, the operand pairs associated with the * or / operator are evaluated first. Whether * or / is evaluated first depends on which comes first in the particular expression. For a listing of the precedence of all operators, see Operator Precedence, Section 11.7.

The expression

$$
-5+3 / 4+6 * 3
$$

is evaluated as

$$
(-5)+(3 / 4)+(6 * 3)
$$

Within a term, operators of equal precedence are evaluated from left to right.
The term

$$
2^{\wedge} 3^{\wedge} 7
$$

is evaluated

In the expression

$$
f 1(x) * f 2(y)
$$

$\mathbf{f 1}$ is evaluated before $\mathbf{f 2}$.
Here are some examples:

| Expression | Evaluation |
| :---: | :---: |
| a+b*c+d | $(a+(b * c))+d$ |
| $-2+4-6 * \operatorname{inv}(8) / 9$ | $((-2)+4)-((6 * \operatorname{inv}(8)) / 9)$ |
| 3.14^5*6/(2+sqrt (3)/4) | $\left(\left(3.14^{5}\right) * 6\right) /(2+(\operatorname{sqrt}(3) / 4))$ |
| $-\mathrm{a}+\mathrm{b}^{*} \mathrm{c}^{\wedge} 2$ | $(-a)+\left(b *\left(c^{2}\right)\right)$ |
| a+b-c+d-e | $(((a+b)-c)+d)-e$ |
| $a^{\wedge} b^{\wedge} c^{*} d$ | $\left(\left(a^{b}\right)^{c}\right) * d$ |
| a*b/d*c | $((a * b) / d) * c$ |
| $a^{\wedge} \mathrm{b}+\mathrm{c}^{*} \mathrm{~d}$ | $\left(a^{b}\right)+(c * d)$ |
| 2^4! | $2^{(4!)}$ |
| 2*3! | 2 * (3!) |

### 10.8 Flow Control

A computer language needs facilities for decision making and looping to control the order in which computations are done. GAUSS has several kinds of flow control statements.

### 10.8.1 Looping

## do loop

The do statement can be used in GAUSS to control looping.

> do while scalar_expression; /* loop if expression is true */

## statements

endo;
also
do until scalar_expression; /* loop if expression is false */

## statements

endo;

The scalar_expression is any expression that returns a scalar result. The expression will be evaluated as TRUE if its real part is nonzero and FALSE if it is zero. There is no counter variable that is automatically incremented in a do loop. If one is used, it must be set to its initial value before the loop is entered and explicitly incremented or decremented inside the loop.

The following example illustrates nested do loops that use counter variables.

```
format /rdn 1,0;
space = " ";
comma = ",";
i = 1;
do while i <= 4;
    j = 1;
```

```
    do while j <= 3;
        print space i comma j;;
        j = j+1;
endo;
i = i+1;
print;
endo;
```

This will print:

| 1,1 | 1,2 | 1,3 |
| :--- | :--- | :--- |
| 2,1 | 2,2 | 2,3 |
| 3,1 | 3,2 | 3,3 |
| 4,1 | 4,2 | 4,3 |

Use the relational and logical operators without the dot '.' in the expression that controls a do loop. These operators always return a scalar result.
break and continue are used within do loops to control execution flow. When break is encountered, the program will jump to the statement following the endo. This terminates the loop. When continue is encountered, the program will jump up to the top of the loop and reevaluate the while or until expression. This allows you to reiterate the loop without executing any more of the statements inside the loop:

```
do until eof(fp); /* continue jumps here */
    x = packr(readr(fp,100));
    if scalmiss(x);
        continue; /* iterate again */
    endif;
    s = s + sumc(x);
    count = count + rows(x);
    if count >= 10000;
        break; /* break out of loop */
    endif;
endo;
mean = s / count; /* break jumps here */
```


## for loop

The fastest looping construct in GAUSS is the for loop:
for counter (start, stop, step);
statements
endfor;
counter is the literal name of the counter variable. start, stop and step are scalar expressions. start is the initial value, stop is the final value and step is the increment.
break and continue are also supported by for loops. (For more information, see for in the GAUSS Language Reference.)

### 10.8.2 Conditional Branching

The if statement controls conditional branching:
if scalar_expression;
statements
elseif scalar_expression;

## statements

```
else;
    statements
endif;
```

The scalar_expression is any expression that returns a scalar result. The expression will be evaluated as TRUE if its real part is nonzero and FALSE if it is zero.

GAUSS will test the expression after the if statement. If it is TRUE, then the first list of statements is executed. If it is FALSE, then GAUSS will move to the expression after the first elseif statement, if there is one, and test it. It will keep testing expressions and will execute the first list of statements that corresponds to a TRUE expression. If no expression is TRUE, then the list of statements following the else statement is executed. After the appropriate list of statements is executed, the program will go to the statement following the endif and continue on.

Use the relational and logical operators without the dot ' ' in the expression that controls an if or elseif statement. These operators always return a scalar result.
if statements can be nested.

One endif is required per if clause. If an else statement is used, there may be only one per if clause. There may be as many elseif's as are required. There need not be any elseif's or any else statement within an if clause.

### 10.8.3 Unconditional Branching

The goto and gosub statements control unconditional branching. The target of both a goto and a gosub is a label.

## goto

A goto is an unconditional jump to a label with no return:

```
label:
goto label;
```

Parameters can be passed with a goto. The number of parameters is limited by available stack space. This is helpful for common exit routines:

```
    goto errout("Matrix singular");
    goto errout("File not found");
        .
errout:
    pop errmsg;
    errorlog errmsg;
    end;
```


## gosub

With a gosub, the address of the gosub statement is remembered and when a return statement is encountered, the program will resume executing at the statement following the gosub.

Parameters can be passed with a gosub in the same way as a goto. With a gosub it is also possible to return parameters with the return statement.

Subroutines are not isolated from the rest of your program and the variables referred to between the label and the return statement can be accessed from other places in your program.

Since a subroutine is only an address marked by a label, there can be subroutines inside of procedures. The variables used in these subroutines are the same variables that are known inside the procedure. They will not be unique to the subroutine, but they may be locals that are unique to
the procedure that the subroutine is in. (For details, see gosub in the GAUSS Language Reference.)

### 10.9 Functions

Single line functions that return one item can be defined with the fn statement.

$$
\mathrm{fn} \operatorname{area}(r)=\mathrm{pi} * r * r
$$

These functions can be called in the same way as intrinsic functions. The above function could be used in the following program sequence.

```
diameter = 3;
radius = 3 / 2;
a = area(radius);
```


### 10.10 Rules of Syntax

This section lists the general rules of syntax for GAUSS programs.

### 10.10.1 Statements

A GAUSS program consists of a series of statements. A statement is a complete expression or command. Statements in GAUSS end with a semicolon with one exception: from the GAUSS command line, the final semicolon in an interactive program is implicit if it is not explicitly given:

$$
\text { (gauss) } \mathrm{x}=5 ; \mathrm{z}=\mathrm{rndn}(3,3) ; \mathrm{y}=\mathrm{x}+\mathrm{z}
$$

Column position is not significant. Blank lines are allowed. Inside a statement and outside of double quotes, the carriage return/line feed at the end of a physical line will be converted to a space character as the program is compiled.

A statement containing a quoted string can be continued across several lines with a backslash as follows.

$$
\begin{aligned}
s= & \text { "This is one really long string that would be " } \backslash \\
& \text { "difficult to assign in just a single line."; }
\end{aligned}
$$

### 10.10.2 Case

GAUSS does not distinguish between uppercase and lowercase except inside double quotes.

### 10.10.3 Comments

// This comments out all text between the '//' and the end of
// the line
/* This kind of comment can be nested */
@ We consider this kind of comment to be obsolete, but it is supported for backwards compatibility @

### 10.10.4 Extraneous Spaces

Extraneous spaces are significant in print and lprint statements where the space is a delimiter between expressions:

```
print x y z;
```

In print and lprint statements, spaces can be used in expressions that are in parentheses:

```
print (x * y) (x + y);
```


### 10.10.5 Symbol Names

The names of matrices, strings, procedures, and functions can be up to 32 characters long. The characters must be alphanumeric or an underscore. The first character must be alphabetic or an underscore.

### 10.10.6 Labels

A label is used as the target of a goto or a gosub. The rules for naming labels are the same as for matrices, strings, procedures, and functions. A label is followed immediately by a colon:

```
here:
```

The reference to a label does not use a colon:

```
goto here;
```


### 10.10.7 Assignment Statements

The assignment operator is the equal sign ' $=$ ':

$$
y=x+z ;
$$

Multiple assignments must be enclosed in braces ' $\}$ ':

```
mant,pow = base10(x);
```

The comparison operator (equal to) is two equal signs ' $==$ ':

```
if x =\,= y;
    print "x is equal to y";
endif;
```


### 10.10.8 Function Arguments

The arguments to functions are enclosed in parentheses '( )':

$$
y=\operatorname{sqrt}(x) ;
$$

### 10.10.9 Indexing Matrices

Brackets '[ ]' are used to index matrices:

$$
\left.\begin{array}{l}
\mathrm{x}=\left\{\begin{array}{lll}
1 & 2 & 3, \\
3 & 7 & 5, \\
3 & 7 & 4, \\
8 & 9 & 5 \\
6 & 1 & 8
\end{array}\right\} ; \\
y=x[3,3] ; \\
z=x[1
\end{array} 2: 4,13\right] ;
$$

Vectors can be indexed with either one or two indices:

$$
\begin{aligned}
& \mathrm{v}=12334356 c c c \\
& \mathrm{k}=\mathrm{v}[3] ; \\
& \mathrm{j}=\mathrm{v}[1,6: 9] ;
\end{aligned}
$$

$\mathbf{x}[2,3]$ returns the element in the second row and the third column of $\mathbf{x}$.
$\mathbf{x}\left[\begin{array}{llll}1 & 3 & 5,4 & 7\end{array}\right]$ returns the submatrix that is the intersection of rows 1,3 , and 5 and columns 4 and 7.
$\mathbf{x}[., 3]$ returns the third column of $\mathbf{x}$.
$\mathbf{x}[3: 5,$.$] returns the submatrix containing the third through the fifth rows of \mathbf{x}$.
The indexing operator will take vector arguments for submatrix extraction or submatrix assignments:

$$
\begin{aligned}
& y=x[r v, c v] ; \\
& y[r v, c v]=x ;
\end{aligned}
$$

$\mathbf{r v}$ and $\mathbf{c v}$ can be any expressions returning vectors or matrices. The elements of $\mathbf{r v}$ will be used as the row indices and the elements of $\mathbf{c v}$ will be used as the column indices. If $\mathbf{r v}$ is a scalar 0 , all rows will be used; if $\mathbf{c v}$ is a scalar 0 , all columns will be used. If a vector is used in an index expression, it is illegal to use the space operator or the colon operator on the same side of the comma as the vector.

### 10.10.10 Arrays of Matrices and Strings

It is possible to index sets of matrices or strings using the varget function.
In this example, a set of matrix names is assigned to mvec. The name $y$ is indexed from mvec and passed to varget which will return the global matrix $\mathbf{y}$. The returned matrix is inverted and assigned to $\mathbf{g}$ :

```
mvec = { x y z a };
i = 2;
g = inv(varget(mvec[i]));
```


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The following procedure can be used to index the matrices in mvec more directly:

```
proc imvec(i);
    retp(varget(mvec[i]));
endp;
```

Then imvec(i) will equal the matrix whose name is in the $i^{\text {th }}$ element of mvec.
In the example above, the procedure imvec() was written so that it always operates on the vector mvec. The following procedure makes it possible to pass in the vector of names being used:

```
proc get(array,i);
    retp(varget(array[i]));
endp;
```

Then get (mvec, 3 ) will return the $3^{r d}$ matrix listed in mvec.

```
proc put(x,array,i);
    retp(varput(x,array[i]));
endp;
```

And put ( $\mathbf{x}$, mvec, 3 ) will assign $\mathbf{x}$ to the $3^{r d}$ matrix listed in mvec and return a 1 if successful or a 0 if it fails.

### 10.10.11 Arrays of Procedures

It is also possible to index procedures. The ampersand operator $(\boldsymbol{\&})$ is used to return a pointer to a procedure.

Assume that $\mathbf{f 1} \mathbf{f} \mathbf{2}$, and $\mathbf{f 3}$ are procedures that take a single argument. The following code defines a procedure fi that will return the value of the $i^{\text {th }}$ procedure, evaluated at $\mathbf{x}$.

```
nms = &f1 | &f2 | &f3;
proc fi(x,i);
    local f;
    f = nms[i];
    local f:proc;
    retp( f(x) );
endp;
```

$\mathbf{f i}(\mathbf{x}, \mathbf{2})$ will return $\mathbf{f} \mathbf{2 ( x )}$. The ampersand is used to return the pointers to the procedures. $\mathbf{n m s}$ is a numeric vector that contains a set of pointers. The local statement is used twice. The first tells the compiler that $\mathbf{f}$ is a local matrix. The $i^{t h}$ pointer, which is just a number, is assigned to $\mathbf{f}$. Then the second local statement tells the compiler to treat $\boldsymbol{f}$ as a procedure from this point on; thus the subsequent statement $\mathbf{f}(\mathbf{x})$ is interpreted as a procedure call.


## Operators

11

### 11.1 Element-by-Element Operators

Element-by-element operators share common rules of conformability. Some functions that have two arguments also operate according to the same rules.

Element-by-element operators handle those situations in which matrices are not conformable according to standard rules of matrix algebra. When a matrix is said to be $\mathrm{E} \times \mathrm{E}$ conformable, it refers to this element-by-element conformability. The following cases are supported:

```
matrix op matrix
matrix op scalar
scalar op matrix
matrix op vector
vector op matrix
vector op vector
```


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In a typical expression involving an element-by-element operator

$$
z=x+y ;
$$

conformability is defined as follows:

- If $x$ and $y$ are the same size, the operations are carried out corresponding element by corresponding element:

$$
\begin{aligned}
& x=\begin{array}{lll}
1 & 3 & 2 \\
4 & 5 & 1 \\
3 & 7 & 4 \\
& & \\
2 & 4 & 3 \\
3 & 1 & 4 \\
6 & 1 & 2 \\
\\
3 & 7 & 5 \\
7 & 6 & 5 \\
9 & 8 & 6
\end{array}
\end{aligned}
$$

- If $x$ is a matrix and $y$ is a scalar, or vice versa, then the scalar is operated on with respect to every element in the matrix. For example, $x+2$ will add 2 to every element of $x$ :

$$
\begin{aligned}
& \begin{array}{lll}
1 & 3 & 2
\end{array} \\
& x=\begin{array}{lll}
4 & 5 & 1
\end{array} \\
& 374 \\
& y=2 \\
& z=\begin{array}{lll}
3 & 5 & 4 \\
6 & 7 & 3 \\
5 & 9 & 6
\end{array}
\end{aligned}
$$

- If $x$ is an $\mathrm{N} \times 1$ column vector and $y$ is an $\mathrm{N} \times \mathrm{K}$ matrix, or vice versa, the vector is swept "across" the matrix:

| vector |  |  | matrix |  |
| :---: | :---: | :---: | :---: | :---: |
| 1 | $\longrightarrow$ | 2 | 4 | 3 |
| 4 | $\longrightarrow$ | 3 | 1 | 4 |
| 3 | $\longrightarrow$ | 6 | 1 | 2 |
|  |  |  | result |  |
|  |  | 3 | 5 | 4 |
|  |  | 7 | 5 | 8 |
|  |  | 9 | 4 | 5 |

- If $x$ is an $1 \times \mathrm{K}$ column vector and $y$ is an $\mathrm{N} \times \mathrm{K}$ matrix, or vice versa, then the vector is swept "down" the matrix:

| vector | 2 | 4 | 3 |
| :--- | :---: | :---: | :---: |
|  | $\downarrow$ | $\downarrow$ | $\downarrow$ |
|  | 2 | 4 | 3 |
| matrix | 3 | 1 | 4 |
|  | 6 | 1 | 2 |
|  | 4 | 8 | 6 |
| result | 5 | 5 | 7 |
|  | 8 | 5 | 5 |

- When one argument is a row vector and the other is a column vector, the result of an element-by-element operation will be the "table" of the two:

| row vector |  | 2 | 4 | 3 | 1 |
| :--- | :--- | :---: | :---: | :---: | :---: |
| column vector | 2 | 4 | 6 | 5 | 3 |
|  | 5 | 7 | 6 | 4 |  |
|  | 5 | 7 | 9 | 8 | 6 |

If $x$ and $y$ are such that none of these conditions apply, the matrices are not conformable to these operations and an error message will be generated.

### 11.2 Matrix Operators

The following operators work on matrices. Some assume numeric data and others will work on either character or numeric data.

### 11.2.1 Numeric Operators

For details on how matrix conformability is defined for element-by-element operators, see Element-by-Element Operators, Section 11.1.
$+\quad$ Addition

$$
y=x+z
$$

Performs element-by-element addition.

- Subtraction or negation

$$
\begin{aligned}
& y=x-z ; \\
& y=-k ;
\end{aligned}
$$

Performs element-by-element subtraction or the negation of all elements, depending on context.

* Matrix multiplication or multiplication

$$
y=x * z
$$

When $z$ has the same number of rows as $x$ has columns, this will perform matrix multiplication (inner product). If $x$ or $z$ are scalar, this performs standard element-by-element multiplication.
/ Division or linear equation solution

$$
x=b / A ;
$$

If $A$ and $b$ are scalars, this performs standard division. If one of the operands is a matrix and the other is scalar, the result is a matrix the same size with the results of the divisions between the scalar and the corresponding elements of the matrix. Use ./ for element-by-element division of matrices.
If $b$ and $A$ are conformable, this operator solves the linear matrix equation

$$
A x=b
$$

Linear equation solution is performed in the following cases:

- If $A$ is a square matrix and has the same number of rows as $b$, this statement will solve the system of linear equations using an LU decomposition.
- If $A$ is rectangular with the same number of rows as $b$, this statement will produce the least squares solutions by forming the normal equations and using the Cholesky decomposition to get the solution:

$$
x=\frac{A^{\prime} b}{A^{\prime} A}
$$

If trap 2 is set, missing values will be handled with pairwise deletion.
\% Modulo division

$$
y=x \% z ;
$$

For integers, this returns the integer value that is the remainder of the integer division of $x$ by $z$. If $x$ or $z$ is noninteger, it will first be rounded to the nearest integer. This is an element-by-element operator.
! Factorial

$$
y=x!;
$$

Computes the factorial of every element in the matrix $x$. Nonintegers are rounded to the nearest integer before the factorial operator is applied. This will not work with complex matrices. If $x$ is complex, a fatal error will be generated.
.* Element-by-element multiplication

$$
y=x . * z ;
$$

If $x$ is a column vector, and $z$ is a row vector (or vice versa), the "outer product" or "table" of the two will be computed. (For comformability rules, see Element-by-Element Operators, Section 11.1.)
./ Element-by-element division

$$
y=x . / z ;
$$

^ Element-by-element exponentiation

$$
y=x^{\wedge} z ;
$$

If $x$ is negative, $z$ must be an integer.
.^ Same as ^
.*. Kronecker (tensor) product

$$
y=x . * . \quad z ;
$$

This results in a matrix in which every element in $x$ has been multiplied (scalar multiplication) by the matrix $z$. For example:

$$
\begin{aligned}
& \mathrm{x}=\{12, \\
& 34 \text { \}; } \\
& z=\{456, \\
& 789\} ; \\
& y=x . * . z ; \\
& x=\begin{array}{ll}
1 & 2 \\
3 & 4
\end{array} \\
& z=\begin{array}{lll}
4 & 5 & 6 \\
7 & 8 & 9
\end{array} \\
& y=\begin{array}{rrrrrr}
4 & 5 & 6 & 8 & 10 & 12 \\
7 & 8 & 9 & 14 & 16 & 18 \\
12 & 15 & 18 & 16 & 20 & 24 \\
21 & 24 & 27 & 28 & 32 & 36
\end{array}
\end{aligned}
$$

*0@*~ Horizontal direct product

$$
\begin{aligned}
& z=x * \sim y ; \\
& x=\begin{array}{ll}
z & 2 \\
3 & 4
\end{array} \\
& y=\begin{array}{rr}
5 & 6 \\
7 & 8
\end{array} \\
& z=\begin{array}{rrrr}
5 & 6 & 10 & 12 \\
21 & 24 & 28 & 32
\end{array}
\end{aligned}
$$

The input matrices $x$ and $y$ must have the same number of rows. The result will have $\operatorname{cols}(x){ }^{*} \operatorname{cols}(y)$ columns.

### 11.2.2 Other Matrix Operators

, Transpose operator

$$
y=x^{\prime} ;
$$

The columns of $y$ will contain the same values as the rows of $x$ and the rows of $y$ will contain the same values as the columns of $x$. For complex matrices this computes the complex conjugate transpose.
If an operand immediately follows the transpose operator, the ' will be interpreted as ${ }^{\prime *}$. Thus $y=x^{\prime} x$ is equivalent to $y=x^{\prime *} x$.
. Bookkeeping transpose operator

$$
y=x .^{\prime} \text {; }
$$

This is provided primarily as a matrix handling tool for complex matrices. For all matrices, the columns of $y$ will contain the same values as the rows of $x$ and the rows of $y$ will contain the same values as the columns of $x$. The complex conjugate transpose is NOT computed when you use . ${ }^{\prime}$.
If an operand immediately follows the bookkeeping transpose operator, the.$^{\prime}$ will be interpreted as.$^{\prime *}$. Thus $y=x .^{\prime} x$ is equivalent to $y=x .^{\prime *} x$.
| Vertical concatenation

$$
\begin{aligned}
& z=x \mid y ; \\
& x=\begin{array}{lll}
z=\begin{array}{lll}
1 & 2 & 3 \\
3 & 4 & 5
\end{array} \\
y=\begin{array}{lll}
7 & 8 & 9
\end{array} \\
z=\begin{array}{lll}
1 & 2 & 3 \\
3 & 4 & 5 \\
7 & 8 & 9
\end{array}
\end{array}, \left.\begin{array}{l} 
\\
y
\end{array} \right\rvert\,
\end{aligned}
$$

Horizontal concatenation

$$
\begin{aligned}
& z=x \sim y ; \\
& x=\begin{array}{ll}
1 & 2 \\
3 & 4
\end{array} \\
& y=\begin{array}{lll}
5 & 6 \\
7 & 8
\end{array} \\
& z=\begin{array}{llll}
1 & 2 & 5 & 6 \\
3 & 4 & 7 & 8
\end{array}
\end{aligned}
$$

### 11.3 Relational Operators

For details on how matrix conformability is defined for element-by-element operators, see Element-by-Element Operators, Section 11.1

Each of these operators has two equivalent representations. Either can be used (for example, < or $\mathbf{l t}$ ), depending only upon preference. The alphabetic form should be surrounded by spaces.

A third form of these operators has a ' $\$$ ' and is used for comparisons between character data and for comparisons between strings or string arrays. The comparisons are done byte by byte starting with the lowest addressed byte of the elements being compared.

The equality comparison operators ( $<=,==,>=, /=$ ) and their dot equivalents can be used to test for missing values and the NaN that is created by floating point exceptions. Less than and greater than comparisons are not meaningful with missings or NaN's, but equal and not equal are valid. These operators are sign-insensitive for missings, NaN's, and zeros.

The string '\$' versions of these operators can also be used to test missings, NaN's and zeros. Because they do a strict byte-to-byte comparison, they are sensitive to the sign bit. Missings, NaN's, and zeros can all have the sign bit set to 0 or 1 , depending on how they were generated and have been used in a program.

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If the relational operator is NOT preceded by a dot ' $:$ ', then the result is always a scalar 1 or 0 , based upon a comparison of all elements of $x$ and $y$. All comparisons must be true for the relational operator to return TRUE.

By this definition, then

$$
\text { if } x /=y \text {; }
$$

is interpreted as: "if every element of $x$ is not equal to the corresponding element of $y$ ". To check if two matrices are not identical, use

$$
\text { if not } x==y \text {; }
$$

For complex matrices, the $==, /=, .==$ and.$/=$ operators compare both the real and imaginary parts of the matrices; all other relational operators compare only the real parts.

- Less than

$$
\begin{aligned}
& z=x<y \\
& z=x \text { lt } y ; \\
& z=x \$<y ;
\end{aligned}
$$

- Less than or equal to

$$
\begin{aligned}
& z=x<=y \\
& z=x \text { le } y \\
& z=x \$<=y
\end{aligned}
$$

- Equal to

$$
z=x==y
$$

$$
\begin{aligned}
& z=x \text { eq } y ; \\
& z=x \$==y ;
\end{aligned}
$$

- Not equal

$$
\begin{aligned}
& z=x /=y \\
& z=x \text { ne } y \\
& z=x \$ /=y
\end{aligned}
$$

- Greater than or equal to

$$
\begin{aligned}
& z=x>=y \\
& z=x \text { ge } y \\
& z=x \$>=y ;
\end{aligned}
$$

- Greater than

$$
\begin{aligned}
& z=x>y ; \\
& z=x \text { gt } y ; \\
& z=x \$>y ;
\end{aligned}
$$

If the relational operator IS preceded by a dot ' $\because$ ', then the result will be a matrix of 1 's and 0 's, based upon an element-by-element comparison of $x$ and $y$.

- Element-by-element less than

$$
\begin{aligned}
& z=x .<y \\
& z=x . \operatorname{lt} y \\
& z=x . \$<y
\end{aligned}
$$

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- Element-by-element less than or equal to

$$
\begin{aligned}
& z=x .<=y \\
& z=x . \text { le } y \\
& z=x . \$<=y
\end{aligned}
$$

- Element-by-element equal to

$$
\begin{aligned}
& z=x .==y \\
& z=x . \text { eq } y \\
& z=x . \$==y ;
\end{aligned}
$$

- Element-by-element not equal to

$$
\begin{aligned}
& z=x . /=y \\
& z=x . \text { ne } y \\
& z=x . \$ /=y
\end{aligned}
$$

- Element-by-element greater than or equal to

$$
\begin{aligned}
& z=x .>=y \\
& z=x \cdot \text { ge } y \\
& z=x . \$>=y
\end{aligned}
$$

- Element-by-element greater than

$$
\begin{aligned}
& z=x .>y ; \\
& z=x . \text { gt } y ; \\
& z=x . \$>y ;
\end{aligned}
$$

### 11.4 Logical Operators

The logical operators perform logical or Boolean operations on numeric values. On input a nonzero value is considered TRUE and a zero value is considered FALSE. The logical operators return a 1 if TRUE and a 0 if FALSE. Decisions are based on the following truth tables:

## Complement

| $X$ | $\operatorname{not} X$ |
| :---: | :---: |
| T | F |
| F | T |

## Conjunction

| $X$ | $Y$ | $X$ and $Y$ |
| :---: | :---: | :---: |
| T | T | T |
| T | F | F |
| F | T | F |
| F | F | F |

## Disjunction

| $X$ | $Y$ | $X$ or $Y$ |
| :---: | :---: | :---: |
| T | T | T |
| T | F | T |
| F | T | T |
| F | F | F |

## Exclusive Or

| $X$ | $Y$ | $X$ xor $Y$ |
| :---: | :---: | :---: |
| T | T | F |
| T | F | T |
| F | T | T |
| F | F | F |

## Equivalence

| $X$ | $Y$ | $X$ eqv $Y$ |
| :---: | :---: | :---: |
| T | T | T |
| T | F | F |
| F | T | F |
| F | F | T |

For complex matrices, the logical operators consider only the real part of the matrices.
The following operators require scalar arguments. These are the ones to use in if and do statements:

- Complement

$$
z=\operatorname{not} x
$$

- Conjunction

$$
z=x \text { and } y ;
$$

- Disjunction

$$
z=x \text { or } y
$$

- Exclusive or

$$
z=x \text { xor } y ;
$$

- Equivalence

$$
z=x \text { eqv } y ;
$$

If the logical operator is preceded by a dot '. ', the result will be a matrix of 1 's and 0 's based upon an element-by-element logical comparison of $x$ and $y$ :

- Element-by-element logical complement

$$
z=. \operatorname{not} x ;
$$

- Element-by-element conjunction

$$
z=x \text {.and } y ;
$$

- Element-by-element disjunction

$$
z=x \text {.or } y ;
$$

- Element-by-element exclusive or

$$
z=x \text {.xor } y ;
$$

- Element-by-element equivalence

$$
z=x \text {.eqv } y ;
$$

### 11.5 Other Operators

## Assignment Operator

Assignments are done with one equal sign:

$$
y=3
$$

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## Comma

Commas are used to delimit lists:

```
clear x,y,z;
```

to separate row indices from column indices within brackets:

$$
y=x[3,5] ;
$$

and to separate arguments of functions within parentheses:

$$
\mathrm{y}=\operatorname{momentd}(\mathrm{x}, \mathrm{~d}) ;
$$

## Period

Dots are used in brackets to signify "all rows" or "all columns":

$$
y=x[., 5] ;
$$

## Space

Spaces are used inside of index brackets to separate indices:

$$
\mathrm{y}=\mathrm{x}\left[\begin{array}{lllll}
1 & 3 & 5,3 & 5 & 9
\end{array}\right] ;
$$

No extraneous spaces are allowed immediately before or after the comma, or immediately after the left bracket or before the right bracket.

Spaces are also used in print and lprint statements to separate the separate expressions to be printed:

```
print x/2 2*sqrt(x);
```

No extraneous spaces are allowed within expressions in print or lprint statements unless the expression is enclosed in parentheses:

```
print (x / 2) (2 * sqrt(x));
```


## Colon

A colon is used within brackets to create a continuous range of indices:

$$
y=x[1: 5, .] ;
$$

## Ampersand

The (\&) ampersand operator will return a pointer to a procedure (proc), function (fn), or structure (struct). It is used when passing procedures or functions to other functions, when indexing procedures, and when initializing structure pointers. (For more information, see Indexing Procedures, Section 12.5 or Structure Pointers, Section 16.2.)

## String Concatenation

$$
\begin{aligned}
& \mathrm{x}=\text { "dog"; } \\
& \mathrm{y}=\text { "cat"; } \\
& \mathrm{z}=\mathrm{x} \$+\mathrm{y} ; \\
& \text { print } \mathrm{z} ; \\
& \text { dogcat }
\end{aligned}
$$

If the first argument is of type string, the result will be of type string. If the first argument is of type matrix, the result will be of type matrix. Here are some examples:
y = 0 \$+ "caterpillar";

The result will be a $1 \times 1$ matrix containing 'caterpil'.
y = zeros(3,1) \$+ "cat";

The result will be a $3 \times 1$ matrix, each element containing 'cat'.
If we use the $y$ created above in the following:
k = y \$+ "fish";

The result will be a $3 \times 1$ matrix with each element containing 'catfish'.
If we then use $k$ created above:

$$
\mathrm{t}=\mathrm{"}=\$ \mathrm{k}[1,1] ;
$$

The result will be a string containing 'catfish'.
If we used the same $k$ to create $z$ as follows:
z = "dog" \$+ k[1,1];

The resulting $z$ will be a string containing 'dogcatfish'.

## String Array Concatenation

\$| Vertical string array concatenation

$$
\begin{aligned}
& \text { x = "dog"; } \\
& \text { y = "fish"; } \\
& \text { k = x \$। y; } \\
& \text { print k; }
\end{aligned}
$$

dog
fish
$\$ \sim$ Horizontal string array concatenation

$$
\begin{aligned}
& \text { x = "dog"; } \\
& \text { y = "fish"; } \\
& \text { k = } \text { \$ }^{\text {y }} \text {; } \\
& \text { print k; }
\end{aligned}
$$

$$
\begin{array}{ll}
\operatorname{dog} & \text { fish }
\end{array}
$$

## String Variable Substitution

In a command like the following:

```
create f1 = olsdat with x,4,2;
```

by default GAUSS will interpret olsdat as the literal name of the GAUSS data file you want to create. It will also interpret $\mathbf{x}$ as the literal prefix string for the variable names $\mathbf{x 1} \mathbf{x} \mathbf{x} \mathbf{x 4}$.

To get the data set name from a string variable, the substitution operator ( ${ }^{\wedge}$ ) could be used as follows:

```
dataset = "olsdat";
create f1 = ^dataset with x,4,2;
```

To get the data set name from a string variable and the variable names from a character vector, use the following:

```
dataset = "olsdat";
vnames = { age, pay, sex };
create f1 = ^dataset with `vnames,0,2;
```

The general syntax is:

```
`variable_name
```

Expressions are not allowed.
The following commands are currently supported with the substitution operator ( ${ }^{\wedge}$ ) in the current version.

```
create f1 = ^dataset with `vnames,0,2;
create f1 = ^dataset using `cmdfile;
open f1 = ^dataset;
output file = ^outfile;
load x = ^datafile;
load path = ``lpath x,y,z,t,w;
loadexe buf = `exefile;
save `name = x;
save path = "spath;
dos ^cmdstr;
run ^prog;
msym ^mstring;
```


### 11.6 Using Dot Operators with Constants

When you use those operators preceded by a '.' (dot operators) with a scalar integer constant, insert a space between the constant and any following dot operator. Otherwise, the dot will be interpreted as part of the scalar; that is, the decimal point. For example:

$$
\begin{aligned}
& \text { let } y=123 ; \\
& x=2 .<y ;
\end{aligned}
$$

will return $\mathbf{x}$ as a scalar 0 , not a vector of 0 's and 1 's, because

$$
x=2 .<y ;
$$

is interpreted as

$$
x=2 .<y
$$

and not as

$$
x=2 .<y
$$

Be careful when using the dot relational operators (. $<, .<=, .==, . /=, .>, .>=$ ). The same problem can occur with other dot operators, also. For example:

$$
\begin{aligned}
& \text { let } x=111 ; \\
& y=x . / 2 . / x ;
\end{aligned}
$$

will return $y$ as a scalar .5 rather than a vector of .5 's, because

$$
y=x . / 2 . / x ;
$$

is interpreted as

$$
y=(x . / 2 .) / x ;
$$

not

$$
y=(x . / 2) . / x ;
$$

The second division, then, is handled as a matrix division rather than an element-by-element division.

### 11.7 Operator Precedence

The order in which an expression is evaluated is determined by the precedence of the operators involved and the order in which they are used. For example, the * and / operators have a higher precedence than the + and - operators. In expressions that contain the above operators, the operand pairs associated with the * or / operator are evaluated first. Whether * or / is evaluated first depends on which comes first in the particular expression.

The expression

$$
-5+3 / 4+6 * 3
$$

is evaluated as

$$
(-5)+(3 / 4)+(6 * 3)
$$

Within a term, operators of equal precedence are evaluated from left to right. The precedence of all operators, from the highest to the lowest, is listed in the following table:

| Operator | Precedence | Operator | Precedence | Operator | Precedence |
| :---: | :---: | :---: | :---: | :---: | :---: |
| .$^{\prime}$ | 90 | . \$>= | 65 | \$>= | 55 |
| , | 90 | ./= | 65 | /= | 55 |
| ! | 89 | . $<$ | 65 | $<$ | 55 |
| .^ | 85 | .<= | 65 | <= | 55 |
| ^ | 85 | .= | 65 | = | 55 |
| (unary -) | 83 | .> | 65 | > | 55 |
| * | 80 | .>= | 65 | >= | 55 |
| * | 80 | .eq | 65 | eq | 55 |
| ** | 80 | .ge | 65 | ge | 55 |
| .*. | 80 | .gt | 65 | gt | 55 |
| ./ | 80 | .le | 65 | le | 55 |
| 1 | 80 | .lt | 65 | 1 t | 55 |
| \% | 75 | .ne | 65 | ne | 55 |
| \$+ | 70 | .not | 64 | not | 49 |
| + | 70 | . and | 63 | and | 48 |
| - | 70 | .or | 62 | or | 47 |
| $\sim$ | 68 | .xor | 61 | xor | 46 |
| \| | 67 | . eqv | 60 | eqv | 45 |
| . \$/= | 65 | \$/= | 55 | (space) | 35 |
| . $\$$ | 65 | \$< | 55 | , | 35 |
| . \$<= | 65 | \$<= | 55 | $=$ | 10 |
| . \$= | 65 | \$= $=$ | 55 |  |  |
| . \$> | 65 | \$> | 55 |  |  |



## Procedures and Keywords

## 12

Procedures are multiple-line, recursive functions that can have either local or global variables. Procedures allow a large computing task to be written as a collection of smaller tasks. These smaller tasks are easier to work with and keep the details of their operation from the other parts of the program that do not need to know them. This makes programs easier to understand and easier to maintain.

A procedure in GAUSS is basically a user-defined function that can be used as if it were an intrinsic part of the language. A procedure can be as small and simple or as large and complicated as necessary to perform a particular task. Procedures allow you to build on your previous work and on the work of others rather than starting over again and again to perform related tasks.

Any intrinsic command or function may be used in a procedure, as well as any user-defined function or other procedure. Procedures can refer to any global variable; that is, any variable in the global symbol table that can be shown with the show command. It is also possible to declare local variables within a procedure. These variables are known only inside the procedure they are defined in and cannot be accessed from other procedures or from the main level program code.

All labels and subroutines inside a procedure are local to that procedure and will not be confused with labels of the same name in other procedures.

### 12.1 Defining a Procedure

A procedure definition consists of five parts, four of which are denoted by explicit GAUSS commands:

1. Procedure declaration proc statement
2. Local variable declaration local statement
3. Body of procedure
4. Return from procedure retp statement
5. End of procedure definition endp statement

There is always one proc statement and one endp statement in a procedure definition. Any statements that come between these two statements are part of the procedure. Procedure definitions cannot be nested. local and retp statements are optional. There can be multiple local and retp statements in a procedure definition. Here is an example:

```
proc (3) = regress(x, y);
    local xxi,b,ymxb,sse,sd,t;
    xxi = invpd(x'x);
    b = xxi * (x'y);
    ymxb = y-xb;
    sse = ymxb'ymxb/(rows(x)-cols(x));
    sd = sqrt(diag(sse*xxi));
    t = b./sd;
    retp(b,sd,t);
endp;
```

This could be used as a function that takes two matrix arguments and returns three matrices as a result. For example: is:

$$
\{b, \operatorname{sd}, \mathrm{t}\}=\operatorname{regress}(\mathrm{x}, \mathrm{y}) ;
$$

Following is a discussion of the five parts of a procedure definition.

### 12.1.1 Procedure Declaration

The proc statement is the procedure declaration statement. The format is:

```
proc \llbracket(rets) =\rrbracket name(\llbracketarg1,\operatorname{arg2,\ldots..argN|);}
```

rets Optional constant, number of values returned by the procedure. Acceptable values here are $0-1023$; the default is 1 .
name Name of the procedure, up to 32 alphanumeric characters or an underscore, beginning with an alpha or an underscore.
arg\# Names that will be used inside the procedure for the arguments that are passed to the procedure when it is called. There can be 0-1023 arguments. These names will be known only in the procedure being defined. Other procedures can use the same names, but they will be separate entities.

### 12.1.2 Local Variable Declarations

The local statement is used to declare local variables. Local variables are variables known only to the procedure being defined. The names used in the argument list of the proc statement are always local. The format of the local statement is:

```
local x,y,f:proc,g:fn,z,h:keyword;
```

Local variables can be matrices or strings. If :proc, : fn, or : keyword follows the variable name in the local statement, the compiler will treat the symbol as if it were a procedure, function, or keyword respectively. This allows passing procedures, functions, and keywords to other procedures. (For more information, see Passing Procedures to Procedures, Section 12.4.

Variables that are global to the system (that is, variables listed in the global symbol table that can be shown with the show command) can be accessed by any procedure without any redundant declaration inside the procedure. If you want to create variables known only to the procedure

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being defined, the names of these local variables must be listed in a local statement. Once a variable name is encountered in a local statement, further references to that name inside the procedure will be to the local rather than to a global having the same name. (See clearg, varget, and varput in the GAUSS Language Reference for ways of accessing globals from within procedures that have locals with the same name.)

The local statement does not initialize (set to a value) the local variables. If they are not passed in as parameters, they must be assigned some value before they are accessed or the program will terminate with a Variable not initialized error message.

All local and global variables are dynamically allocated and sized automatically during execution. Local variables, including those that were passed as parameters, can change in size during the execution of the procedure.

Local variables exist only when the procedure is executing and then disappear. Local variables cannot be listed with the show command.

The maximum number of locals is limited by stack space and the size of workspace memory. The limiting factor applies to the total number of active local symbols at any one time during execution. If cat has 10 locals and it calls dog which has 20 locals, there are 30 active locals whenever cat is called.

There can be multiple local statements in a procedure. They will affect only the code in the procedure that follows. Therefore, for example, it is possible to refer to a global $\mathbf{x}$ in a procedure and follow that with a local statement that declares a local $\mathbf{x}$. All subsequent references to $\mathbf{x}$ would be to the local $\mathbf{x}$. (This is not good programming practice, but it demonstrates the principle that the local statement affects only the code that is physically below it in the procedure definition.) Another example is a symbol that is declared as a local and then declared as a local procedure or function later in the same procedure definition. This allows doing arithmetic on local function pointers before calling them. (For more information, see Indexing Procedures, Section 12.5.

### 12.1.3 Body of Procedure

The body of the procedure can have any GAUSS statements necessary to perform the task the procedure is being written for. Other user-defined functions and other procedures can be referenced as well as any global matrices and strings.

GAUSS procedures are recursive, so the procedure can call itself as long as there is logic in the procedure to prevent an infinite recursion. The process would otherwise terminate with either an Insufficient workspace memory message or a Procedure calls too deep message, depending on the space necessary to store the locals for each separate invocation of the procedure.

### 12.1.4 Returning from the Procedure

The return from the procedure is accomplished with the retp statement:

```
retp;
retp(expression1,expression2,\ldots,expressionN);
```

The retp statement can have multiple arguments. The number of items returned must coincide with the number of rets in the proc statement.

If the procedure was defined with no items returned, the retp statement is optional. The endp statement that ends the procedure will generate an implicit retp with no objects returned. If the procedure returns one or more objects, there must be an explicit retp statement.

There can be multiple retp statements in a procedure, and they can be anywhere inside the body of the procedure.

### 12.1.5 End of Procedure Definition

The endp statement marks the end of the procedure definition: endp;

An implicit retp statement that returns nothing is always generated here so it is impossible to run off the end of a procedure without returning. If the procedure was defined to return one or more objects, executing this implicit return will result in a Wrong number of returns error message and the program will terminate.

### 12.2 Calling a Procedure

Procedures are called like this:

```
dog(i,j,k); /* no returns */
y = cat(i,j,k); /* one return */
{ x,y,z } = bat(i,j,k); /* multiple returns */
call bat(i,j,k); /* ignore any returns */
```

Procedures are called in the same way that intrinsic functions are called. The procedure name is followed by a list of arguments in parentheses. The arguments must be separated by commas.

If there is to be no return value, use

```
proc (0) = dog(x,y,z);
```

when defining the procedure and use

```
dog(ak,4,3);
```

or

$$
\text { call } \operatorname{dog}(a k, 4,3) ;
$$

when calling it.
The arguments passed to procedures can be complicated expressions involving calls to other functions and procedures. This calling mechanism is completely general. For example,

$$
y=\operatorname{dog}(\operatorname{cat}(3 * x, \operatorname{bird}(x, y))-2,2,1) ;
$$

is legal.

### 12.3 Keywords

A keyword, like a procedure, is a subroutine that can be called interactively or from within a GAUSS program. A keyword differs from a procedure in that a keyword accepts exactly one string argument, and returns nothing. Keywords can perform many tasks not as easily accomplished with procedures.

### 12.3.1 Defining a Keyword

A keyword definition is much like a procedure definition. Keywords always are defined with 0 returns and 1 argument. The beginning of a keyword definition is the keyword statement:
keyword name (strarg);
name Name of the keyword, up to 32 alphanumeric characters or an underscore, beginning with an alpha or an underscore.
strarg Name that will be used inside of the keyword for the argument that is passed to the keyword when it is called. There is always one argument. The name is known only in the keyword being defined. Other keywords can use the same name, but they will be separate entities. This will always be a string. If the keyword is called with no characters following the name of the keyword, this will be a null string.

The rest of the keyword definition is the same as a procedure definition. (For more information, see Defining a Procedure, Section 12.1. Keywords always return nothing. Any retp statements, if used, should be empty. For example:

```
keyword add(s);
    local tok, sum;
    if s $=\,= "";
        print "The argument is a null string";
        retp;
    endif;
    print "The argument is: '" s "'";
    sum = 0;
    do until s $=\,= "";
        { tok, s } = token(s);
        sum = sum + stof(tok);
    endo;
    format /rd 1,2;
    print "The sum is: " sum;
endp;
```

The keyword defined above will print the string argument passed to it. The argument will be printed enclosed in single quotes.

### 12.3.2 Calling a Keyword

When a keyword is called, every character up to the end of the statement, excluding the leading spaces, is passed to the keyword as one string argument. For example, if you type

```
add 1 2 3 4 5;
```

the keyword will respond

The sum is: 15.00

Here is another example:
add;
the keyword will respond

The argument is a null string

### 12.4 Passing Procedures to Procedures

Procedures and functions can be passed to procedures in the following way:

```
proc max(x,y); /* procedure to return maximum */
    if x>y;
        retp(x);
    else;
        retp(y);
    endif;
endp;
proc min(x,y); /* procedure to return minimum */
    if x<y;
        retp(x);
    else;
        retp(y);
    endif;
endp;
fn lgsqrt(x) = ln(sqrt(x)); /* function to return
    :: log of square root
    */
```

```
proc myproc(&f1,&f2,x,y);
    local f1:proc, f2:fn, z;
    z = f1(x,y);
    retp(f2(z));
endp;
```

The procedure myproc takes four arguments. The first is a procedure $f 1$ that has two arguments. The second is a function $\mathbf{f 2}$ that has one argument. It also has two other arguments that must be matrices or scalars. In the local statement, $\mathbf{f 1}$ is declared to be a procedure and $\mathbf{f 2}$ is declared to be a function. They can be used inside the procedure in the usual way. $\mathbf{f 1}$ will be interpreted as a procedure inside myproc, and $£ 2$ will be interpreted as a function. The call to myproc is made as follows:

```
k = myproc(&max,&lgsqrt,5,7); /* log of square root of 7 */
k = myproc(&min,&lgsqrt,5,7); /* log of square root of 5 */
```

The ampersand (\&) in front of the function or procedure name in the call to myproc causes a pointer to the function or procedure to be passed. No argument list should follow the name when it is preceded by the ampersand.

Inside myproc, the symbol that is declared as a procedure in the local statement is assumed to contain a pointer to a procedure. It can be called exactly like a procedure is called. It cannot be save'd but it can be passed on to another procedure. If it is to be passed on to another procedure, use the ampersand in the same way.

### 12.5 Indexing Procedures

This example assumes there are a set of procedures named $f 1-£ 5$ that are already defined. A $1 \times 5$ vector procvec is defined by horizontally concatenating pointers to these procedures. A new procedure, $\mathbf{g}(x, i)$ is then defined to return the value of the $i^{t h}$ procedure evaluated at $x$ :

```
procvec = &f1 ~ &f2 ~ &f3 ~ &f4 ~ &f5;
```

```
proc g(x,i);
    local f;
    f = procvec[i];
    local f:proc;
    retp( f(x) );
endp;
```

The local statement is used twice. The first time, $\mathbf{f}$ is declared to be a local matrix. After $\mathbf{f}$ has been set equal to the $i^{\text {th }}$ pointer, $\mathbf{f}$ is declared to be a procedure and is called as a procedure in the retp statement.

### 12.6 Multiple Returns from Procedures

Procedures can return multiple items, up to 1023. The procedure is defined like this example of a complex inverse:

```
proc (2) = cminv(xr,xi); /* (2) specifies number of
                                    :: return values
                                    */
    local ixy, zr, zi;
    ixy = inv(xr)*xi;
    zr = inv(xr+xi*ixy); /* real part of inverse. */
    zi = -ixy*zr; /* imaginary part of inverse. */
    retp(zr,zi); /* return: real part, imaginary part */
endp;
```

It can then be called like this:

$$
\{\mathrm{zr}, \mathrm{zi}\}=\operatorname{cminv}(\mathrm{xr}, \mathrm{xi}) ;
$$

To make the assignment, the list of targets must be enclosed in braces.

Also, a procedure that returns more than one argument can be used as input to another procedure or function that takes more than one argument:

```
proc (2) = cminv(xr,xi);
    local ixy, zr, zi;
    ixy = inv(xr)*xi;
    zr = inv(xr+xi*ixy); /* real part of inverse. */
    zi = -ixy*zr; /* imaginary part of inverse. */
    retp(zr,zi);
endp;
proc (2) = cmmult(xr,xi,yr,yi);
    local zr,zi;
    zr = xr*yr-xi*yi;
    zi = xr*yi+xi*yr;
    retp(zr,zi);
endp;
{ zr,zi } = cminv( cmmult(xr,xi,yr,yi) );
```

The two returned matrices from cmmult() are passed directly to cminv() in the statement above. This is equivalent to the following statements:

```
{ tr,ti } = cmmult(xr,xi,yr,yi);
{ zr,zi } = cminv(tr,ti);
```

This is completely general so the following program is legal:

```
proc (2) \(=\operatorname{cmcplx}(x)\);
    local r,c;
    \(r=\operatorname{rows}(x)\);
    c \(=\operatorname{cols}(x)\);
    \(\operatorname{retp}(x, \operatorname{zeros}(r, c))\);
endp;
```

```
proc (2) = cminv(xr,xi);
    local ixy, zr, zi;
    ixy = inv(xr)*xi;
    zr = inv(xr+xi*ixy); /* real part of inverse. */
    zi = -ixy*zr; /* imaginary part of inverse. */
    retp(zr,zi);
endp;
proc (2) = cmmult(xr,xi,yr,yi);
    local zr,zi;
    zr = xr*yr-xi*yi;
    zi = xr*yi+xi*yr;
    retp(zr,zi);
endp;
{ xr,xi } = cmcplx(rndn(3,3));
{ yr,yi } = cmcplx(rndn(3,3));
{ zr,zi } = cmmult( cminv(xr,xi),cminv(yr,yi) );
{ qr,qi } = cmmult( yr,yi,cminv(yr,yi) );
{ wr,wi } = cmmult(yr,yi,cminv(cmmult(cminv(xr,xi),yr,yi)));
```


### 12.7 Saving Compiled Procedures

When a file containing a procedure definition is run, the procedure is compiled and is then resident in memory. The procedure can be called as if it were an intrinsic function. If the new command is executed or you quit GAUSS and exit to the operating system, the compiled image of the procedure disappears and the file containing the procedure definition will have to be compiled again.

If a procedure contains no global references, that is, if it does not reference any global matrices or strings and it does not call any user-defined functions or procedures, it can be saved to disk in compiled form in a .fcg file with the save command, and loaded later with the loadp command
whenever it is needed. This will usually be faster than recompiling. For example:

```
save path = c:\gauss\cp proc1,proc2,proc3;
loadp path = c:\gauss\cp proc1,proc2,proc3;
```

The name of the file will be the same as the name of the procedure, with a. fcg extension. (For details, see loadp and save in the GAUSS Language Reference.)

All compiled procedures should be saved in the same subdirectory, so there is no question where they are located when it is necessary to reload them. The loadp path can be set in your startup file to reflect this. Then, to load in procedures, use
loadp proc1, proc2, proc3;

Procedures that are saved in . fcg files will NOT be automatically loaded. It is necessary to explicitly load them with loadp. This feature should be used only when the time necessary for the autoloader to compile the source is too great. Also, unless these procedures have been compiled with \#lineson, debugging will be more complicated.

## Sparse Matrices

## 13

The sparse matrix data type stores only the non-zero values of a 2-dimensional sparse matrix, which makes working with sparse matrices faster and more efficient.

### 13.1 Defining Sparse Matrices

The sparse matrix data type is strongly typed in GAUSS, which means that a variable must be defined as a sparse matrix variable before it may be used as such. Once a variable has been defined as a sparse matrix, it may not be used as another data type. Similarly, once a variable has been used as a matrix, array, or other non-sparse data type, it may not be redefined as a sparse matrix.

To define a global sparse matrix, you may use either the declare or the let command:

```
declare sparse matrix sm1;
let sparse matrix sm1;
```

or the following implicit let statement:

```
sparse matrix sm1;
```

declare may be used to define multiple sparse matrices in a single statement:

```
declare sparse matrix sm1, sm2, sm3;
```

To define a local sparse matrix inside of a procedure, use an implicit let statement:

```
sparse matrix lsm1;
```

As neither let nor declare support the initialization of a sparse matrix at this time, you must initialize a sparse matrix with an assignment after defining it.

### 13.2 Creating and Using Sparse Matrices

Several new functions have been added to allow you to create and manipulate sparse matrices. These functions are:

| denseToSp | Converts a dense matrix to a sparse matrix. |
| :--- | :--- |
| denseToSpRE | Converts a dense matrix to a sparse matrix, using a relative epsilon. |
| packedToSp | Creates a sparse matrix from a packed matrix of non-zero values and <br> row and column indices. |
| spBiconjGradSol | Solves the system of linear equations $\mathrm{Ax}=\mathrm{b}$ using the biconjugate <br> gradient method. |
| spConjGradSol | Solves the system of linear equations $\mathrm{Ax}=\mathrm{b}$ for symmetric matrices <br> using the conjugate gradient method. |


| spCreate | Creates a sparse matrix from vectors of non-zero values, row <br> indices, and column indices. |
| :--- | :--- |
| spDenseSubmat | Returns a dense submatrix of sparse matrix. |
| spDiagRvMat | Inserts submatrices along the diagonal of a sparse matrix. |
| spEye | Creates a sparse identity matrix. |
| spGetNZE | Returns the non-zero values in a sparse matrix, as well as their <br> corresponding row and column indices. |
| spLDL | Computes the LDL decomposition of a symmetric sparse matrix A. |
| spLU | Computes the LU decomposition of a sparse matrix A with partial <br> pivoting. |
| spNumNZE | Returns the number of non-zero elements in a sparse matrix. |
| spOnes | Returns a sparse submatrix of sparse matrix. |
| spSubmat | Converts a sparse matrix to a dense matrix. |
| spToDense | Multiplies a sparse matrix transposed by a dense matrix. |
| spTrTDense | Multiplies a sparse matrix by a scalar. |
| spTScalar | Creates a sparse matrix containing no non-zero values. |

See Command Reference, Chapter 33, for detailed information on each command.

### 13.3 Sparse Support in Matrix Functions and Operators

Support for the sparse matrix data type has also been added to many matrix functions and operators. The following is a complete list of the matrix functions and operators that currently support the new sparse matrix type:

|  | .$/=$ | .$<=$ |
| :--- | :--- | :--- |
| $\sim$ | $==$ | abs |
| $\mid$ | .$==$ | cols |
| $*$ | $>$ | maxc |
| .$*$ | .$>$ | minc |
| + | $>=$ | print |
| - | .$>=$ | rows |
| $/$ | .$<$ | scalerr |
| .$/$ | $<=$ | show |
| $/=$ |  | type |

Indexing is also supported for sparse matrices, using the same syntax as matrix indexing.
Note that printing a sparse matrix results in a table of the non-zero values contained in the sparse matrix, followed by their corresponding row and column indices, respectively.

### 13.3.1 Return Types for Dyadic Operators

The types of the returns for the dyadic operators were decided on a case-by-case basis, using the following general principles:

1. The return type for dyadic operations on two dense arguments is always dense.
2. The return type for dyadic operations on two sparse arguments is always sparse unless the result is likely to be significantly less sparse than the sparse arguments.
3. The return type for dyadic operations on a dense argument and a sparse argument (regardless of order) is dense unless the return is likely to be at least as sparse as the sparse argument.

These general principles have led to the following decisions regarding return types (note that only the cases that are displayed in these tables have been implemented at this point):

## Element-by-Element Numeric Operators

Element-by-Element Addition

| Result | $=$ | Left | Operator | Right |
| :--- | :--- | :---: | :---: | :---: |
| dense | $=$ | sparse | + | dense |
| dense | $=$ | dense | + | dense |
| sparse | $=$ | sparse | + | sparse |
| dense | $=$ | dense | + | sparse |

Element-by-Element Subtraction

| Result | $=$ | Left | Operator | Right |
| :--- | :--- | :---: | :---: | :---: |
| dense | $=$ | sparse | - | dense |
| dense | $=$ | dense | - | dense |
| sparse | $=$ | sparse | - | sparse |
| dense | $=$ | dense | - | sparse |


| Element-by-Element Multiplication |  |  |  |  |
| :--- | :--- | :---: | :---: | :---: |
| Result | $=$ | Left | Operator | Right |
| sparse | $=$ | sparse | .$^{*}$ | dense |
| dense | $=$ | dense | .$^{*}$ | dense |
| sparse | $=$ | sparse | .$^{*}$ | sparse |
| sparse | $=$ | dense | .$^{*}$ | sparse |


| Element-by-Element Division |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
| Result | $=$ | Left | Operator | Right |
| sparse | $=$ | sparse | .$/$ | dense |
| dense | $=$ | dense | .$/$ | dense |
| dense | $=$ | sparse | ./ | sparse |
| dense | $=$ | dense | .$/$ | sparse |

## Other Numeric Operators

| Matrix Multiplication |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
| Result | $=$ | Left | Operator | Right |
| dense | $=$ | sparse | $*$ | dense |
| dense | $=$ | dense | $*$ | dense |
| sparse | $=$ | sparse | $*$ | sparse |


| Linear Solve |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
| Result | $=$ | Left | Operator | Right |
| dense | $=$ | dense | $/$ | dense |
| dense | $=$ | dense | $/$ | sparse |

Note that at this time, the dense $=$ dense $/$ sparse case is defined only for real data.
When either of its arguments are sparse, the / operator uses a tolerance to determine the result, which may be read or set using the sysstate function, case 39 . The default tolerance is $1 \mathrm{e}-14$.

## Relational Operators

Since the results of element-by-element 'dot' comparison operators depend largely on the kind of data inputted, there are both both dense-returning and sparse-returning versions of the dot comparison operators when one or both arguments is a sparse matrix. The regular dot comparison operators and their alphabetic counterparts always return dense matrices, and there is a new set of alphabetic dot comparison operators that all return sparse matrices:

| Element-by-Element Dot Comparison Operators |  |  |  |
| :--- | :--- | :--- | :---: |
| Operation | Dense-Returning |  | Sparse-Returning |
| Equal to | .$==$ | .eq | .speq |
| Not equal to | .$/=$ | .ne | .spne |
| Less than | .$<$ | .lt | .splt |
| Less than or equal to | .$<=$ | .$l e$ | .sple |
| Greater than | .$>$ | .$g t$ | .spgt |
| Greater than or equal to | .$>=$ | .$g e$ | .spge |

Since the element-by-element 'non-dot' comparison operators ( $==, /=,<,<=,>,>=$ ) and their alphabetic counterparts (eq, ne, lt, le, gt, ge) all return scalars, there are no sparse-returning versions of them.

## Other Matrix Operators

| Horizontal Concatenation |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
| Result | $=$ | Left | Operator | Right |
| dense | $=$ | dense | $\sim$ | dense |
| sparse | $=$ | sparse | $\sim$ | sparse |
|  |  |  |  |  |
| Vertical Concatenation |  |  |  |  |
| Result | $=$ | Left | Operator | Right |
| dense | $=$ | dense | $\mid$ | dense |
| sparse | $=$ | sparse | $\mid$ | sparse |



## N-Dimensional Arrays

## 14

In GAUSS, internally, matrices and arrays are separate data types. Matrices, which are 2 -dimensional objects, are stored in memory in row major order. Therefore, a $3 \times 2$ matrix is stored as follows:

$$
[1,1] \quad[1,2] \quad[2,1] \quad[2,2] \quad[3,1] \quad[3,2]
$$

The slowest moving dimension in memory is indexed on the right, and the fastest moving dimension is indexed on the left. This is true of N -dimensional arrays as well. A $4 \times 3 \times 2$ array is stored in the following way:

| $[1,1,1]$ | $[1,1,2]$ | $[1,2,1]$ | $[1,2,2]$ | $[1,3,1]$ | $[1,3,2]$ |
| :--- | :--- | :--- | :--- | :--- | :--- |
| $[2,1,1]$ | $[2,1,2]$ | $[2,2,1]$ | $[2,2,2]$ | $[2,3,1]$ | $[2,3,2]$ |
| $[3,1,1]$ | $[3,1,2]$ | $[3,2,1]$ | $[3,2,2]$ | $[3,3,1]$ | $[3,3,2]$ |
| $[4,1,1]$ | $[4,1,2]$ | $[4,2,1]$ | $[4,2,2]$ | $[4,3,1]$ | $[4,3,2]$ |

A complex N -dimensional array is stored in memory in the same way. Like complex matrices, complex arrays are stored with the entire real part first, followed by the entire imaginary part.

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Every N -dimensional array has a corresponding $\mathrm{N} \times 1$ vector of orders that contains the sizes of each dimension of the array. This is stored with the array and can be accessed with getorders. The first element of the vector of orders corresponds to the slowest moving dimension, and the last element corresponds to the fastest moving dimension (refer to the sectionnameGlossary of Terms at the end of the chapter for clear definitions of these terms). The vector of orders for a $6 \times 5 \times 4 \times 3 \times 2$ array, which has 5 dimensions, is the following $5 \times 1$ vector:

Two terms that are important in working with N -dimensional arrays are "dimension index" and "dimension number." A dimension index specifies a dimension based on indexing the vector of orders. It is a scalar, 1-to-N, where 1 corresponds to the dimension indicated by the first element of the vector of orders of the array (the slowest moving dimension) and N corresponds to the dimension indicated by the last element of the vector of orders (the fastest moving dimension).

A dimension number specifies dimensions by numbering them in the same order that one would add dimensions to an array. In other words, the dimensions of an N -dimensional array are numbered such that the fastest moving dimension has a dimension number of 1 , and the slowest moving dimension has a dimension number of N .

A $6 \times 5 \times 4 \times 3 \times 2$ array has 5 dimensions, so the first element of the vector of orders (in this case, 6) refers to the size of dimension number 5 . Since the index of this element in the vector of orders is 1 , the dimension index of the corresponding dimension (dimension number 5) is also 1 .

You will find references to both dimension index and dimension number in the documentation for the functions that manipulate arrays.

There are a number of functions that have been designed to manipulate arrays. These functions allow you to manipulate a subarray within the array by passing in a locator vector to index any subarray that comprises a contiguous block of memory within the larger block. A vector of indices of an N -dimensional array is a [1-to- N$] \times 1$ vector of base 1 indices into the array, where the first element corresponds to the first element in a vector of orders. An $N \times 1$ vector of indices locates the
scalar whose position is indicated by the indices. For a $4 \times 3 \times 2$ array $x$, the $3 \times 1$ vector of indices:

3
2
1
indexes the [3,2,1] element of $x$. A $2 \times 1$ vector of indices for this 3-dimensional example, references the 1 -dimensional array whose starting location is given by the indices.

Because the elements of the vector of indices are always in the same order (the first element of the vector of indices corresponds to the slowest moving dimension of the array, the second element to the second slowest moving dimension, and so on), each unique vector of indices locates a unique subarray.

In general, an $[\mathrm{N}-\mathrm{K}] \times 1$ vector of indices locates a K-dimensional subarray that begins at the position indicated by the indices. The sizes of the dimensions of the K-dimensional subarray correspond to the last K elements of the vector of orders of the N -dimensional array. For a $6 \times 5 \times 4 \times 3 \times 2$ array $y$, the $2 \times 1$ vector of indices:

2
5
locates the $4 \times 3 \times 2$ subarray in $y$ that begins at $[2,5,1,1,1]$ and ends at $[2,5,4,3,2]$.

### 14.1 Bracketed Indexing

Brackets '[ ]' can be used to index N -dimensional arrays in virtually the same way that they are used to index matrices. Bracketed indexing is slower than the convenience array functions, such as getarray and setarray; however, it can be used to index non-contiguous elements. In order to index an N -dimensional array with brackets, there must be N indices located within the brackets, where the first index corresponds to the slowest moving dimension of the array and the last index corresponds to the fastest moving dimension.

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For a $2 \times 3 \times 4$ array $\mathbf{x}$, such that
[1,1,1] through [1,3,4] =

| 1 | 2 | 3 | 4 |
| :--- | :--- | :--- | :--- |

$\begin{array}{llll}5 & 6 & 7 & 8\end{array}$
$\begin{array}{llll}9 & 10 & 11 & 12\end{array}$
[2,1,1] through $[2,3,4]=$
$\begin{array}{llll}13 & 14 & 15 & 16\end{array}$
$\begin{array}{llll}17 & 18 & 19 & 20\end{array}$
$\begin{array}{llll}21 & 22 & 23 & 24\end{array}$
$\mathbf{x}[1,2,3]$ returns a $1 \times 1 \times 1$ array containing the $[1,2,3]$ element of $\mathbf{x}:$

7
$x[., 3,2]$ returns a $2 \times 1 \times 1$ array containing

$$
10
$$

22
$x[2, \ldots, 14]$ returns a $1 \times 3 \times 2$ array containing
$13 \quad 16$
$17 \quad 20$
$21 \quad 24$

14-4
$\mathbf{x}[., 2,1: 3]$ returns a $2 \times 1 \times 3$ array containing
$5 \quad 6 \quad 7$
$17 \quad 18 \quad 19$

### 14.2 E×E Conformability

The following describes rules for $\mathrm{E} \times \mathrm{E}$ conformability of arrays for operators and functions with two or more arguments.

- Any N -dimensional array is conformable to a scalar.
- An array is conformable to a matrix only if the array has fewer than 3 dimensions, and the array and matrix follow the standard rules of $\mathrm{E} \times \mathrm{E}$ conformability.
- Two arrays are $\mathrm{E} \times \mathrm{E}$ conformable if they comply with one of the following requirements:
- The two arrays have the same number of dimensions, and each dimension has the same size.
- The two arrays have the same number of dimensions, and each of the $\mathrm{N}-2$ slowest moving dimensions has the same size. In this case, the 2 fastest moving dimensions of the arrays must follow the $\mathrm{E} \times \mathrm{E}$ comformability rules that apply to matrices.
- Both of the arrays have fewer than 3 dimensions, and they follow the $\mathrm{E} \times \mathrm{E}$ conformability rules that apply to matrices.


### 14.3 Glossary of Terms

dimensions The number of dimensions of an object.
vector of orders $\mathrm{N} \times 1$ vector of the sizes of the dimensions of an object, where N is the number of dimensions, and the first element corresponds to the slowest moving dimension.
vector of indices [1-to-N] $\times 1$ vector of indices into an array, where the first element corresponds to the first element in a vector of orders.
dimension number Scalar [1-to-N], where 1 corresponds to the fastest moving dimension and N to the slowest moving dimension.
dimension index Scalar [1-to-N], where 1 corresponds to the first element of the vector of orders or vector of indices.
locator $[1-$ to-N] $\times 1$ vector of indices into an array used by array functions to locate a contiguous block of the array.

## Working with Arrays

## 15

### 15.1 Initializing Arrays

The use of N-dimensional arrays in GAUSS is an additional tool for reducing development time and increasing execution speed of programs. There are multiple ways of handling N -dimensional arrays and using them to solve problems, and these ways sometimes have implications for a trade-off between speed of execution and development time. We will try to make this clear in this chapter.

The term "arrays" specifically refers to N-dimensional arrays and must not be confused with matrices. Matrices and arrays are distinct types even if in fact they contain identical information. Functions for conversion from one to the other are described below.

There are five basic ways of creating an array depending on how the contents are specified:
areshape $\quad$ Create array from specified matrix .
aconcat Create array from matrices and arrays.
aeye Create array of identity matrices.
arrayinit Allocate array filled with specified scalar value.
arrayalloc Allocate array with no specified contents.

### 15.1.1 areshape

areshape is a method for creating an array with specified contents. arrayinit creates an array filled with a selected scalar value: areshape will do the same, but with a matrix. For example, given a matrix, areshape will create an array containing multiple copies of that matrix:

```
x = reshape(seqa(1,1,4),2,2);
ord = 3 | 2 | 2;
a = areshape(x,ord);
print a;
    Plane [1,.,.]
    1.0000 2.0000
    3.00004.0000
    Plane [2,.,.]
    1.0000 2.0000
    3.0000 4.0000
    Plane [3,.,.]
    1.0000 2.0000
    3.0000 4.0000
```


## Reading Data from the Disk into an Array

areshape is a fast way to re-dimension a matrix or array already in memory. For example, suppose we have a GAUSS data set containing panel data and that it's small enough to be read in all at once:

```
panel = areshape(loadd("panel"),5|100|10);
mn = amean(panel,2); /* 5x1x10 array of means */
    /*of each panel */
mm = moment (panel,0); /* 5x10x10 array of moments */
    /* of each panel */
/*
** vc is a 5x10x10 array of
** covariance matrices
*/
vc = mm / 100 - amult(atranspose(mn,1|3|2),mn);
```

panel is a $5 \times 100 \times 10$ array, and in this context is 5 panels of 100 cases measured on 10 variables.

## Inserting Random Numbers into Arrays

A random array of any dimension or size can be quickly created using areshape. Thus, for a $10 \times 10 \times 5 \times 3$ array:

```
ord = { 10, 10, 5, 3 };
y = areshape(rndu(prodc(ord),1),ord);
```

The quick and dirty method above uses the linear congruential generator, which is fast but doesn't have the properties required for serious Monte Carlo work. For series simulation you will need to use the KM generator:

```
sdO = 345678;
ord = { 10, 10, 5, 3 };
{ z,sdQ } = rndKMu(prodc(ord),1,sdQ);
y = areshape(z,ord);
```


## Expanding a Matrix into an Array Vector of Matrices

For computing the log-likelihood of a variance components model of panel data, it is necessary to expand a $\mathrm{T} \times \mathrm{T}$ matrix into an $\mathrm{NT} \times \mathrm{T}$ array of these matrices. This is easily accomplished using areshape. For example:

```
m={ 1.0 0.3 0.2,
    0.3 1.0 0.1,
    0.2 0.1 1.0 };
r = areshape(m,3|3|3);
print r;
```

Plane [1,.,.]
$1.0000 \quad 0.3000 \quad 0.2000$
$0.3000 \quad 1.0000 \quad 0.1000$
$0.2000 \quad 0.1000 \quad 1.0000$
Plane [2,.,.]
$1.0000 \quad 0.3000 \quad 0.2000$
$0.3000 \quad 1.0000 \quad 0.1000$
$0.2000 \quad 0.10001 .0000$
Plane [3,.,.]
$1.0000 \quad 0.3000 \quad 0.2000$
$0.3000 \quad 1.0000 \quad 0.1000$
$0.2000 \quad 0.10001 .0000$

### 15.1.2 aconcat

aconcat creates arrays from conformable sets of matrices or arrays. With this function, contents are completely specified by the user. This example tries three concatenations, one along each
dimension:

```
rndseed 345678;
x1 = rndn(2,2);
x2 = arrayinit(2|2,1);
/*
** along the first dimension or rows
*/
a = aconcat(x1,x2,1);
print a;
    -0.4300 -0.2878 1.0000 1.0000
    -0.1327-0.0573 1.0000 1.0000
/*
** along the second dimension or columns
*/
a = aconcat(x1,x2,2);
print a;
    -0.4300 -0.2878
    -0.1327 -0.0573
    1.0000 1.0000
    1.0000 1.0000
/*
** along the third dimension
*/
a = aconcat(x1,x2,3);
print a;
    Plane [1,.,.]
```

$$
\begin{aligned}
& \begin{aligned}
-0.4300 & -0.2878 \\
-0.1327 & -0.0573
\end{aligned} \\
& \text { Plane }[2, \ldots, .] \\
& 1.0000 \\
& 1.0000 \\
& 1.0000
\end{aligned}
$$

### 15.1.3 aeye

aeye creates an array in which the principal diagonal of the two trailing dimensions is set to one. For example:

```
ord = 2 | 3 | 3;
a = aeye(ord);
print a;
    Plane [1,.,.]
    1.00000 0.00000 0.00000
    0.00000 1.00000 0.00000
    0.00000 0.00000 1.00000
    Plane [2,.,.]
        1.00000 0.00000 0.00000
        0.00000 1.00000 0.00000
        0.00000 0.00000 1.00000
```


### 15.1.4 arrayinit

arrayinit creates an array with all elements set to a specified value. For example:

$$
\text { ord }=3|2| 3 ;
$$

```
a = arrayinit(ord,1);
print a;
Plane [1,.,.]
    1.0000 1.0000 1.0000
    1.0000 1.0000 1.0000
Plane [2,.,.]
    1.0000 1.0000 1.0000
    1.0000 1.0000 1.0000
Plane [3,.,.]
    1.0000 1.0000 1.0000
    1.0000 1.0000 1.0000
```


### 15.1.5 arrayalloc

arrayalloc creates an array with specified number and size of dimensions without setting elements to any values. This requires a vector specifying the order of the array. The length of the vector determines the number of dimensions, and each element determines the size of the corresponding dimensions. The array will then have to be filled using any of several methods described later in this chapter.

For example, to allocate a $2 \times 2 \times 3$ array:

```
rndseed 345678;
ord = 3 | 2 | 2;
a = arrayalloc(ord,0);
for i(1,ord[1],1);
    a[i,.,.] = rndn(2,3);
endfor;
```

```
print a;
    Plane [1,.,.]
        -0.4300 -0.2878 -0.1327
        -0.0573 -1.2900 0.2467
Plane [2,.,.]
        -1.4249 -0.0796 1.2693
        -0.7530 -1.7906 -0.6103
    Plane [3,.,.]
        1.2586 -0.4773 0.7044
        -1.2544 0.5002 0.3559
```

The second argument in the call to arrayalloc specifies whether the created array is real or complex. arrayinit creates only real arrays.

### 15.2 Assigning to Arrays

There are three methods used for assignment to an array:

| index operator | The same method as matrices, generalized to arrays. |
| :--- | :--- |
| putArray | Put a subarray into an N-dimensional array and returns the result. |
| setArray | Set a subarray of an N-dimensional array in place. |

And there are several ways to extract parts of arrays:
index operator The same method as matrices, generalized to arrays.

| getArray | Get a subarray from an array. |
| :--- | :--- |
| getMatrix | Get a matrix from an array. |
| getMatrix4D | Get a matrix from a 4-dimensional array. |
| getScalar4D | Get a scalar from a 4-dimensional array. |

The index operator is the slowest way to extract parts of arrays. The specialized functions are the fastest when the circumstances are appropriate for their use.

### 15.2.1 index operator

The index operator will put a subarray into an array in a manner analogous to the use of index operators on matrices:

```
a = arrayinit(3|2|2,0);
b = arrayinit(3|1|2,1);
a[.,2,.] = b;
print a;
    Plane [1,.,.]
    0.00000 0.00000
    1.0000 1.0000
    Plane [2,.,.]
    0.00000 0.00000
    1.0000 1.0000
    Plane [3,.,.]
    0.00000 0.00000
    1.0000 1.0000
```


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As this example illustrates, the assignment doesn't have to be contiguous. putMatrix and setMatrix require a contiguous assignment, but for that reason they are faster.

The right hand side of the assignment can also be a matrix:

```
a[1,.,.] = rndn(2,2);
print a;
    Plane [1,.,.]
        -1.7906502 -0.61038103
            1.2586160-0.47736360
Plane [2,.,.]
            0.00000 0.00000
            1.00000 1.00000
            Plane [3,.,.]
            0.00000 0.00000
            1.00000 1.00000
```

The index operator will extract an array from a subarray in a manner analogous to the use of index operators on matrices:

```
a = areshape(seqa(1,1,12),3|2|2);
b = a[.,1,.];
print a;
    Plane [1,.,.]
    1.0000 2.0000
    3.0000 4.0000
```

```
    Plane [2,.,.]
    5.0000 6.0000
        7.0000 8.0000
        Plane [3,.,.]
        9.0000 10.000
        11.000 12.000
print b;
Plane [1,.,.]
    1.0000 2.0000
Plane [2,.,.]
    5.0000 6.0000
Plane [3,.,.]
    9.0000 10.000
```

It is important to note that the result is always an array even if it's a scalar value:

```
c = a[1,1,1];
print c;
    Plane [1,.,.]
    1.0000
```

If you require a matrix result, and if the result has one or two dimensions, use arraytomat to convert to a matrix, or use getMatrix, or getMatrix4D. Or, if the result is a scalar, use getScalar3D or getScalar4D.

### 15.2.2 getArray

getArray is an additional method for extracting arrays:

```
a = areshape(seqa(1,1,12),3|2|2);
b = getarray(a,2|1);
print a;
    Plane [1,.,.]
            1.0000 2.0000
            3.0000 4.0000
            Plane [2,.,.]
            5.0000 6.0000
            7.0000 8.0000
            Plane [3,.,.]
            9.0000 10.000
            11.000 12.000
print b;
    5.0000 6.0000
```

getArray can only extract a contiguous part of an array. To get non-contiguous parts you must use the index operator.

### 15.2.3 getMatrix

If the result is one or two dimensions, getMatrix returns a portion of an array converted to a matrix. getMatrix is about 20 percent faster than the index operator:

```
a = areshape(seqa(1,1,12),3|2|2);
b = getMatrix(a,2);
print b;
    5.0000 6.0000
    7.0000 8.0000
```


### 15.2.4 getMatrix4D

This is a specialized version of getMatrix for 4-dimensional arrays. It behaves just like getMatrix but is dramatically faster for that type of array. The following illustrates the difference in timing:

```
a = arrayinit(100|100|10|10,1);
tQ = date;
for i(1,100,1);
    for j(1,100,1);
        b = a[i,j,.,.];
    endfor;
endfor;
t1 = date;
e1 = ethsec(t0,t1);
print e1;
print;
t2=date;
for i(1,100,1);
```

```
        for j(1,100,1);
        b = getMatrix4d(a,i,j);
        endfor;
endfor;
t3 = date;
e2 = ethsec(t2,t3);
print e2;
print;
print ftostrC(100*((e1-e2)/e1),
    "percent difference - %6.2lf%%");
    13.000000
    5.0000000
percent difference - 61.54%
```


### 15.2.5 getScalar3D, getScalar4D

These are specialized versions of getMatrix for retrieving scalar elements of 3-dimensional and 4-dimensional arrays, respectively. They behave just like getMatrix, with scalar results, but are much faster. For example:

```
a = arrayinit(100|10|10,1);
tQ = date;
for i(1,100,1);
    for j(1,10,1);
        for k(1, 10, 1);
            b = a[i,j,k];
        endfor;
    endfor;
endfor;
```

```
t1 = date;
e1 = ethsec(t0,t1);
print e1;
print;
t2=date;
for i(1,100,1);
    for j(1,10,1);
        for k(1, 10,1);
            b = getscalar3d(a,i,j,k);
            endfor;
    endfor;
endfor;
t3 = date;
e2 = ethsec(t2,t3);
print e2;
print;
print ftostrC(100*((e1-e2)/e1),
    "percent difference - %6.2lf%%");
    7.0000000
    2.0000000
percent difference - 71.43%
```


### 15.2.6 putArray

putArray enters a subarray, matrix, or scalar into an N-dimensional array and returns the result in an array. This function is much faster than the index operator, but it requires the part of the array being assigned to be contiguous:

```
a = arrayinit(3|2|2,3);
b = putarray(a,2,eye(2));
```

```
print b;
    Plane [1,.,.]
        3.0000 3.0000
        3.0000 3.0000
    Plane [2,.,.]
        1.0000 0.00000
        0.00000 1.0000
    Plane [3,.,.]
        3.0000 3.0000
        3.0000 3.0000
```


### 15.2.7 setArray

setArray enters a subarray, matrix, or scalar into an N-dimensional array in place:

```
a = arrayinit(3|2|2,3);
setarray a,2,eye(2);
print b;
    Plane [1,.,.]
    3.0000 3.0000
        3.0000 3.0000
    Plane [2,.,.]
    1.0000 0.0000
    0.0000 1.0000
```

```
Plane [3,.,.]
3.0000 3.0000
3.0000 3.0000
```


### 15.3 Looping with Arrays

When working with arrays, for loops and do loops may be used in the usual way. In the following, let $\mathbf{Y}$ be an $\mathrm{N} \times 1 \times \mathrm{L}$ array of L time series, $\mathbf{X}$ an $\mathrm{N} \times 1 \times \mathrm{K}$ array of K independent variables, $\mathbf{B}$ a $K \times L$ matrix of regression coefficients, phi a $\mathrm{P} \times \mathrm{L} \times \mathrm{L}$ array of garch coefficients, theta a $\mathrm{Q} \times \mathrm{L} \times \mathrm{L}$ array of arch coefficients, and omega a $\mathrm{L} \times \mathrm{L}$ symmetric matrix of constants. The log-likelihood for a multivariate garch BEKK model can be computed using the index operator:

```
yord = getOrders(Y);
xord = getOrders(X);
gord = getOrders(phi);
aord = getOrders(theta);
N = yord[1]; /* No. of observations */
L = yord[3]; /* No. of time series */
K = xord[3]; /* No. of independent variables */
        /* in mean equation */
P = gord[1]; /* order of garch parameters */
Q = aord[1]; /* order of arch parameters */
r = maxc(P|Q);
E = Y - amult(X,areshape(B,N|K|L));
sigma = areshape(omega,N|L|L);
for i(r+1,N,1);
    for j(1,Q,1);
        W = amult(theta[j,.,.],
        atranspose(E[i-j,.,.],1|3|2));
        sigma[i,.,.] = sigma[i,.,.] + amult(W,atranspose(W,1|3|2));
```

endfor;
for $\mathrm{j}(1, \mathrm{P}, 1)$;
sigma[i,.,.] $=\operatorname{sigma[i,.,.]~+~amult(amult(phi[j,.,.],~}$ sigma[i-j,.,.]),phi[j,.,.]);
endfor;
endfor;
sigmai = invpd(sigma);
lndet $=\ln (\operatorname{det}($ sigma $)) ;$
$\ln 1=-0.5 *(\mathrm{~L} *(\mathrm{~N}-\mathrm{R}) * \operatorname{asum}(\ln (\operatorname{det}($ sigmai $)), 1)+$ asum(amult(amult(E, sigmai), atranspose(E, 1|3|2)), 3);

Instead of index operators, the above computation can be done using getArray and setArray:

```
yord = getOrders(Y);
xord = getOrders(X);
gord = getOrders(phi);
aord = getOrders(theta);
N = yord[1]; /* No. of observations */
L = yord[3]; /* No. of time series */
K = xord[3]; /* No. of independent variables */
        /* in mean equation */
P = gord[1]; /* order of garch parameters */
Q = aord[1]; /* order of arch parameters */
r = maxc(P|Q);
E = Y - amult(X,areshape(B,N|K|L));
sigma = areshape(omega,N|L|L);
for i(r+1,N,1);
    for j(1,Q,1);
        W = amult(getArray(theta,j),
        atranspose(getArray(E,i-j),2|1));
        setarray sigma,i,getArray(sigma,i)+
```

```
    amult(W,atranspose(W,2|1));
endfor;
for j(1,P,1);
        setarray sigma,i,getArray(sigma,i)+
            areshape(amult(amult(getArray(phi,j),
            getArray(sigma,i-j)),getArray(phi,j)),3|3);
        endfor;
endfor;
sigmai = invpd(sigma);
lndet = ln(det(sigma));
lnl = -0.5*( L*(N-R)*asum(ln(det(sigmai)),1)+
    asum(amult(amult(E,sigmai),atranspose(E,1|3|2)),3)
```

Putting the two code fragments above into loops that called them a hundred times and measuring the time, produced the following results:
index operator: 2.604 seconds
getArray, setArray: 1.092 seconds

Thus, the getArray and setArray methods are more than twice as fast.

### 15.3.1 loopnextindex

Several keyword functions are available in GAUSS for looping with arrays. The problem in the previous section, for example, can be written using these functions rather than with for loops:

```
sigind = r + 1;
sigloop:
```

```
    sig0ind = sigind[1];
    thetaind = 1;
thetaloop:
    sig0ind = sig0ind - 1;
    W = amult(getArray(theta,thetaind),
        atranspose(getArray(E,sig0ind),2|1));
    setarray sigma,sigind,getArray(sigma,sigind)+
        amult(W,atranspose(W,2|1));
    loopnextindex thetaloop,thetaind,aord;
    sig0ind = sigind;
    phiind = 1;
philoop:
    sig0ind[1] = sig0ind[1] - 1;
    setarray sigma,sigind,getArray(sigma,sigind)+
        areshape(amult(amult(getArray(phi,phiind),
        getArray(sigma,sig0ind)),
        getArray(phi,phiind)),3|3);
    loopnextindex philoop,phiind,gord;
    loopnextindex sigloop,sigind,sigord;
```

The loopnextindex function in this example isn't faster than the for loop used in the previous section primarily because the code is looping only through the first dimension in each loop. The advantages of loopnextindex, previousindex, nextindex, and walkindex are when the code is looping through the higher dimensions of a highly dimensioned array. In this case, looping through an array can be very complicated and difficult to manage using for loops.
loopnextindex can be faster and more useful.
The next example compares two ways of extracting a subarray from a 5 -dimensional array:

```
ord = 3|3|3|3|3;
a = areshape(seqa(1,1,prodc(ord)),ord);
b = eye(3);
for i(1,3,1);
```

```
        for j(1,3,1);
        for k(1,3,1);
            setarray a,i|j|k,b;
            endfor;
        endfor;
endfor;
ind = { 1,1,1 };
loopi:
    setarray a,ind,b;
    loopnextindex loopi,ind,ord;
```

Calling each loop 10,000 times and measuring the time each takes, we get
for loop: 1.171 seconds
loopnextindex: . 321 seconds

In other words, loopnextindex is about four times faster, a very significant difference.

### 15.4 Miscellaneous Array Functions

### 15.4.1 atranspose

This function changes the order of the dimensions. For example:

```
a = areshape(seqa(1,1,12),2|3|2);
print a;
```

    Plane [1,.,.]
    ```
        1.0000 2.0000
        3.0000 4.0000
        5.0000 6.0000
    Plane [2,.,.]
        7.0000 8.0000
        9.0000 10.000
        11.000 12.000
/*
** swap 2nd and 3rd dimension
*/
print atranspose(a,1|3|2);
    Plane [1,.,.]
        1.0000 3.0000 5.0000
        2.0000 4.0000 6.0000
        Plane [2,.,.]
        7.0000 9.0000 11.000
        8.0000 10.000 12.000
/*
** swap 1st and 3rd dimension
*/
print atranspose(a,3|2|1);
    Plane [1,.,.]
    1.0000 7.0000
    3.0000 9.0000
    5.0000 11.000
```

```
            Plane [2,.,.]
            2.0000 8.0000
            4.0000 10.000
            6.0000 12.000
/*
** move 3rd into the front
*/
print atranspose(a,3|1|2);
    Plane [1,.,.]
            1.0000 3.0000 5.0000
            7.0000 9.0000 11.000
            Plane [2,.,.]
            2.0000 4.0000 6.0000
            8.0000 10.000 12.000
```


### 15.4.2 amult

This function performs a matrix multiplication on the last two trailing dimensions of an array. The leading dimensions must be strictly conformable, and the last two trailing dimensions must be conformable in the matrix product sense. For example:

```
\(a=\operatorname{areshape}(\operatorname{seqa}(1,1,12), 2|3| 2)\);
b = areshape(seqa \((1,1,16), 2|2| 4)\);
c = amult(a,b);
print a;
```

```
Plane [1,.,.]
```

```
        1.0000 2.0000
        3.0000 4.0000
        5.0000 6.0000
        Plane [2,.,.]
        7.0000 8.0000
        9.0000 10.000
        11.000 12.000
print b;
    Plane [1,.,.]
    1.0000 2.0000 3.0000 4.0000
    5.0000 6.0000 7.0000 8.0000
    Plane [2,.,.]
    9.0000 10.000 11.000 12.000
    13.000 14.000 15.000 16.000
print c;
    Plane [1,.,.]
    11.000 14.000 17.000 20.000
    23.000 30.000 37.000 44.000
    35.000 46.000 57.000 68.000
    Plane [2,.,.]
        167.00 182.00 197.00 212.00
        211.00 230.00 249.00 268.00
        255.00 278.00 301.00 324.00
```

Suppose we have a matrix of data sets, a $2 \times 2$ matrix of $100 \times 5$ data sets that we've stored in a $2 \times 2 \times 100 \times 5$ array called $\mathbf{x}$. The moment matrices of these data sets can easily and quickly be computed using atranspose and amult:

```
vc = amult(atranspose(x,1|2|4|3),x);
```


### 15.4.3 amean, amin, amax

These functions compute the means, minimums, and maximums, respectively, across a dimension of an array. The size of the selected dimension of the resulting array is shrunk to one and contains the means, minimums, or maximums depending on the function called. For example:

```
a = areshape(seqa(1,1,12),2|3|2);
print a;
    Plane [1,.,.]
        1.0000 2.0000
        3.0000 4.0000
        5.0000 6.0000
    Plane [2,.,.]
        7.0000 8.0000
        9.0000 10.000
        11.000 12.000
/*
** compute means along third dimension
*/
print amean(a,3);
```

    Plane [1,.,.]
    ```
        4.0000 5.0000
        6 . 0 0 0 0 ~ 7 . 0 0 0 0 ~
        8.0000 9.0000
/*
** print means along the second dimension, i.e.,
    down the columns
*/
print amean(a,2);
    Plane [1,.,.]
            3.00004.0000
        Plane [2,.,.]
            9.0000 10.000
/*
** print the minimums down the columns
*/
print amin(a,2);
        Plane [1,.,.]
        1.0000 2.0000
        Plane [2,.,.]
        7.0000 8.0000
/*
** print the maximums along the third dimension
*/
```

```
print amax(a,3);
Plane [1,.,.]
7.0000 8.0000
9.0000 10.000
11.000 12.000
```


### 15.4.4 getDims

This function returns the number of dimensions of an array:

```
a = arrayinit(4|4|5|2,0);
print getdims(a);
```

4.00

### 15.4.5 getOrders

This function returns the sizes of each dimension of an array. The length of the vector returned by getOrders is the dimension of the array:

```
a = arrayinit(4|4|5|2,0);
print getOrders(a);
```

4.00
4.00
5.00
2.00

### 15.4.6 arraytomat

This function converts an array with two or fewer dimensions to a matrix:

```
a = arrayinit(2|2,0);
b = arraytomat(a);
type(a);
    21.000
type(b);
    6.0000
```


### 15.4.7 mattoarray

This function converts a matrix to an array:

```
b = rndn(2,2);
a = mattoarray(b);
type(b);
    6 . 0 0 0 0
type(a);
```

    21.000
    
### 15.5 Using Arrays with GAUSS functions

Many of the GAUSS functions have been re-designed to work with arrays. There are two general approaches to this implementation. There are exceptions, however, and you are urged to refer to the documention if you are not sure how a particular GAUSS function handles array input.

In the first approach, the function returns an element-by-element result that is strictly conformable to the input. For example, cdfnc returns an array of identical size and shape to the input array:

```
a = areshape(seqa(-2,.5,12),2|3|2);
b = cdfnc(a);
print b;
    Plane [1,.,.]
    0.9772 0.9331
    0.8413 0.6914
    0.5000 0.3085
    Plane [2,.,.]
    0.1586 0.0668
    0.0227 0.0062
    0.0013 0.0002
```

In the second approach, which applies generally to GAUSS matrix functions, the function operates on the matrix defined by the last two trailing dimensions of the array. Thus, given a $5 \times 10 \times 3$ array, moment returns a $5 \times 3 \times 3$ array of five moment matrices computed from the five $10 \times 3$ matrices in the input array.

Only the last two trailing dimensions matter; i.e., given a $2 \times 3 \times 4 \times 5 \times 10 \times 6$ array, moment returns a $2 \times 3 \times 4 \times 5 \times 6 \times 6$ array of moment matrices.

For example, in the following the result is a $2 \times 3$ array of $3 \times 1$ vectors of singular values of a $2 \times 3$ array of $6 \times 3$ matrices:

```
a = areshape(seqa(1,1,108),2|3|6|3);
b=svds(a);
print b;
```

```
Plane [1,1,.,.]
            45.894532
            1.6407053
            1.2063156e-015
Plane [1,2,.,.]
            118.72909
            0.63421188
            5.8652600e-015
Plane [1,3,.,.]
            194.29063
            0.38756064
            1.7162751e-014
Plane [2,1,.,.]
            270.30524
            0.27857175
            1.9012118e-014
Plane [2,2,.,.]
            346.47504
            0.21732995
    1.4501098e-014
Plane [2,3,.,.]
            422.71618
            0.17813229
        1.6612287e-014
```

It might be tempting to conclude from this example that, in general, a GAUSS function's behavior
on the last two trailing dimensions of an array is strictly analogous to the GAUSS function's behavior on a matrix. This may be true with some of the functions, but not all. For example, the GAUSS meanc function returns a column result for matrix input. However, the behavior for the GAUSS amean function is not analogous. This function takes a second argument that specifies on which dimension the mean is to be taken. That dimension is then collapsed to a size of 1 . Thus:

```
a = areshape(seqa(1,1,24),2|3|4);
print a;
    Plane [1,.,.]
        1.000 2.000 3.000 4.000
        5.000 6.000 7.000 8.000
        9.000 10.000 11.000 12.000
    Plane [2,.,.]
        13.000 14.000 15.000 16.000
        17.000 18.000 19.000 20.000
        21.000 22.000 23.000 24.000
/*
** means computed across rows
*/
b = amean(a,1);
print b;
    Plane [1,.,.]
        2.500
        6.500
        10.500
    Plane [2,.,.]
```

    14.500
    ```
        18.500
        22.500
/*
** means computed down columns
*/
c = amean(a,2);
print c;
        Plane [1,.,.]
            5.000 6.000 7.000 8.000
        Plane [2,.,.]
            17.000 18.000 19.000 20.000
/*
** means computed along 3rd dimension
*/
d = amean(a,3);
print d;
        Plane [1,.,.]
            7.000 8.000 9.000 10.000
            11.000 12.000 13.000 14.000
            15.000 16.000 17.000 18.000
```


### 15.6 A Panel Data Model

Suppose we have N cases observed at T times. Let $y_{i t}$ be an observation on a dependent variable for the $i^{t h}$ case at time $t, X_{i t}$ an observation of $k$ independent variables for the $i^{t h}$ case at time $t, B$, a
$K \times 1$ vector of coefficients. Then

$$
y_{i t}=X_{i t} B+\mu_{i}+\epsilon_{i t}
$$

is a variance components model where $\mu_{i}$ is a random error term uncorrelated with $\epsilon_{i t}$, but which is correlated within cases. This implies an $\mathrm{NT} \times \mathrm{NT}$ residual moment matrix that is block diagonal with $\mathrm{N} \mathrm{T} \times \mathrm{T}$ moment matrices with the following form:

$$
\left[\begin{array}{cccc}
\sigma_{\mu}^{2}+\sigma_{\epsilon}^{2} & \sigma^{2} & \ldots & \sigma_{\mu}^{2} \\
\sigma_{\mu}^{2} & \sigma_{\mu}^{2}+\sigma_{\epsilon}^{2} & \ldots & \sigma_{\mu}^{2} \\
\vdots & \vdots & \vdots & \vdots \\
\sigma_{\mu}^{2} & \sigma_{\mu}^{2} & \ldots & \sigma_{\mu}^{2}+\sigma_{\epsilon}^{2}
\end{array}\right]
$$

The log-likelihood for this model is

$$
\ln L=-0.5\left(N T \ln (2 \pi)-\ln |\Omega|+(Y-X B) / \Omega^{-1}(Y-X B)\right)
$$

where $\Omega$ is the block-diagonal moment matrix of the residuals.

## Computing the Log-likelihood

Using GAUSS arrays, we can compute the log-likelihood of this model without resorting to do loops. Let $\mathbf{Y}$ be a $100 \times 3 \times 1$ array of observations on the dependent variable, and $\mathbf{X}$ a $100 \times 3 \times 5$ array of observations on the independent variables. Further let B be a $5 \times 1$ vector of coefficients, and sigu and sige be the residual variances of $\mu$ and $\epsilon$ respectively. Then, in explicit steps we compute

$$
\begin{aligned}
& \mathrm{N}=100 ; \\
& \mathrm{T}=3 ; \\
& \mathrm{K}=5 ;
\end{aligned}
$$

```
sigma = sigu * ones(T,T) + sige * eye(T); /* TxT sigma */
sigmai = invpd(sigma); /* sigma inverse */
lndet = N*ln(detl);
E = Y - amult(X,areshape(B,N|K|1)); /* residuals */
Omegai = areshape(sigmai,N|T|T); /* diagonal blocks */
    /* stacked in a vector array */
R1 = amult(atranspose(E,1|3|2),Omegai); /* E'Omegai */
R2 = amult(R1,E); /* R1*E */
lnL = -0.5*(N*T*ln(2*pi) - lndet + asum(R2,3)); /* log-likelhood */
```

All of this can be made more efficient by nesting statements, which eliminates copying of temporary intervening arrays to local arrays. It is also useful to add a check for the positive definiteness of sigma:

```
N = 100;
T = 3;
K = 5;
const = -0.5*N*T*\operatorname{ln}(2*pi);
oldt = trapchk(1);
trap 1,1;
sigmai = invpd(sigu*ones(T,T)+sige*eye(T));
trap oldt,1;
if not scalmiss(sigmai);
    E = Y - amult(X,areshape(B,N|K|1));
    lnl = const + 0.5*N*ln(detl)-
        0.5*asum(amult(amult(atranspose(E,1|3|2),
        areshape(sigmai,N|T|T)),E),3);
else;
    lnl = error(0);
endif;
```


### 15.7 Appendix

This is an incomplete list of special functions for working with arrays. Many GAUSS functions have been modified to handle arrays and are not listed here. For example, cdfnc computes the complement of the Normal cdf for each element of an array just as it would for a matrix. See the documentation for these GAUSS functions for information about their behavior with arrays.

| aconcat | Concatenate conformable matrices and arrays in a user-specified <br> dimension. |
| :--- | :--- |
| aeye | Create an array of identity matrices. |
| amax | Compute the maximum elements across a dimension of an array. |
| amean | Compute the mean along one dimension of an array. |
| amin | Compute the minimum elements across a dimension of an array. |
| amult | Perform a matrix multiplication on the last two trailing dimensions <br> of an array. |
| areshape | Reshape a scalar, matrix, or array into an array of user-specified size. |
| arrayalloc | Create an N-dimensional array with unspecified contents. |
| arrayinit | Create an N-dimensional array with a specified fill value. |
| arraytomat | Change an array to type matrix. |
| asum | Compute the sum across one dimension of an array. |
| atranspose | Transpose an N-dimensional array. |
| getarray | Get a contiguous subarray from an N-dimensional array. |
| getdims | Get the number of dimensions in an array. |
| getmatrix | Get a contiguous matrix from an N-dimensional array. |
| getmatrix4D | Get a contiguous matrix from a 4-dimensional array. |
| getorders | Get the vector of orders corresponding to an array. |

getscalar3D Get a scalar from a 3-dimensional array.
getscalar4D Get a scalar form a 4-dimensional array.
loopnextindex Increment an index vector to the next logical index and jump to the specified label if the index did not wrap to the beginning.
mattoarray Change a matrix to a type array.
nextindex Return the index of the next element or subarray in an array.
previousindex Return the index of the previous element or subarray in an array.
putarray Put a contiguous subarray into an N-dimensional array and return the resulting array.
setarray Set a contiguous subarray of an N-dimensional array.
walkindex Walk the index of an array forward or backward through a specified dimension.

## Structures

## 16

### 16.1 Basic Structures

### 16.1.1 Structure Definition

The syntax for a structure definition is

```
struct A { /* list of members */ };
```

The list of members can include scalars, arrays, matrices, strings, and string arrays, as well as other structures. As a type, scalars are unique to structures and don't otherwise exist.

For example, the following defines a structure containing the possible contents:

```
struct generic_example {
    scalar x;
    matrix y;
```

```
string s1;
string array s2
struct other_example t;
};
```

A useful convention is to put the structure definition into a file with a . sdf extension. Then, for any command file or source code file that requires this definition, put

## \#include filename.sdf

For example:

```
#include example.sdf
```

These statements create structure definitions that persist until the workspace is cleared. They do not create structures, only structure-type definitions. The next section describes how to create an instance of a structure.

### 16.1.2 Declaring an Instance

To use a structure, it is necessary to declare an instance. The syntax for this is
struct structure_type structure_name;

For example:

```
#include example.sdf
struct generic_example p0;
```


### 16.1.3 Initializing an Instance

Members of structures are referenced using a "dot" syntax:

```
p0.x = rndn(20,3);
```

The same syntax applies when referred to on the right-hand side:

```
mn = meanc(p0.x);
```


## Initialization of Global Structures

Global structures are initialized at compile time. Each member of the structure is initialized according to the following schedule:

| scalar | 0, a scalar zero |
| :--- | :--- |
| matrix | , an empty matrix with zero rows and zero columns |
| array | 0, a 1 -dimensional array set to zero |
| string | '", a null string |
| string array | $\prime "$, a $1 \times 1$ string array set to null |

If a global already exists in memory, it will not be reinitialized. It may be the case in your program that when it is rerun, the global variables may need to be reset to default values. That is, your program may depend on certain members of a structure being set to default values that are set to some other value later in the program. When you rerun this program, you will want to reinitialize the global structure. To do this, make an assignment to at least one of the members. This can be made convenient by writing a procedure that declares a structure and initializes one of its members to a default value, and then returns it. For example:

```
/* ds.src */
#include ds.sdf
proc dsCreate;
```

```
        struct DS dO;
        d0.dataMatrix = 0;
        retp(dQ);
endp;
```

Calling this function after declaring an instance of the structure will ensure initialization to default values each time your program is run:

```
struct DS d0;
dQ = dsCreate;
```


## Initializing Local Structures

Local structures, which are structures defined inside procedures, are initialized at the first assignment. The procedure may have been written in such a way that a subset of structures are used an any one call, and in that case time is saved by not initializing the unused structures. They will be initialized to default values only when the first assignment is made to one of its members.

### 16.1.4 Arrays of Structures

To create a matrix of instances, use the reshape command:

```
#include ds.sdf
struct DS p0;
p0 = reshape(dsCreate,5,1);
```

This creates a $5 \times 1$ vector of instances of DS structures, with all of the members initialized to default values.

When the instance members have been set to some other values, reshape will produce multiple copies of that instance set to those values.

Matrices or vectors of instances can also be created by concatenation:

```
#include trade.sdf
struct option p0,p1,p2;
p0 = optionCreate;
p1 = optionCreate;
p2 = p1 | p0;
```


### 16.1.5 Structure Indexing

Structure indexing may be used to reference a particular element in a structure array. The syntax follows that of matrix indexing. For example, given the following structure definition:

```
struct example1 {
    matrix x;
    matrix y;
    string str;
};
```

you could create an array of example1 structures and index it as follows:

```
struct example1 e1a;
struct example1 e1b;
e1a = e1a | e1b;
e1a[2,1].y = rndn(25,10);
```

In this example, e1a and $\mathbf{e} 1 \mathbf{b}$ are concatenated to create a $2 \times 1$ array of example1 structures that is assigned back to e1a. Then the $\mathbf{y}$ member of the [2,1] element of $\mathbf{e} 1 \mathrm{a}$ is set to a random matrix.

Indexing of structure arrays can occur on multiple levels. For example, let's define the following structures:

```
struct example3 {
```

```
        matrix w;
        string array sa;
};
struct example2 {
    matrix z;
    struct example3 e3;
};
```

and let's redefine example1 to include an instance of an example2 structure:

```
struct example1 {
    matrix x;
    matrix y;
    string str;
    struct example2 e2;
};
```

Let's assume that we have an example1 structure e1 like the one displayed in Figure 16.1. We could then index the structure as follows:
r = e1.e2[3,1].e3[2,1].w

You can also use indexing to reference the structure itself, rather than a member of that structure:

```
struct example3 e3tmp;
e3tmp = e1.e2[3,1].e3[2,1];
```

Or you can use indexing to reference a subarray of structures:

$$
\text { e3tmp }=\text { e1.e2[3,1].e3[., 1]; }
$$



Figure 16.1: Structure tree for e1

In this case, e 3 tmp would be an array of $3 \times 1$ example 3 structures, since the [ 3,1 ] member of e1.e2 contains a $3 \times 1$ array of example 3 structures.

It is important to remember, however, that when indexing a structure array on multiple levels, only the final index may resolve to an array of structures. For example:

$$
\text { e3tmp }=\text { e1.e2[., 1].e3[2, 1]; }
$$

would be invalid, since e1.e2[.,1] resolves to a $3 \times 1$ array of example 2 structures.

### 16.1.6 Saving an Instance to the Disk

Instances and vectors or matrices of instances of structures can be saved in a file on the disk, and later loaded from the file onto the disk. The syntax for saving an instance to the disk is

```
ret = savestruct(instance,filename);
```

The file on the disk will have an .fsr extension.

For example:

```
#include ds.sdf
struct DS p0;
p0 = reshape(dsCreate,2,1);
retc = saveStruct(p2,"p2");
```

This saves the vector of instances in a file called p2.fsr. retc will be zero if the save was successful; otherwise, nonzero.

### 16.1.7 Loading an Instance from the Disk

The syntax for loading a file containing an instance or matrix of instances is

```
instance, retc = loadstruct(file_name,structure_name);
```

For example:

```
#include trade.sdf;
struct DS p3;
{ p3, retc } = loadstruct("p2","ds");
```


### 16.1.8 Passing Structures to Procedures

Structures or members of structures can be passed to procedures. When a structure is passed as an argument to a procedure, it is passed by value. The structure becomes a local copy of the structure that was passed. The data in the structure is not duplicated unless the local copy of the structure has a new value assigned to one of its members. Structure arguments must be declared in the procedure definition:

```
struct rectangle {
    matrix ulx;
    matrix uly;
    matrix lrx;
    matrix lry;
    };
proc area(struct rectangle rect);
    retp((rect.lrx - rect.ulx).*(rect.uly - rect.lry));
endp;
```

Local structures are defined using a struct statement inside the procedure definition:

```
proc center(struct rectangle rect);
    struct rectangle cent;
    cent.lrx = (rect.lrx - rect.ulx) / 2;
    cent.ulx = -cent.lrx;
    cent.uly = (rect.uly - rect.lry) / 2;
    cent.lry = -cent.uly;
    retp(cent);
endp;
```


### 16.2 Structure Pointers

A structure pointer is a separate data type that contains the address of a structure and is used to reference that structure.

### 16.2.1 Creating and Assigning Structure Pointers

Given the following structure type definition:

```
struct example_struct {
    matrix x;
    matrix y;
};
```

a pointer to an example_struct structure can be created with the following syntax:

```
struct example_struct *esp;
```

However, at this point, esp is not yet pointing at anything. It has only been defined to be the kind of pointer that points at example_struct structures. To set it to point at a particular structure instance, we must first create the structure instance:

```
struct example_struct es;
```

and then we can set esp to point at es betting esp to the address of es:
esp = \&es;

The following code:

```
struct example_struct es2;
es2 = *esp;
```

copies the contents of the structure that esp is pointing at (i.e., the contents of es) to es2. It is the same as

```
struct example_struct es2;
es2 = es;
```


### 16.2.2 Structure Pointer References

To reference a member of a structure, we use a "dot" syntax. For example, we might use the following code to set the $\mathbf{x}$ member of es.

```
es.x = rndn(3,3);
```

To reference a member of a structure using a pointer to that structure, we use an "arrow" syntax. For example, we might use the following code to set the $\mathbf{x}$ member of es using the pointer esp:

$$
\text { esp }->x=\operatorname{rndn}(10,5) ;
$$

This code will modify es, since esp is merely a pointer to es.
Structure pointers cannot be members of a structure. The following is illegal:

```
struct example_struct_2 {
    matrix z;
    struct example_struct *ep;
};
```

Therefore, since a structure pointer will never be a member of a structure, neither
sp1->sp2->x;
nor
s.sp1->x;
will ever be valid ( $\mathbf{s p 1}$ and $\mathbf{s p 2}$ are assumed to be structure pointers, $\mathbf{s}$ a structure instance, and $\mathbf{x}$ a matrix). The "arrow" ( $->$ ) will only be valid if it is used for the first (or furthest left) dereference, as in:

```
sp1->st.x;
```

At this point we do not support indexing of structure pointers. Thus, a structure pointer should point at a scalar structure instance, not a matrix of structures. However, you may index members of that scalar structure instance. So, for example, let us suppose that you defined the following structure types:

```
struct sb {
    matrix y;
    matrix z;
```

```
};
struct sa {
    matrix x;
    struct structb s;
};
```

and then created an instance of an sa structure, a0, setting a0.s to a $3 \times 2$ matrix of sb structures. The following would be legal:

```
struct sa *sap
sap = &a0;
sap->s[3,1].y = rndn(3,3);
```


### 16.2.3 Using Structure Pointers in Procedures

Structure pointers are especially useful in cases where structures are passed into and out of procedures. If a procedure takes a structure as an argument and modifies any members of that structure, then it makes a local copy of the entire structure before modifying it. Thus if you want to have the modified copy of the structure after running the procedure, you need to pass the structure out of the procedure as one of its return arguments. For example:

```
struct example_struct {
    matrix x;
    matrix y;
    matrix z;
};
proc product(struct example_struct es);
    es.z = (es.x).*(es.y);
    retp(es);
endp;
struct example_struct es1;
```

```
es1.x = rndn(1000,100);
es1.y = rndn(1000,1);
es1 = product(es1);
```

In this example, the structure es1 is passed into the procedure, copied and modified. The modified structure is then passed out of the procedure and assigned back to es1.

Structure pointers allow you to avoid such excessive data copying and eliminate the need to pass a structure back out of a procedure in cases like this. When you pass a structure pointer into a procedure and then modify a member of the structure that it references, the actual structure is modified rather than a local copy of it. Thus there is no need to pass the modifed structure back out of the procedure. For example, the above example could be accomplished using structure pointers as follows:

```
struct example_struct {
    matrix x;
    matrix y;
    matrix z;
};
proc(0) = product(struct example_struct *esp);
    esp->z = (esp->x).*(esp->y);
endp;
struct example_struct es1;
struct example_struct *es1p;
es1p = &es1;
es1.x = rndn(1000,100);
es1.y = rndn(1000,1);
product(es1p);
```

In this case, the procedure modifies the structure es1, which esip is pointing at, instead of a local copy of the structure.

### 16.3 Special Structures

There are three common types of structures that will be found in the GAUSS Run-Time Library and applications.

The DS and PV structures are defined in the GAUSS Run-Time Library. Their definitions are found in ds.sdf and pv.sdf, respectively, in the src source code subdirectory.

Before structures, many procedures in the Run-Time Library and all applications had global variables serving a variety of purposes, such as setting and altering defaults. Currently, these variables are being entered as members of "control" structures.

### 16.3.1 The DS Structure

The DS structure, or "data" structure, is a very simple structure. It contains a member for each GAUSS data type. The following is found in ds.sdf:

```
struct DS {
    scalar type;
    matrix dataMatrix;
    array dataArray;
    string dname;
    string array vnames;
    };
```

This structure was designed for use by the various optimization functions in GAUSS, in particular, sqpSolvemt, as well as a set of gradient procedures, gradmt, hessmt, et al.

These procedures all require that the user provide a procedure computing a function (to be optimized or take the derivative of, etc.), which takes the DS structure as an argument. The Run-Time Library procedures such as sqpSolvemt take the DS structure as an argument and pass it on to the user-provided procedure without modification. Thus, the user can put into that structure whatever might be needed as data in the procedure.

To initialize an instance of a DS structure, the procedure dsCreate is defined in ds. src:

```
#include ds.sdf
struct DS d0;
dQ = dsCreate;
```


### 16.3.2 The PV Structure

The PV structure, or parameter vector structure, is used by various optimization, modelling, and gradient procedures, in particular sqpSolvemt, for handling the parameter vector. The GAUSS Run-Time Library contains special functions that work with this structure. They are prefixed by "pv" and defined in pv.src. These functions store matrices and arrays with parameters in the structure, and retrieve various kinds of information about the parameters and parameter vector from it.

## "Packing" into a PV Structure

The various procedures in the Run-Time Library and applications for optimization, modelling, derivatives, etc., all require a parameter vector. Parameters in complex models, however, often come in matrices of various types, and it has been the responsibility of the programmer to generate the parameter vector from the matrices and vice versa. The PV procedures make this problem much more convenient to solve.

The typical situation involves two parts: first, "packing" the parameters into the PV structure, which is then passed to the Run-Time Library procedure or application; and second, "unpacking" the PV structure in the user-provided procedure for use in computing the objective function. For example, to pack parameters into a PV structure:

```
#include sqpsolvemt.sdf
/* starting values */
b0 = 1; /* constant in mean equation */
garch = { .1, .1 }; /* garch parameters */
arch = { .1, .1 }; /* arch parameters */
omega = .1; /* constant in variance equation */
```

```
struct PV p0;
p0 = pvPack(pvCreate,b0,"b\otimes");
p0 = pvPack(p0,garch,"garch");
p0 = pvPack(p0,arch,"arch");
p0 = pvPack(p0,omega,"omega");
/* data */
z = loadd("tseries");
struct DS d0;
dQ.dataMatrix = z;
```

Next, in the user-provided procedure for computing the objective function, in this case minus the log-likelihood, the parameter vector is unpacked:

```
proc ll(struct PV pQ, struct DS dQ);
    local b0,garch,arch,omega,p,q,h,u,vc,w;
    b0 = pvUnpack(p0,"b0");
    garch = pvUnpack(p0,"garch");
    arch = pvUnpack(p0,"arch");
    omega = pvUnpack(p0,"omega");
    p = rows(garch);
    q = rows(arch);
    u = d0.dataMatrix - b0;
    vc = moment(u,0)/rows(u);
    w = omega + (zeros(q,q) | shiftr((u.*ones(1,q))',
        seqa(q-1,-1,q))) * arch;
    h = recserar(w,vc*ones(p,1),garch);
    logl = -0.5 * ((u.*u)./h + ln(2*pi) + ln(h));
    retp(logl);
```

endp;

## Masked Matrices

The pvUnpack function unpacks parameters into matrices or arrays for use in computations. The first argument is a PV structure containing the parameter vector. Sometimes the matrix or vector is partly parameters to be estimated (that is, a parameter to be entered in the parameter vector) and partly fixed parameters. To distinguish between estimated and fixed parameters, an additional argument is used in the packing function called a "mask", which is strictly conformable to the input matrix. Its elements are set to 1 for an estimated parameter and 0 for a fixed parameter. For example:

```
p0 = pvPackm(p0,.1*eye(3),"theta",eye(3));
```

Here just the diagonal of a $3 \times 3$ matrix is added to the parameter vector.

When this matrix is unpacked, the entire matrix is returned with current values of the parameters on the diagonal:

```
print pvUnpack(p0,"theta");
```

| 0.1000 | 0.0000 | 0.0000 |
| :--- | :--- | :--- |
| 0.0000 | 0.1000 | 0.0000 |
| 0.0000 | 0.0000 | 0.1000 |

## Symmetric Matrices

Symmetric matrices are a special case because even if the entire matrix is to be estimated, only the nonredundant portion is to be put into the parameter vector. Thus, for them there are special procedures. For example:

```
vc = { 1 . 6 .4, . 6 1 .2, .4 .2 1 };
p0 = pvPacks(p0,vc,"vc");
```

There is also a procedure for masking in case only a subset of the nonredundant elements are to be included in the parameter vector:

```
vc = { 1 .6 .4, .6 1 .2, .4 . 2 1 };
mask = {11 0, 1 10, 0 0 1 };
p0 = pvPacksm(p0,vc,"vc",mask);
```


## Fast Unpacking

When unpacking matrices using a matrix name, pvUnpack has to make a search through a list of names, which is relatively time-consuming. This can be alleviated by using an index rather than a name in unpacking. To do this, though, requires using a special pack procedure that establishes the index:

```
p0 = pvPacki(p0,b0,"b0",1);
p0 = pvPacki(p0,garch,"garch",2);
p0 = pvPacki(p0,arch,"arch",3);
p0 = pvPacki(p0,omega,"omega",4);
Now they may be unpacked using the index number:
b0 = pvUnpack(p0,1);
garch = pvUnpack(p0,2);
arch = pvUnpack(p0,3);
omega = pvUnpack(p0,4);
```

When packed with an index number, they may be unpacked either by index or by name, but unpacking by index is faster.

### 16.3.3 Miscellaneous PV Procedures

## pvList

This procedure generates a list of the matrices or arrays packed into the structure:

```
p0 = pvPack(p0,b0,"b0");
p0 = pvPack(p0,garch,"garch");
p0 = pvPack(p0,arch,"arch");
p0 = pvPack(p0,omega,"omega");
print pvList(p0);
    b0
    garch
    arch
    omega
```


## pvLength

This procedure returns the length of the parameter vector:

```
print pvLength(p0);
```

6.0000

## pvGetParNames

This procedure generates a list of parameter names:
print pvGetParNames(p0);

```
b0[1,1]
garch[1,1]
garch[2,1]
arch[1,1]
arch[2,1]
omega[1,1]
```


## pvGetParVector

This procedure returns the parameter vector itself:
print pvGetParVector (p0);
1.0000
0.1000
0.1000
0.1000
0.1000
1.0000

## pvPutParVector

This procedure replaces the parameter vector with the one in the argument:

```
newp = { 1.5, .2, .2, .3, .3, . 8 };
p0 = pvPutParVector(newp);
print pvGetParVector(p0);
```

1.5000
0.2000
0.2000
0.3000
0.3000
0.8000

## pvGetIndex

This procedure returns the indices in the parameter vector of the parameters in a matrix. These indices are useful when setting linear constraints or bounds in sqpSolvemt. Bounds, for example, are set by specifying a $\mathrm{K} \times 2$ matrix where K is the length of the parameter vector and the first column are the lower bounds and the second the upper bounds. To set the bounds for a particular parameter, then, requires knowing where that parameter is in the parameter vector. This information can be found using pvGetIndex. For example:

```
// get indices of lambda parameters in parameter vector
lind = pvGetIndex(par0,"lambda");
// set bounds constraint matrix to unconstrained default
c0.bounds = ones(pvLength(par0),1).*(-1e250^1e250);
// set bounds for lambda parameters to be positive
c0.bounds[lind,1] = zeros(rows(lind),1);
```


### 16.3.4 Control Structures

Another important class of structures is the "control" structure. Applications developed before structures were introduced into GAUSS typically handled some program specifications by the use of global variables which had some disadvantages, in particular, preventing the nesting of calls to procedures.

Currently, the purposes served by global variables are now served by the use of a control structure. For example, for sqpSolvemt:

```
struct sqpSolvemtControl {
    matrix A;
    matrix B;
    matrix C;
    matrix D;
    scalar eqProc;
    scalar ineqProc;
```

```
matrix bounds;
scalar gradProc;
scalar hessProc;
scalar maxIters;
scalar dirTol;
scalar CovType;
scalar feasibleTest;
scalar maxTries;
scalar randRadius;
scalar trustRadius;
scalar seed;
scalar output;
scalar printIters;
matrix weights;
};
```

The members of this structure determine optional behaviors of sqpSolvemt.

## 16.4 sqpSolvemt

sqpSolvemt is a procedure in the GAUSS Run-Time Library that solves the general nonlinear programming problem using a Sequential Quadratic Programming descent method, that is, it solves

```
minf(0)
```

subject to

| $A \theta=B$ | linear equality |
| :--- | :--- |
| $C \theta>=D$ | linear inequality |
| $H(\theta)=0$ | nonlinear equality |
| $G(\theta)>=0$ | nonlinear inequality |
| $\theta_{l b}<=\theta<=\theta_{u b}$ | bounds |

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The linear and bounds constraints are redundant with respect to the nonlinear constraints, but are treated separately for computational convenience.

The call to sqpSolvemt has four input arguments and one output argument:

$$
\text { out }=\text { SQPsolveMT }(\mathcal{E} f c t, P, D, C) ;
$$

### 16.4.1 Input Arguments

The first input argument is a pointer to the objective function to be minimized. The procedure computing this objective function has two arguments: a PV structure containing the start values, and a DS structure containing data, if any. For example:

```
proc fct(struct PV p0, struct DS dQ);
    local y, x, bQ, b, e, s;
    y = dQ[1].dataMatrix;
    x = dQ[2].dataMatrix;
    b0 = pvUnpack(p0,"constant");
    b = pvUnpack(p0,"coefficients");
    e = y - b0 - x * b;
    s = sqrt(e'e/rows(e));
    retp(-pdfn(e/s);
endp;
```

Note that this procedure returns a vector rather than a scalar. When the objective function is a properly defined log-likelihood, returning a vector of minus log-probabilities permits the calculation of a QML covariance matrix of the parameters.

The remaining input arguments are structures:
$P \quad$ a PV structure containing starting values of the parameters
$D \quad$ a DS structure containing data, if any

C an sqpSolvemtControl structure

The DS structure is optional. sqpSolvemt passes this argument on to the user-provided procedure that $\mathcal{E f f c t}$ is pointing to without modification. If there is no data, a default structure can be passed to it.

## sqpSolvemtControl Structure

A default sqpSolvemtControl structure can be passed in the fourth argument for an unconstrained problem. The members of this structure are as follows:

A $\quad \mathrm{M} \times \mathrm{K}$ matrix, linear equality constraint coecients: $A \theta=B$, where p is a vector of the parameters.

B $\quad \mathrm{M} \times 1$ vector, linear equality constraint constants: $A \theta=B$, where p is a vector of the parameters.

C $\quad \mathrm{M} \times \mathrm{K}$ matrix, linear inequality constraint coefficients: $C \theta=D$, where p is a vector of the parameters.

D $\quad \mathrm{M} \times 1$ vector, linear inequality constraint constants: $C \theta=D$, where p is a vector of the parameters.
eqProc scalar, pointer to a procedure that computes the nonlinear equality constraints. When such a procedure has been provided, it has two input arguments, instances of PV and DS structures, and one output argument, a vector of computed inequality constraints.

Default $=$.; i.e., no inequality procedure.
IneqProc scalar, pointer to a procedure that computes the nonlinear inequality constraints. When such a procedure has been provided, it has two input arguments, instances of PV and DS structures, and one output argument, a vector of computed inequality constraints.
Default = .; i.e., no inequality procedure .

| Bounds | $1 \times 2$ or $\mathrm{K} \times 2$ matrix, bounds on parameters. If $1 \times 2$ all parameters have same bounds. |
| :---: | :---: |
|  | Default $=-1 \mathrm{e} 2561 \mathrm{e} 256$. |
| GradProc | scalar, pointer to a procedure that computes the gradient of the function with respect to the parameters. When such a procedure has been provided, it has two input arguments, instances of PV and DS structures, and one output argument, the derivatives. If the function procedure returns a scalar, the gradient procedure returns a $1 \times \mathrm{K}$ row vector of derivatives. If function procedure turns an $N \times 1$ vector, the gradient procedure returns an $N \times K$ matrix of derivatives. |
|  | This procedure may compute a subset of the derivatives. sqpSolvemt will compute numerical derivatives for all those elements set to missing values in the return vector or matrix. |
|  | Default = .; i.e., no gradient procedure has been provided. |
| HessProc | scalar, pointer to a procedure that computes the Hessian; i.e., the matrix of second order partial derivatives of the function with respect to the parameters. When such a procedure has been provided, it has two input arguments, instances of PV and DS structures, and one output argument, a vector of computed inequality constraints. |
|  | Default = .; i.e., no Hessian procedure has been provided. |
|  | Whether the objective function procedure returns a scalar or vector, the Hessian procedure must return a $\mathrm{K} \times \mathrm{K}$ matrix. Elements set to missing values will be computed numerically by sqpSolvemt. |
| MaxIters | scalar, maximum number of iterations. Default $=1 \mathrm{e}+5$. |
| MaxTries | scalar, maximum number of attemps in random search. Default $=100$. |
| DirTol | scalar, convergence tolerance for gradient of estimated coefficients. Default $=1 \mathrm{e}-5$. When this criterion has been satisifed, sqpSolvemt exits the iterations. |
| CovType | scalar, if 2, QML covariance matrix, else if 0 , no covariance matrix is computed, else ML covariance matrix is computed. For a QML covariance matrix, the objective function procedure must return an $N \times 1$ vector of minus log-probabilities. |


| FeasibleTest | scalar, if nonzero, parameters are tested for feasibility before computing <br> function in line search. If function is defined outside inequality boundaries, <br> then this test can be turned off. Default $=1$. |
| :--- | :--- |
| randRadius | scalar, if zero, no random search is attempted. If nonzero, it is the radius of <br> the random search. Default $=.001$. |
| seed | scalar, if nonzero, seeds random number generator for random search, <br> otherwise time in seconds from midnight is used. |
| trustRadius | scalar, radius of the trust region. If scalar missing, trust region not applied. <br> The trust sets a maximum amount of the direction at each iteration. Default <br> $=.001$. |
| output | scalar, if nonzero, results are printed. Default $=0$. |
| PrintIters | scalar, if nonzero, prints iteration information. Default $=0$. |
| weights | vector, weights for objective function returning a vector. Default $=1$. |

### 16.4.2 Output Argument

The single output argument is an sqpSolvemtOut structure. Its definition is:

```
struct SQPsolveMTOut {
    struct PV par;
    scalar fct;
    struct SQPsolveMTLagrange lagr;
    scalar retcode;
    matrix moment;
    matrix hessian;
    matrix xproduct;
    };
```

The members of this structure are:
par instance of a PV structure containing the parameter estimates are placed in the matrix member par.
fct
scalar, function evaluated at final parameter estimates.
lagr an instance of an SQPLagrange structure containing the Lagrangeans for the constraints. For an instance named lagr, the members are:

| lagr.lineq | $\mathrm{M} \times 1$ vector, Lagrangeans of linear equality <br> constraints |
| :---: | :--- |
| lagr.nlineq | $\mathrm{N} \times 1$ vector, Lagrangeans of nonlinear equality <br> constraints |
| lagr.linineq | $\mathrm{P} \times 1$ vector, Lagrangeans of linear inequality <br> constraints |
| lagr.nlinineq | $\mathrm{Q} \times 1$ vector, Lagrangeans of nonlinear inequality <br> constraints |

lagr.bounds $\mathrm{K} \times 2$ matrix, Lagrangeans of bounds
Whenever a constraint is active, its associated Lagrangean will be nonzero. For any constraint that is inactive throughout the iterations as well as at convergence, the corresponding Lagrangean matrix will be set to a scalar missing value.
retcode return code:

## 0 normal convergence

1 forced exit
2 maximum number of iterations exceeded
3 function calculation failed
4 gradient calculation failed
5 Hessian calculation failed
6 line search failed
7 error with constraints
8 function complex
9 feasible direction couldn't be found

### 16.4.3 Example

Define

$$
Y=\Lambda \eta+\theta
$$

where $\Lambda$ is a $\mathrm{K} \times \mathrm{L}$ matrix of loadings, $\eta$ an $\mathrm{L} \times 1$ vector of unobserved "latent" variables, and $\theta$ a $K \times 1$ vector of unobserved errors. Then

$$
\Sigma=\Lambda \Phi \Lambda \prime \Psi
$$

where $\Phi$ is the $\mathrm{L} \times \mathrm{L}$ covariance matrix of the latent variables, and $\Psi$ is the $\mathrm{K} \times \mathrm{K}$ covariance matrix of the errors.

The log-likelihood of the $i^{t h}$ observation is

$$
\log P(i)=-\frac{1}{2}[K \ln (2 \pi)+\ln |\pi|+Y(i) \Sigma Y(i) \prime]
$$

Not all elements of $\Lambda, \Phi$, and $\Psi$ can be estimated. At least one element of each column of $\Lambda$ must be fixed to 1 , and $\Psi$ is usually a diagonal matrix.

## Constraints

To ensure a well-defined log-likelihood, constraints on the parameters are required to guarantee positive definite covariance matrices. To do this, a procedure is written that returns the eigenvalues of $\Sigma$ and $\Phi$ minus a small number. sqpSolvemt then finds parameters such that these eigenvalues are greater than or equal to that small number.

### 16.4.4 The Command File

This command file can be found in the file sqpfact.e in the examples subdirectory:

```
#include sqpsolvemt.sdf
lambda = { 1.0 0.0,
    0.5 0.0,
    0.0 1.0,
    0.0 0.5 };
lmask = { 0 0,
    10,
    0 0,
    0 1 };
phi = { 1.0 0.3,
    0.3 1.0 };
psi = { 0.6 0.0 0.0 0.0,
    0.0 0.6 0.0 0.0,
    0.0 0.0 0.6 0.0,
    0.0 0.0 0.0 0.6 };
tmask={1000,
        0 100,
        0 0 1 0,
        0 0 0 1 };
struct PV par0;
par0 = pvCreate;
par@ = pvPackm(par@,lambda,"lambda",lmask);
par@ = pvPacks(par@,phi,"phi");
par@ = pvPacksm(par0,psi,"psi",tmask);
struct SQPsolveMTControl cQ;
```

```
c0 = sqpSolveMTcontrolCreate;
lind = pvGetIndex(par0,"lambda"); /* get indices of lambda */
                                    /* parameters in parameter */
                                    /* vector */
tind = pvGetIndex(par0,"psi"); /* get indices of psi */
                                    /* parameters in parameter */
                                    /* vector */
CQ.bounds = ones(pvLength(par0),1).*(-1e250~1e250);
cQ.bounds[lind,1] = zeros(rows(lind),1);
cQ.bounds[lind,2] = 10*ones(rows(lind),1);
c0.bounds[tind,1] = .001*ones(rows(tind),1);
cQ.bounds[tind,2] = 100*ones(rows(tind),1);
cQ.output = 1;
CQ.printIters = 1;
cQ.trustRadius = 1;
cQ.ineqProc = &ineq;
cQ. covType = 1;
struct DS dD;
dQ = dsCreate;
dQ.dataMatrix = loadd("maxfact");
output file = sqpfact.out reset;
struct SQPsolveMTOut out0;
out0 = SQPsolveMT(&lpr,par0,dQ,c0);
lambdahat = pvUnpack(out0.par,"lambda");
phihat = pvUnpack(out0.par,"phi");
psihat = pvUnpack(out0.par,'psi");
print "estimates";
print;
print "lambda" lambdahat;
```

```
print;
print "phi" phihat;
print;
print "psi" psihat;
struct PV stderr;
stderr = out0.par;
if not scalmiss(outQ.moment);
    stderr = pvPutParVector(stderr,sqrt(diag(outQ.moment)));
    lambdase = pvUnpack(stderr,"lambda");
    phise = pvUnpack(stderr,"phi");
    psise = pvUnpack(stderr,"psi");
    print "standard errors";
    print;
    print "lambda" lambdase;
    print;
    print "phi" phise;
    print;
    print "psi" psise;
endif;
output off;
proc lpr(struct PV par1, struct DS data1);
    local lambda,phi,psi,sigma,logl;
    lambda = pvUnpack(par1,"lambda");
    phi = pvUnpack(par1,"phi");
    psi = pvUnpack(par1,"psi");
    sigma = lambda*phi*lambda' + psi;
    logl = -lnpdfmvn(data1.dataMatrix,sigma);
    retp(logl);
endp;
proc ineq(struct PV par1, struct DS data1);
```

```
    local lambda,phi,psi,sigma,e;
    lambda = pvUnpack(par1,"lambda");
    phi = pvUnpack(par1,"phi");
    psi = pvUnpack(par1,"psi");
    sigma = lambda*phi*lambda' + psi;
    e = eigh(sigma) - .001; /* eigenvalues of sigma */
    e = e | eigh(phi) - .001; /* eigenvalues of phi */
    retp(e);
endp;
```



## Run-Time Library Structures

## 17

Two structures are used by several GAUSS Run-Time Library functions for handling parameter vectors and data: the PV parameter structure and the DS data structure.

### 17.1 The PV Parameter Structure

The members of an instance of structure of type PV are all "private," that is, not accessible directly to the user. It is designed to handle parameter vectors for threadsafe optimization functions. Entering and receiving parameter vectors, and accessing properties of this vector, are accomplished using special functions.

Suppose you are optimizing a function containing a $K \times L$ matrix of coefficients. The optimization function requires a parameter vector but your function uses a $\mathrm{K} \times \mathrm{L}$ matrix. Your needs and the needs of the optimization function can be both satisfied by an instance of the structure of type PV. For example:

```
struct PV p1;
p1 = pvCreate;
```

```
x = zeros(4,3); /* on input contains start values, */
    /* on exit contains estimates */
p1 = pvPack(p1,x,"coefficients");
```

The pvCreate function initializes p1 to default values. pvPack enters the $4 \times 3$ matrix stored row-wise as a $12 \times 1$ parameter vector for the optimization function. The optimization program will pass the instance of the structure of type PV to your objective function.

By calling pvUnpack your $4 \times 3$ coefficient matrix is retrieved from the parameter vector. For example, in your procedure you have

```
x = pvUnpack(p1,"coefficients");
```

and now $\mathbf{x}$ is a $4 \times 3$ matrix of coefficients for your use in calculating the object function.
Suppose that your objective function has parameters to be estimated in a covariance matrix. The covariance matrix is a symmetric matrix where only the lower left portion contains unique values for estimation. To handle this, use pvPacks. For example:

```
struct PV p1;
p1 = pvCreate;
cov={ 1 . 1 . 1,
    .1 1 .1,
    .1.1 1};
p1 = pvPacks(p1,cov,"covariance");
```

Only the lower left portion of cov will be stored in the parameter vector. When the covariance matrix is unpacked, the parameters in the parameter vector will be entered into both the lower and upper portions of the matrix.

There may be cases where only a portion of a matrix being used to compute the objective function are parameters to be estimated. In this case use pvPackm with a "mask" matrix that contains ones where parameters are to be estimated and zeros otherwise. For example,

```
struct PV p1;
p1 = pvCreate;
cov = { 1 .5,
    . 5 1 };
mask = { 0 1,
    10};
p1 = pvPacksm(p1,cov,"correlation",mask);
```

Here only the one element in the lower left of cov is stored in the parameter vector. Suppose the optimization program sends a trial value for that parameter of, say, .45 . When the matrix is unpacked in your procedure it will contain the fixed values associated with the zeros in the mask as well as the trial value in that part of the matrix associated with the ones. Thus,

```
print unpack(p1,"correlation");
    1.0000 .4500
    .4500 1.0000
```

A mask may also be used with general matrices to store a portion of a matrix in the parameter vector.

```
struct PV p1;
p1 = pvCreate;
m = { 0 . 5 1,
    . 5 0 . 3 };
mask = { 0 1 1,
    1 0 0};
p1 = pvPackm(p1,m,"coefficients",mask);
```

A PV instance can, of course, hold parameters from all these types of matrices: symmetric, masked symmetric, rectangular, and masked rectangular. For example:

```
lambda = { 1.0 0.0,
    0.5 0.0,
    0.0 1.0,
    0.0 0.5};
lmask = { 0 0,
        1 0,
        0 0,
        0 1};
phi = { 1.0 0.3,
        0.3 1.0 };
theta ={ {0.6 0.0 0.0 0.0,
    0.0 0.6 0.0 0.0,
    0.0 0.0 0.6 0.0,
    0.0 0.0 0.0 0.6};
tmask = {\begin{array}{lllll}{1}&{0}&{0}&{0,}\end{array},
        0 1 0 0,
        0 0 1 0,
        0 0 0 1};
struct PV par0;
par0 = pvCreate;
par0 = pvPackm(par@,lambda,"lambda",lmask);
par@ = pvPacks(par0,phi,"phi");
par0 = pvPacksm(par0,theta,"theta",tmask);
```

It isn't necessary to know where in the parameter vector the parameters are located in order to use them in your procedure calculating the objective function. Thus:

```
lambda = pvUnpack(par1,"lambda");
```

```
phi = pvUnpack(par1,"phi");
theta = pvUnpack(par1,"theta");
sigma = lambda*phi*lambda' + theta;
```

Additional functions are available to retrieve information on the properties of the parameter vector. pvGetParVector and pvPutParVector get and put parameter vector from and into the PV instance, pvGetParNames retrieves names for the elements of the parameter vector, pvList returns the list of matrix names in the PV instance, pvLength the length of the parameter vector.

```
struct PV p1;
p1 = pvCreate;
cov = { 1 . 5,
    . 5 1 };
mask = { 0 1,
    10 };
```

p1 = pvPacksm(p1,cov,"correlation",mask);
print pvGetParVector(p1);
.5000
p1 = pvPutParVector (p1,.8);
print pvGetParVector(p1);
.8000
print pvUnpack(p1,"correlation");
1.0000 .8000
.80001 .0000
print pvGetParNames(p1);
correlation[2,1]

```
print pvLength(p1);
```

1.0000

Also, pvTest tests an instance to make sure it is properly constructed. pvCreate generates an initialized instance, and pvGetIndex returns the indices of the parameters of an input matrix in the parameter vector. This last function is most useful when constructing linear constraint indices for the optimization programs.

### 17.2 Fast Pack Functions

Unpacking matrices using matrix names is slow because it requires a string search through a string array of names. A set of special packing functions are provided that avoid the search altogether. These functions use a "table" of indices that you specify to find the matrix in the PV instance. For example:

```
struct PV p1;
p1 = pvCreate(2);
y = rndn(4,1);
x = rndn(4,4);
p1 = pvPacki(p1,y,"Y",1);
p1 = pvPacki(p1,x,"X",2);
print pvUnpack(p1,1);
    . }342
    .0407
    . }561
    .0953
print pvUnpack(p1,"Y");
```

The call to pvPacki puts an entry in the table associating the matrix in its second argument with the index 1 . As indicated above the matrix can be unpacked either by index or by name. Unpacking by index, however, is much faster than by name.

Note that the matrix can be unpacked using either the index or the matrix name.

There are index versions of all four of the packing functions, pvPacki, pvPackmi, pvPacksi, and pvPacksmi.

### 17.3 The DS Data Structure

An instance of the DS data structure contains the following members:

```
struct DS d0;
```

| dQ.dataMatrix | $\mathrm{M} \times \mathrm{K}$ matrix, data |
| :--- | :--- |
| dQ.dataArray | N -dimensional array, data |
| dQ.type | scalar |
| dQ.dname | string |
| dQ.vnames | string array |

The definition and use of the elements of $\mathbf{d O}$ are determined by the particular application and are mostly up to the user. A typical use might use a vector of structures. For example, suppose the objective function requires a vector of observations on a dependent variable as well as on $K$ independent variables. Then:

```
struct DS d0;
d0 = dsCreate;
y = rndn(20,1);
x = rndn(20,5);
d0 = reshape(d0,2,1);
dQ[1].dataMatrix = y;
d0[2].dataMatrix = X;
```

The dO instance would be passed to the optimization program which then passes it to your procedure computing the objective function. For example:

```
proc lpr(struct PV p1, struct DS d1);
    local u;
    u = dQ[1].dataMatrix - dQ[2].dataMatrix * pvUnpack(p1,"beta");
    retp(u'u);
endp;
```

A particular application may require setting other members of the DS instance for particular purposes, but in general you may use them for your own purposes. For example, dQ . dname could be set to a GAUSS dataset name from which you read the data in the objective function procedure, or dQ. vnames could be set to the variable names of the columns of the data stored in dQ. dataMatrix, or dQ.type could be an indicator variable for the elements of a vector of DS instances.

The following are complete examples of the use of the PV and DS structures. The first example fits a set of data to the Micherlitz model. It illustrates packing and unpacking by index.

```
#include sqpsolvemt.sdf
struct DS Y;
Y = dsCreate;
Y.dataMatrix = 3.183|
```

```
    3.059|
    2.871|
    2.622|
    2.541|
    2.184|
    2.110|
    2.075|
    2.018|
    1.903|
    1.770|
    1.762|
    1.550;
struct DS X;
X = dsCreate;
X.dataMatrix = seqa(1,1,13);
struct DS Z;
Z = reshape(Z,2,1);
Z[1] = Y;
Z[2] = X;
struct SQPsolveMTControl c1;
c1 = sqpSolveMTcontrolCreate; /* initializes */
                                    /* default values */
c1.bounds = 0~100; /* constrains parameters */
c1.CovType = 1;
c1.output = 1;
c1.printIters = 0;
c1.gradProc = &grad;
struct PV par1;
par1 = pvCreate(1);
```

```
start = { 2, 4, 2 };
par1 = pvPacki(par1,start,"Parameters",1);
struct SQPsolveMTout out1;
out1 = SQPsolveMT(&Micherlitz,par1,Z,c1);
estimates = pvGetParVector(out1.par);
print " parameter estimates ";
print estimates;
print;
print " standard errors ";
print sqrt(diag(out1.moment));
proc Micherlitz(struct PV par1,struct DS Z);
    local p0,e,s2;
    p@ = pvUnpack(par1,1);
    e = Z[1].dataMatrix - p0[1] - p0[2]*exp(-p0[3]
        *Z[2].dataMatrix);
    s2 = moment(e,0)/(rows(e)-1);
    retp( (2/rows(e))*(e.*e/s2 + ln(2*pi*s2)));
endp;
proc grad(struct PV par1, struct DS Z);
    local p0,e,e1,e2,e3,w,g,s2;
    p0 = pvUnpack(par1,1);
    w = exp(-p0[3]*Z[2].dataMatrix);
    e = z[1].dataMatrix - p0[1] - p0[2] * w;
    s2 = moment(e,0) / rows(e);
    e1 = - ones(rows(e),1);
    e2 = -w;
    e3 = p0[2]*Z[2].dataMatrix.*w;
    w = (1 - e.*e / s2) / rows(e);
    g = e.*e1 + w*(e'e1);
    g = g ~ (e.*e2 + w* (e'e2));
    g = g ~ (e.*e3 + w*(e'e3));
```

```
    retp(4*g/(rows(e)*s2));
endp;
```

This example estimates parameters of a "confirmatory factor analysis" model.

```
\#include sqpsolvemt.sdf
lambda = { 1.0 0.0,
    0.5 0.0,
    0.0 1.0,
    0.0 0.5 };
lmask = { 0 0,
    1 0,
    0 0,
    0 1 };
phi = { 1.0 0.3,
    0.3 1.0 };
theta = { 0.6 0.0 0.0 0.0,
    0.0 0.6 0.0 0.0,
    0.0 0.0 0.6 0.0,
    0.0 0.0 0.0 0.6 };
tmask = {\begin{array}{lllll}{1}&{0}&{0}&{0,}\end{array}]
    0 1 0 0,
        0 0 1 0,
        0 0 0 1};
```

struct PV par0;
par0 = pvCreate;
par@ = pvPackm(par@,lambda,"lambda",lmask);
par0 = pvPacks(parQ,phi,"phi");
par@ = pvPacksm(par@,theta,"theta",tmask);

```
struct SQPsolveMTControl cQ;
cQ = sqpSolveMTcontrolCreate;
lind = pvGetIndex(par0,"lambda"); /* get indices of */
    /* lambda parameters */
    /* in parameter vector */
tind = pvGetIndex(par0,"theta"); /* get indices of */
                                    /* theta parameters */
                                    /* in parameter vector */
cQ.bounds = ones(pvLength(par0),1).*(-1e250~1e250);
cQ.bounds[lind,1] = zeros(rows(lind),1);
cQ.bounds[lind,2] = 10*ones(rows(lind),1);
c0.bounds[tind,1] = .001*ones(rows(tind),1);
cQ.bounds[tind,2] = 100*ones(rows(tind),1);
cQ.ineqProc = &ineq;
cQ.covType = 1;
struct DS dD;
dQ = dsCreate;
dQ.dataMatrix = loadd("maxfact");
struct SQPsolveMTOut out0;
out0 = SQPsolveMT(&lpr,par0,d0,c0);
lambdahat = pvUnpack(out0.par,"lambda");
phihat = pvUnpack(out0.par,"phi");
thetahat = pvUnpack(out0.par,"theta");
print "estimates";
print;
print "lambda" lambdahat;
print;
```

```
print "phi" phihat;
print;
print "theta" thetahat;
struct PV stderr;
stderr = out0.par;
if not scalmiss(outQ.moment);
    stderr =
        pvPutParVector(stderr,sqrt(diag(out0.moment)));
    lambdase = pvUnpack(stderr,"lambda");
    phise = pvUnpack(stderr,"phi");
    thetase = pvUnpack(stderr,"theta");
    print "standard errors";
    print;
    print "lambda" lambdase;
    print;
    print "phi" phise;
    print;
    print "theta" thetase;
endif;
proc lpr(struct PV par1, struct DS data1);
    local lambda,phi,theta,sigma,logl;
    lambda = pvUnpack(par1,"lambda");
    phi = pvUnpack(par1,"phi");
    theta = pvUnpack(par1,"theta");
    sigma = lambda*phi*lambda' + theta;
    logl = -lnpdfmvn(data1.dataMatrix,sigma);
    retp(logl);
endp;
```

```
proc ineq(struct PV par1, struct DS data1);
    local lambda, phi, theta, sigma,e;
    lambda = pvUnpack(par1,"lambda");
    phi = pvUnpack(par1,"phi");
    theta = pvUnpack(par1,"theta");
    sigma \(=\) lambda*phi*lambda' + theta;
    e = eigh(sigma) - .001; /* eigenvalues of sigma */
    e = e | eigh(phi) - . 001 ; /* eigenvalues of phi */
    retp(e);
endp;
```


## Multi-Threaded Programming in GAUSS

## 18

The term thread comes from the phrase "thread of execution"-simply, it denotes a section of code that you want to execute. A single-threaded program has only one thread of execution, i.e., the program itself. A multi-threaded program is one that can have multiple threads-sections of code-executing simultaneously. Since these threads are part of the same program, they share the same workspace, and see and operate on the same symbols. Threads allow you to take full advantage of the hardware processing resources available on hyper-threaded, multi-core, and multi-processor systems, executing independent calculations simultaneously, combining and using the results of their work when done.

### 18.1 The Functions

GAUSS includes four keywords for multi-threading your programs:

ThreadStat Marks a single statement to be executed as a thread.
ThreadBegin Marks the beginning of a block of code to be executed as a thread.

ThreadEnd Marks the end of a block of code to be executed as a thread.
ThreadJoin Completes the definition of a set of threads, waits until they are done.

ThreadStat defines a single statement to be executed as a thread:

```
ThreadStat n = m'm;
```

ThreadBegin and ThreadEnd define a multi-line block of code to be executed as a thread:

$$
\begin{aligned}
& \text { ThreadBegin; } \\
& \qquad \begin{array}{l}
y=x \\
z=y
\end{array} \\
& \text { ThreadEnd; }
\end{aligned}
$$

Together these define sets of threads to be executed concurrently:

```
ThreadStat n = m'm; // Thread 1
ThreadBegin; // Thread 2
    y = x'x;
    z = y'y;
ThreadEnd;
ThreadBegin; // Thread 3
    q = r'r;
    r = q'q;
ThreadEnd;
ThreadStat p = o'o; // Thread 4
```

Finally, ThreadJoin completes the definition of a set of threads. It waits for the threads in a set to finish and rejoin the creating (the parent) thread, which can then continue, making use of their individual calculations:

| ThreadBegin; $\begin{aligned} & y=x \prime x ; \\ & z=y \prime y ; \end{aligned}$ | // Thread 1 |
| :---: | :---: |
| ThreadEnd; |  |
| $\begin{aligned} & \text { ThreadBegin; } \\ & \begin{array}{c} q=r \prime r \\ r=q \end{array} \end{aligned}$ | // Thread 2 |
| ThreadEnd; |  |
| ThreadStat $\mathrm{n}=\mathrm{m}$ 'm; | // Thread 3 |
| ThreadStat p = o'o; | // Thread 4 |
| ThreadJoin; | // waits for Threads 1-4 to finish |
| $\mathrm{b}=\mathrm{z}+\mathrm{r}+\mathrm{n}$ 'p; | // Using the results |

### 18.2 GAUSS Threading Concepts

This is really the one and only thing you need to know about threads: threads are separate sections of the same program, executing simultaneously, operating on the same data. In fact, it's so fundamental it's worth saying again: threads are separate sections of code in a program, running at the same time, using the same workspace, referencing and operating on the same symbols.

This raises basic issues of workflow and data integrity. How do you manage the creation and execution of threads, and make use of the work they do? And how do you maintain data integrity? (You do not want two threads assigning to the same symbol at the same time.)

To handle thread workflow, GAUSS employs a split-and-join approach. At various points in your program (as many as you like), you define a set of threads that will be created and run as a group. When created, the threads in the set execute simultaneously, each doing useful work. The parent thread waits for the created threads to complete, then continues, the results of their work now available for further use.

To maintain data integrity, we introduce the writer-must-isolate (informally, the any-thread-can-read-unless-some-thread-writes) programming rule. That is to say, symbols that are read from but not assigned to can be referenced by as many threads in a set as you like. Symbols that are assigned to, however, must be wholly owned by a single thread. No other thread in the set can reference that symbol. They cannot assign to it, nor can they read from it. They cannot refer to it at all.

Note: the writer-must-isolate rule only applies to the threads within a given set (including any child thread sets they may create). It does not apply between thread sets that have no chance of running simultaneously.

For threads defined in the main code, the writer-must-isolate rule applies to the global symbols. For threads defined in procedures or keywords, it applies to the global symbols, local symbols, and the procedure/keyword arguments.

### 18.3 Coding With Threads

There are two main points to coding with threads.
One-you can define threads anywhere. You can define them in the main code, you can define them in proc's and keyword's, and yes, you can define them inside other threads.

Two-you can call proc's and keyword's from threads. This is what really ties everything together. You can call a proc from a thread, and that proc can create threads, and any of those threads can call proc's, and any of those proc's can create threads, and ... you get the picture.

So-you can do things like this:

```
\(\mathrm{q}=\operatorname{chol}(\mathrm{b})\);
ThreadBegin;
    \(\mathrm{x}=\mathrm{q}+\mathrm{m}\);
    ThreadBegin;
        \(y=x\) ' \(x\);
        z = q'm;
    ThreadEnd;
    ThreadBegin;
        \(\mathrm{a}=\mathrm{b}+\mathrm{x}\);
        \(\mathrm{c}=\mathrm{a}+\mathrm{m}\);
    ThreadEnd;
    ThreadJoin;
    q = m'c;
ThreadEnd;
```

ThreadBegin;
ThreadStat $r=m$ 'm;
ThreadStat $s=m+i n v(b)$;
ThreadJoin;
t = r's;
ThreadEnd;
ThreadJoin;
$\mathrm{x}=\mathrm{r}+\mathrm{s}+\mathrm{q}+\mathrm{z}-\mathrm{t}$;

More importantly, you can do things like this:

```
proc bef(x);
    local y,t;
    ThreadStat y = nof(x);
    ThreadStat t = dof(x'x);
    ThreadJoin;
    t = t+y;
    retp(t);
endp;
proc abr(m);
    local x,y,z,a,b;
    a = m'm;
    ThreadStat x = inv(m);
    ThreadStat y = bef(m);
    ThreadStat z = dne(a);
    ThreadJoin;
    b = chut(x,y,z,a);
    retp(inv(b));
endp;
s = rndn(500,500);
```

```
ThreadStat t = abr(s);
ThreadStat q = abr(s^2);
ThreadStat r = che(s);
ThreadJoin;
w = del(t,q,r);
print w[1:10,1:10];
```

This means you can multi-thread anything you want, and call it from anywhere. You can multi-thread all the proc's and keyword's in your libraries, and call them freely anywhere in your multi-threaded programs.

### 18.4 Coding Restrictions

A few points on coding restrictions. First, you can't interlace thread definition statements and regular statements. You can't do this:

```
ThreadStat a = b'b;
n = q;
ThreadStat c = d'd;
ThreadJoin;
```

Or this:

```
if k == 1;
    ThreadStat a = b'b;
elseif k == 2;
    ThreadStat a = c'c;
endif;
if j == 1;
    ThreadStat d = e'e;
elseif j == 2;
    ThreadStat d = f'f;
endif;
ThreadJoin;
```

Each set of threads is defined as a group, and always completed by a ThreadJoin, like this:

```
n = q;
ThreadStat a = b'b;
ThreadStat c = d'd;
ThreadJoin;
```

And this:

```
ThreadBegin;
    if \(k=1\);
        \(\mathrm{a}=\mathrm{b}\) 'b;
        elseif k == 2;
            a = c'c;
    endif;
ThreadEnd;
ThreadBegin;
    if \(\mathrm{j}=1\);
        d = e'e;
        elseif j == 2;
            \(d=f\) 'f;
    endif;
ThreadEnd;
ThreadJoin;
```

Second-as stated above, you can reference read-only symbols in as many threads within a set as you like, but any symbols that are assigned to must be wholly owned by a single thread. A symbol that is assigned to by a thread cannot be written or read by any other thread in that set. This is the writer-must-isolate rule.

So, you can do this:

ThreadStat $x=y$ y;
ThreadStat $z=y+y$;

```
ThreadStat a = b-y;
ThreadJoin;
```

You cannot do this:

```
ThreadStat x = y'y;
Threadstat z = x'x;
ThreadStat a = b-y;
ThreadJoin;
```

This is because the threads within a set run simultaneously. Thus, there is no way of knowing when an assignment to a symbol has taken place, no way of knowing in one thread the "state" of a symbol in another.

Let's revisit the nested thread example for a minute and see how the writer-must-isolate rule applies to it:
$\mathrm{q}=$ chol(b); // main code, no threads yet
ThreadBegin; // Th1: isolates $x, y, z, a, c, q$ from Th2
$\mathrm{x}=\mathrm{q}+\mathrm{m}$;
ThreadBegin; // Th1.1: isolates $y, z$ from 1.2
$y=x$ 'x;
z = q'm;
ThreadEnd;
ThreadBegin; // Th1.2: isolates a,c from 1.1
$\mathrm{a}=\mathrm{b}+\mathrm{x} ;$
$\mathrm{c}=\mathrm{a}+\mathrm{m}$;
ThreadEnd;
ThreadJoin; // Joins 1.1, 1.2
$\mathrm{q}=\mathrm{m}$ ' ;
ThreadEnd;
ThreadBegin; // Th2: isolates r,s,t from Th1
ThreadStat $\mathrm{r}=\mathrm{m}$ 'm; // Th2.1: isolates r from 2.2
ThreadStat // Th2.2: isolates s from 2.1
$s=m+\operatorname{inv}(b) ;$

```
    ThreadJoin; // Joins 2.1, 2.1
    t = r's;
ThreadEnd;
ThreadJoin; // Joins Th1, Th2
x = r+S+q+z-t;
```

The main point here is that any symbols a thread or its children assign to must be isolated from all the other threads (and their children) of the same nesting level in that set. On the other hand, the children of a thread can freely read/write symbols that are read/written by their parent, because there is no risk of simultaneity; they must only isolate written symbols from their siblings and siblings' offspring.

If you break the writer-must-isolate rule, your program (and probably GAUSS) will crash. Worse, until it crashes, it will be happily producing indeterminate results.

Finally-the ThreadEnd command is what tells a thread to terminate, so you mustn't write code that keeps a thread from reaching it. For example, don't retp from the middle of a thread:

```
ThreadStat m = imt( 9 );
ThreadBegin;
    x = q[1];
    if x = 1;
        retp(z);
    else;
        r = z + 2;
    endif;
ThreadEnd;
ThreadJoin;
```

And don't use goto to jump into or out of the middle of a thread:

## retry:

ThreadBegin;
\{ err, x $\}=$ fna(q);
if err;

```
            goto badidea;
    endif;
    x = fnb(x);
ThreadEnd;
ThreadStat y = fnb(y);
ThreadJoin;
z = fnc(x,y);
save z;
end;
badidea:
errorlog "Error computing fna(q)";
q = fnd(q);
goto retry;
```

Basically, don't do anything that will keep a thread from reaching its ThreadEnd command. That's the only way it knows its work is done. end and stop are okay to call, though-they will bring the program to an end as usual, and terminate all running threads in the process.
(You can use goto and labels to jump around within a thread-that is, within the confines of a ThreadBegin/ThreadEnd pair.)

## Libraries

## 19

The GAUSS library system allows for the creation and maintenance of modular programs. The user can create "libraries" of frequently used functions that the GAUSS system will automatically find and compile whenever they are referenced in a program.

### 19.1 Autoloader

The autoloader resolves references to procedures, keywords, matrices, and strings that are not defined in the program from which they are referenced. The autoloader automatically locates and compiles the files containing the symbol definitions that are not resolved during the compilation of the main file. The search path used by the autoloader is first the current directory, and then the paths listed in the src_path configuration variable in the order they appear. src_path can be defined in the GAUSS configuration file.

### 19.1.1 Forward References

When the compiler encounters a symbol that has not previously been defined, that is called a "forward reference". GAUSS handles forward references in two ways, depending on whether they are "left-hand side" or "right-hand side" references.

## Left-Hand Side

A left-hand side reference is usually a reference to a symbol on the left-hand side of the equal sign in an expression.

$$
x=5 ;
$$

Left-hand side references, since they are assignments, are assumed to be matrices. In the statement above, $\mathbf{x}$ is assumed to be a matrix and the code is compiled accordingly. If, at execution time, the expression actually returns a string, the assignment is made and the type of the symbol $\mathbf{x}$ is forced to string.

Some commands are implicit left-hand side assignments. There is an implicit left-hand side reference to $\mathbf{x}$ in each statement below:

```
clear x;
load x;
open x = myfile;
```


## Right-Hand Side

A right-hand side reference is usually a reference to a symbol on the right-hand side of the equal sign in an expression such as:

```
z = 6;
y = z + dog;
print y;
```

In the program above, since dog is not previously known to the compiler, the autoloader will search for it in the active libraries. If it is found, the file containing it will be compiled. If it is not found in a library, the autoload/autodelete state will determine how it is handled.

### 19.1.2 The Autoloader Search Path

If the autoloader is OFF, no forward references are allowed. Every procedure, matrix, and string referenced by your program must be defined before it is referenced. An external statement can be used above the first reference to a symbol, but the definition of the symbol must be in the main file or in one of the files that are \#include'd. No global symbols are deleted automatically.

If the autoloader is ON, GAUSS searches for unresolved symbol references during compilation using a specific search path as outlined below. If the autoloader is OFF, an Undefined symbol error message will result for right-hand side references to unknown symbols.

When autoload is ON, the autodelete state controls the handling of references to unknown symbols.

The following search path will be followed to locate any symbols not previously defined:

## Autodelete ON

1. user library
2. user-specified libraries.
3. gauss library
4. current directory, then src_path for files with a .g extension.

Forward references are allowed and .g files need not be in a library. If there are symbols that cannot be found in any of the places listed above, an Undefined symbol error message will be
generated and all uninitialized variables and all procedures with global references will be deleted from the global symbol table. This autodeletion process is transparent to the user, since the symbols are automatically located by the autoloader the next time the program is run. This process results in more compile time, which may or may not be significant, depending on the speed of the computer and the size of the program.

## Autodelete OFF

1. user library
2. user-specified libraries.
3. gauss library

All .g files must be listed in a library. Forward references to symbols that are not listed in an active library are not allowed. For example:

```
x = rndn(10,10);
y = sym(x); /* Forward reference to sym */
proc sym(x);
    retp(x+x');
endp;
```

Use an external statement for anything referenced above its definition if autodelete is OFF:

```
external proc sym;
```

$\mathrm{x}=\operatorname{rndn}(10,10)$;
$\mathrm{y}=\operatorname{sym}(\mathrm{x})$;
proc sym(x);
$\operatorname{retp}(x+x$ ') ;
endp;

When autodelete is OFF, symbols not found in an active library will not be added to the symbol table. This prevents the creation of uninitialized procedures in the global symbol table. No deletion of symbols from the global symbol table will take place.

## Libraries

The first place GAUSS looks for a symbol definition is in the "active" libraries. A GAUSS library is a text file that serves as a dictionary to the source files that contain the symbol definitions. When a library is active, GAUSS will look in it whenever it is looking for a symbol it is trying to resolve. The library statement is used to make a library active. Library files should be located in the subdirectory listed in the lib_path configuration variable. Library files have an .lcg extension.

Suppose you have several procedures that are all related and you want them all defined in the same file. You can create such a file, and, with the help of a library, the autoloader will be able to find the procedures defined in that file whenever they are called.

First, create the file that is to contain your desired procedure definitions. By convention, this file is usually named with a . src extension, but you may use any name and any file extension. In this file, put all the definitions of related procedures you wish to use. Here is an example of such a file. It is called norm. src:

```
/*
** norm.src
**
** This is a file containing the definitions of three
** procedures which return the norm of a matrix x.
** The three norms calculated are the 1-norm, the
** inf-norm and the E-norm.
*/
proc onenorm(x);
    retp(maxc(sumc(abs(x))));
endp;
proc infnorm(x);
```

```
    retp(maxc(sumc(abs(x'))));
endp;
proc Enorm(x);
    retp(sumc(sumc(x.*x)));
endp;
```

Next, create a library file that contains the name of the file you want access to, and the list of symbols defined in it. This can be done with the lib command. (For details, see lib in the GAUSS Language Reference.)

A library file entry has a filename that is flush left. The drive and path can be included to speed up the autoloader. Indented below the filename are the symbols included in the file. There can be multiple symbols listed on a line, with spaces between. The symbol type follows the symbol name, with a colon delimiting it from the symbol name. The valid symbol types are:

| fn | user-defined single line function. |
| :--- | :--- |
| keyword | keyword. |
| proc | procedure. |
| matrix | matrix, numeric or character. |
| array | N-dimensional array. |
| string | string. |
| sparse matrix | sparse matrix. |
| struct | structure. |

A structure is always denoted by struct followed by the structure type name.
If the symbol type is missing, the colon must not be present and the symbol type is assumed to be proc. Both library files below are valid:

Example 1

```
/*
** math
**
** This library lists files and procedures for mathematical routines.
*/
norm.src
    onenorm:proc infnorm:proc Enorm:proc
complex.src
    cmmult:proc cmdiv:proc cmadd:proc cmsoln:proc
poly.src
    polychar:proc polyroot:proc polymult:proc
```


## Example 2

```
/*
** math
**
*/
c:\gauss\src\norm.src
    onenorm : proc
    infnorm : proc
    Enorm : proc
c:\gauss\src\complex.src
    cmmult : proc
    cmdiv : proc
    cmadd : proc
    cmsoln : proc
c:\gauss\src\fcomp.src
    feq : proc
    fne : proc
    flt : proc
    fgt : proc
    fle : proc
```

** This library lists files and procedures for mathematical routines.

fge : proc<br>$\mathrm{c}: \backslash$ gauss $\backslash \mathrm{src} \backslash$ fcomp.dec<br>_fcmptol : matrix

Once the autoloader finds, via the library, the file containing your procedure definition, everything in that file will be compiled. For this reason, you should combine related procedures in the same file in order to minimize the compiling of procedures not needed by your program. In other words, you should not combine unrelated functions in one . src file because if one function in a . src file is needed, the whole file will be compiled.

## user Library

This is a library for user-created procedures. If the autoloader is ON, the user library is the first place GAUSS looks when trying to resolve symbol references.

You can update the user library with the lib command as follows:

> lib user myfile.src

This will update the user library by adding a reference to myfile.src.
No user library is shipped with GAUSS. It will be created the first time you use the lib command to update it.

For details on the parameters available with the lib command, see the GAUSS Language Reference.

## .g Files

If autoload and autodelete are ON and a symbol is not found in a library, the autoloader will assume it is a procedure and look for a file that has the same name as the symbol and a.g extension. For example, if you have defined a procedure called square, you could put the definition in a file called square.g in one of the subdirectories listed in your src_path. If autodelete is OFF, the .g file must be listed in an active library; for example, in the user library.

### 19.2 Global Declaration Files

If your application makes use of several global variables, create a file containing declare statements. Use files with the extension .dec to assign default values to global matrices and strings with declare statements and to declare global N-dimensional arrays, sparse matrices, and structures, which will be initialized as follows:

| Variable Type | Initializes To |
| :--- | :--- |
| N-dimensional array | 1 -dimensional array of 1 containing 0 |
| sparse matrix | empty sparse matrix |
| structure | $1 \times 1$ structure containing empty and/or zeroed out members |

In order to declare structures in a . dec file, you must \#include the file(s) containing the definitions of the types of structures that you wish to declare at the top of your . dec file. For example, if you have the following structure type definition in a file called mystruct. sdf:

```
struct mystruct {
    matrix m;
    array a;
    scalar scal;
    string array sa;
};
```

You could declare an instance of that structure type, called ms, in a . dec file as follows:
\#include mystruct.sdf
declare struct mystruct ms;

See declare in the Command Reference, Chapter 33, for more information.
A file with a . ext extension containing the same symbols in external statements can also be created and \#include'd at the top of any file that references these global variables. An
appropriate library file should contain the name of the . dec files and the names of the globals they declare. This allows you to reference global variables across source files in an application.

Here is an example that illustrates the way in which . dec, .ext, .lcg and .src files work together. Always begin the names of global matrices or strings with '_, to distinguish them from procedures.
.src File:

```
/*
** fcomp.src
**
** These functions use _fcmptol to fuzz the comparison operations
** to allow for roundoff error.
**
** The statement: y = feq(a,b);
**
** is equivalent to: y = a eq b;
**
** Returns a scalar result, 1 (true) or 0 (false)
**
** y = feq(a,b);
** y = fne(a,b);
*/
#include fcomp.ext
proc feq(a,b);
    retp(abs(a-b) <= _fcmptol);
endp;
proc fne(a,b);
    retp(abs(a-b) > _fcmptol);
endp;
```

. dec File:

```
/*
** fcomp.dec - global declaration file for fuzzy comparisons.
*/
declare matrix _fcmptol != 1e-14;
```

.ext File:

```
/*
** fcomp.ext - external declaration file for fuzzy comparisons.
*/
external matrix _fcmptol;
```

.lcg File:

```
/*
** fcomp.lcg - fuzzy compare library
*/
fcomp.dec
    _fcmptol:matrix
fcomp.src
    feq:proc
    fne:proc
```

With the exception of the library (.lcg) files, these files must be located along your src_path. The library files must be on your lib_path. With these files in place, the autoloader will be able to find everything needed to run the following programs:

```
library fcomp;
x = rndn(3,3);
xi = inv(x);
```

```
xix = xi*x;
if feq(xix,eye(3));
    print "Inverse within tolerance.";
else;
    print "Inverse not within tolerance.";
endif;
```

If the default tolerance of $1 \mathrm{e}-14$ is too tight, the tolerance can be relaxed:

```
library fcomp;
x = rndn(3,3);
xi = inv(x);
xix = xi*x;
_fcmptol = 1e-12; /* reset tolerance */
if feq(xix,eye(3));
    print "Inverse within tolerance.";
else;
    print "Inverse not within tolerance.";
endif;
```


### 19.3 Troubleshooting

Below is a partial list of errors you may encounter in using the library system, followed by the most probable cause.
(4) : error G0290 : '/gauss/lib/prt.lcg' : Library not found

The autoloader is looking for a library file called prt.lcg, because it has been activated in a library statement. Check the subdirectory listed in your lib_path configuration variable for a file called prt.lcg.
(0) : error G0292 : 'prt.dec' : File listed in library not found

The autoloader cannot find a file called prt. dec. Check for this file. It should exist somewhere along your src_path, if you have it listed in prt.lcg.

## Undefined symbols: <br> PRTVEC /gauss/src/tstprt.g(2)

The symbol prtvec could not be found. Check if the file containing prtvec is in the src_path. You may have not activated the library that contains your symbol definition. Do so in a library statement.

```
/gauss/src/prt.dec(3) : Redefinition of '__vnames' (proc)__vnames being
    declared external matrix
```

You are trying to illegally force a symbol to another type. You probably have a name conflict that needs to be resolved by renaming one of the symbols.

```
/gauss/lib/prt.lcg(5) : error G0301 : 'prt.dec' : Syntax error in
    library
Undefined symbols:
    __VNAMES /gauss/src/prt.src(6)
```

Check your library to see that all filenames are flush left and that all the symbols defined in that file are indented by at least one space.

### 19.3.1 Using .dec Files

Below is some advice you are encouraged to follow when constructing your own library system:

- Whenever possible, declare variables in a file that contains only declare statements. When your program is run again without clearing the workspace, the file containing the variable declarations will not be compiled and declare warnings will be prevented.
- Provide a function containing regular assignment statements to reinitialize the global variables in your program if they ever need to be reinitialized during or between runs. Put this in a separate file from the declarations:

```
proc (0) = globset;
    _vname = "X";
    _con = 1;
    _row = 0;
    _title = "";
endp;
```

- Never declare any global in more than one file.
- To avoid meaningless redefinition errors and declare warnings, never declare a global more than once in any one file. Redefinition error messages and declare warnings are meant to help you prevent name conflicts, and will be useless to you if your code generates them normally.

By following these guidelines, any declare warnings and redefinition errors you get will be meaningful. By knowing that such warnings and errors are significant, you will be able to debug your programs more efficiently.

## Compiler

20

GAUSS allows you to compile your large, frequently used programs to a file that can be run over and over with no compile time. The compiled image is usually smaller than the uncompiled source. GAUSS is not a native code compiler; rather, it compiles to a form of pseudocode. The file will have a .gcg extension.

The compile command will compile an entire program to a compiled file. An attempt to edit a compiled file will cause the source code to be loaded into the editor if it is available to the system. The run command assumes a compiled file if no extension is given, and that a file with a.gcg extension is in the src_path. A saveall command is available to save the current contents of memory in a compiled file for instant recall later. The use command will instantly load a compiled program or set of procedures at the beginning of an ASCII program before compiling the rest of the ASCII program file.

Since the compiled files are encoded binary files, the compiler is useful for developers who do not want to distribute their source code.

### 20.1 Compiling Programs

Programs are compiled with the compile command.

### 20.1.1 Compiling a File

Source code program files that can be run with the run command can be compiled to .gcg files with the compile command:

```
compile qxy.e;
```

All procedures, global matrices, arrays, strings and string arrays, and the main program segment will be saved in the compiled file. The compiled file can be run later using the run command. Any libraries used in the program must be present and active during the compile, but not when the program is run. If the program uses the dlibrary command, the . dll files must be present when the program is run and the dlibrary path must be set to the correct subdirectory. This will be handled automatically in your configuration file. If the program is run on a different computer than it was compiled on, the . dll files must be present in the correct location. sysstate (case 24) can be used to set the dlibrary path at run-time.

### 20.2 Saving the Current Workspace

The simplest way to create a compiled file containing a set of frequently used procedures is to use saveall and an external statement:
library pgraph;
external proc xy,logx,logy,loglog,hist;
saveall pgraph;

Just list the procedures you will be using in an external statement and follow it with a saveall statement. It is not necessary to list procedures that you do not explicitly call, but are called from
another procedure, because the autoloader will automatically find them before the saveall command is executed. Nor is it necessary to list every procedure you will be calling, unless the source will not be available when the compiled file is use'd.

Remember, the list of active libraries is NOT saved in the compiled file, so you may still need a library statement in a program that is use'ing a compiled file.

### 20.3 Debugging

If you are using compiled code in a development situation in which debugging is important, compile the file with line number records. After the development is over, you can recompile without line number records if the maximum possible execution speed is important. If you want to guarantee that all procedures contain line number records, put a new statement at the top of your program and turn line number tracking on.


## File I/O

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The following is a partial list of the I/O commands in the GAUSS programming language:

| close | Close a file. |
| :--- | :--- |
| closeall | Close all open files. |
| colsf | Number of columns in a file. |
| create | Create GAUSS data set. |
| eof | Test for end of file. |
| fcheckerr | Check error status of a file. |
| fclearerr | Check error status of a file and clear error flag. |
| $\mathbf{f f l u s h}$ | Flush a file's output buffer. |
| fgets | Read a line of text from a file. |
| fgetsa | Read multiple lines of text from a file. |


| fgetsat | Read multiple lines of text from a file, discarding newlines. |
| :--- | :--- |
| fgetst | Read a line of text from a file, discarding newline. |
| fileinfo | Return names and information of files matching a specification. |
| files | Return a directory listing as a character matrix. |
| filesa | Return a list of files matching a specification. |
| fopen | Open a file. |
| fputs | Write strings to a file. |
| fputst | Write strings to a file, appending newlines. |
| fseek | Reposition file pointer. |
| fstrerror | Get explanation of last file I/O error. |
| ftell | Get position of file pointer. |
| getf | Load a file into a string. |
| getname | Get variable names from data set. |
| iscplxf | Return whether a data set is real or complex. |
| load | Load matrix file or small ASCII file (same as loadm). |
| loadd | Load a small GAUSS data set into a matrix. |
| loadm | Load matrix file or small ASCII file. |
| loads | Load string file. |
| open | Open a GAUSS data set. |
| output | Control printing to an auxiliary output file or device. |
| readr | Read a specified number of rows from a file. |
| rowsf | Number of rows in file. |
| save | Save matrices, strings, procedures. |

saved Save a matrix in a GAUSS data set.
seekr $\quad$ Reset read/write pointer in a data set.
sortd Sort a data set.
typef $\quad$ Return type of data set (bytes per element).
writer Write data to a data set.

### 21.1 ASCII Files

GAUSS has facilities for reading and writing ASCII files. Since most software can also read and write ASCII files, this provides one method of sharing data between GAUSS and many other kinds of programs.

### 21.1.1 Matrix Data

## Reading

Files containing numeric data that are delimited with spaces or commas and are small enough to fit into a single matrix or string can be read with load. Larger ASCII data files can be converted to GAUSS data sets with the ATOG utility program (see ATOG, Chapter 28). ATOG can convert packed ASCII files as well as delimited files.

For small delimited data files, the load statement can be used to load the data directly into a GAUSS matrix. The resulting GAUSS matrix must be no larger than the limit for a single matrix.

For example,

$$
\text { load } x[]=\text { dat1.asc; }
$$

will load the data in the file dat1. asc into an $N \times 1$ matrix $\mathbf{x}$. This method is preferred because rows ( $\mathbf{x}$ ) can be used to determine how many elements were actually loaded, and the matrix can be reshape'd to the desired form:

```
load x[] = dat1.asc;
if rows(x) eq 500;
    x = reshape(x,100,5);
else;
    errorlog "Read Error";
    end;
endif;
```

For quick interactive loading without error checking, use

```
load x[100,5] = dat1.asc;
```

This will load the data into a $100 \times 5$ matrix. If there are more or fewer than 500 numbers in the data set, the matrix will automatically be reshaped to $100 \times 5$.

## Writing

To write data to an ASCII file the print or printfm command is used to print to the auxiliary output. The resulting files are standard ASCII files and can be edited with GAUSS's editor or another text editor.

The output and outwidth commands are used to control the auxiliary output. The print or printfm command is used to control what is sent to the output file.

The window can be turned on and off using screen. When printing a large amount of data to the auxiliary output, the window can be turned off using the command

```
screen off;
```

This will make the process much faster, especially if the auxiliary output is a disk file.
It is easy to forget to turn the window on again. Use the end statement to terminate your programs; end will automatically perform screen on and output off.

The following commands can be used to control printing to the auxiliary output:

| format | Specify format for printing a matrix. |
| :--- | :--- |
| output | Open, close, rename auxiliary output file or device. |
| outwidth | Set auxiliary output width. |
| printfm | Formatted matrix print. |
| print | Print matrix or string. |
| screen | Turn printing to the window on and off. |

This example illustrates printing a matrix to a file:

```
format /rd 8,2;
outwidth 132;
output file = myfile.asc reset;
screen off;
print x;
output off;
screen on;
```

The numbers in the matrix $\mathbf{x}$ will be printed with a field width of 8 spaces per number, and with 2 places beyond the decimal point. The resulting file will be an ASCII data file. It will have 132 column lines maximum.

A more extended example follows. This program will write the contents of the GAUSS file mydata.dat into an ASCII file called mydata. asc. If there is an existing file by the name of mydata.asc, it will be overwritten:

```
output file = mydata.asc reset;
screen off;
format /rd 1,8;
open fp = mydata;
do until eof(fp);
    print readr(fp,200);;
```

```
endo;
fp = close(fp);
end;
```

The output ... reset command will create an auxiliary output file called mydata.asc to receive the output. The window is turned off to speed up the process. The GAUSS data file mydata.dat is opened for reading and 200 rows are read per iteration until the end of the file is reached. The data read are printed to the auxiliary output mydata. asc only, because the window is off.

### 21.1.2 General File I/O

getf will read a file and return it in a string variable. Any kind of file can be read in this way as long as it will fit into a single string variable.

To read files sequentially, use fopen to open the file and use fgets, fputs, and associated functions to read and write the file. The current position in a file can be determined with ftell. The following example uses these functions to copy an ASCII text file:

```
proc copy(src, dest);
    local fin, fout, str;
    fin = fopen(src, "rb");
    if not fin;
        retp(1);
    endif;
    fout = fopen(dest, "wb");
    if not fin;
        call close(fin);
        retp(2);
    endif;
    do until eof(fin);
```

```
            str = fgets(fin, 1024);
            if fputs(fout, str) /= 1;
            call close(fin);
            call close(fout);
            retp(3);
            endif;
        endo;
        call close(fin);
        call close(fout);
        retp(0);
endp;
```


### 21.2 Data Sets

GAUSS data sets are the preferred method of storing data contained in a single matrix for use within GAUSS. Use of these data sets allows extremely fast reading and writing of data. Many library functions are designed to read data from these data sets.

If you want to store multiple variables of various types in a single file, see GAUSS Data Archives, Section 21.3.

### 21.2.1 Layout

GAUSS data sets are arranged as matrices; that is, they are organized in terms of rows and columns. The columns in a data file are assigned names, and these names are stored in the header, or, in the case of the v89 format, in a separate header file.

The limit on the number of rows in a GAUSS data set is determined by disk size. The limit on the number of columns is limited by RAM. Data can be stored in 2,4 , or 8 bytes per number, rather than just 8 bytes as in the case of GAUSS matrix files.

The ranges of the different formats are:

| Bytes | Type | Significant Digits | Range |
| :--- | :--- | :--- | :---: |
|  |  |  |  |
| 2 | integer | 4 | $-32768<=\mathrm{X}<=32767$ |
| 4 | single | $6-7$ | $8.43 \mathrm{E}-37<=\|\mathrm{X}\|<=3.37 \mathrm{E}+38$ |
| 8 | double | $15-16$ | $4.19 \mathrm{E}-307<=\|\mathrm{X}\|<=1.67 \mathrm{E}+308$ |

### 21.2.2 Creating Data Sets

Data sets can be created with the create or datacreate command. The names of the columns, the type of data, etc., can be specified. (For details, see create in the GAUSS Language Reference.)

Data sets, unlike matrices, cannot change from real to complex, or vice-versa. Data sets are always stored a row at a time. The rows of a complex data set, then, have the real and imaginary parts interleaved, element by element. For this reason, you cannot write rows from a complex matrix to a real data set-there is no way to interleave the data without rewriting the entire data set. If you must, explicitly convert the rows of data first, using the real and imag functions (see the GAUSS Language Reference), and then write them to the data set. Rows from a real matrix CAN be written to a complex data set; GAUSS simply supplies 0's for the imaginary part.

To create a complex data set, include the complex flag in your create command.

### 21.2.3 Reading and Writing

The basic functions in GAUSS for reading data files are open and readr:

```
open f1 = dat1;
x = readr(f1,100);
```

The call to readr in this example will read in 100 rows from dat1. dat. The data will be assigned to a matrix $\mathbf{x}$.
loadd and saved can be used for loading and saving small data sets.

The following example illustrates the creation of a GAUSS data file by merging (horizontally concatenating) two existing data sets:

```
file1 = "dat1";
file2 = "dat2";
outfile = "daty";
open fin1 = `file1 for read;
open fin2 = ` file2 for read;
varnames = getname(file1)|getname(file2);
otyp = maxc(typef(fin1)|typef(fin2));
create fout = `outfile with ^varnames,0,otyp;
nr = 400;
do until eof(fin1) or eof(fin2);
    y1 = readr(fin1,nr);
    y2 = readr(fin2,nr);
    r = maxc(rows(y1)|rows(y2));
    y = y1[1:r,.] ~ y2[1:r,.];
    call writer(fout,y);
endo;
closeall fin1,fin2,fout;
```

In this example, data sets dat1.dat and dat2. dat are opened for reading. The variable names from each data set are read using getname, and combined in a single vector called varnames. A variable called otyp is created, which will be equal to the larger of the two data types of the input files. This will insure that the output is not rounded to less precision than the input files. A new data set daty.dat is created using the create ... with ... command. Then, on every iteration of the loop, 400 rows are read in from each of the two input data sets, horizontally concatenated, and written out to daty.dat. When the end of one of the input files is reached, reading and writing will stop. The closeall command is used to close all files.

### 21.2.4 Distinguishing Character and Numeric Data

Although GAUSS itself does not distinguish between numeric and character columns in a matrix or data set, some of the GAUSS Application programs do. When creating a data set, it is important to indicate the type of data in the various columns. The following discusses two ways of doing this.

## Using Type Vectors

The v89 data set format distinguished between character and numeric data in data sets by the case of the variable names associated with the columns. The v96 data set format, however, stores this type information separately, resulting in a much cleaner and more robust method of tracking variable types, and greater freedom in the naming of data set variables.

When you create a data set, you can supply a vector indicating the type of data in each column of the data set. For example:

```
data = { M 32 21500,
    F 27 36000,
    F 28 19500,
    M 25 32000 };
vnames = { "Sex" "Age" "Pay" };
vtypes = { 0 1 1 };
create f = mydata with ^vnames, 3, 8, vtypes;
call writer(f,data);
f = close(f);
```

To retrieve the type vector, use vartypef.

```
open f = mydata for read;
vn = getnamef(f);
vt = vartypef(f);
print vn';
print vt';
\begin{tabular}{ccc} 
Sex & Age & Pay \\
0 & 1 & 1
\end{tabular}
```

The call to getnamef in this example returns a string array rather than a character vector, so you can print it without the '\$' prefix.

## Using the Uppercase/Lowercase Convention (v89 Data Sets)

Historically, some GAUSS Application programs recognized an "uppercase/lowercase" convention: if the variable name was uppercase, the variable was assumed to be numeric, and if it was lowercase, the variable was assumed to be character.

However, this is now obsolete; use vartypef and v96 data sets to be compatible with future versions.

### 21.3 GAUSS Data Archives

The GAUSS Data Archive (GDA) is extremely powerful and flexible, giving you much greater control over how you store your data. There is no limitation on the number of variables that can be stored in a GDA, and the only size limitation is the amount of available disk space. Moreover, GDA's are designed to hold whatever type of data you want to store in them. You may write matrices, arrays, strings, string arrays, sparse matrices, and structures to a GDA, and the GDA will keep track of the type, size and location of each of the variables contained in it. Since GAUSS now supports reading and writing to GDA's that were created on other platforms, GDA's provide a simple solution to the problem of sharing data across platforms.

See Section 21.5.12 for information on the layout of a GDA.

### 21.3.1 Creating and Writing Variables to GDA's

To create a GAUSS Data Archive, call gdaCreate, which creates a GDA containing only header information. It is recommended that file names passed into gdaCreate have a . gda extension; however, gdaCreate will not force an extension.

To write variables to the GDA, you must call gdaWrite. A single call to gdaWrite writes only one variable to the GDA. Writing multiple variables requires multiple calls to gdaWrite.

For example, the following code:

```
ret = gdaCreate("myfile.gda",1);
```

```
ret = gdaWrite("myfile.gda",rndn(100,50),"x1");
ret = gdaWrite("myfile.gda","This is a string","str1");
ret = gdaWrite('myfile.gda",394,"x2");
```

produces a GDA containing the following variables:
Index Name Type Size

| 1 | x 1 | matrix | $100 \times 50$ |
| :--- | :--- | :--- | :--- |
| 2 | str 1 | string | 16 chars |
| 3 | x 2 | matrix | $1 \times 1$ |

### 21.3.2 Reading Variables from GDA's

The following table details the commands that you may use to read various types of variables from a GAUSS Data Archive:

| Variable Type | Read Command(s) |
| :--- | :--- |
| matrix <br> array <br> string <br> string array | gdaRead <br> gdaReadByIndex |
| sparse matrix | gdaReadSparse |
| structure | gdaReadStruct |

gdaRead, gdaReadSparse, and gdaReadStruct take a variable name and return the variable data. gdaReadByIndex returns the variable data for a specified variable index.

For example, to get the variable $\mathbf{x 1}$ out of myfile.gda, you could call:
y = gdaRead("myfile.gda", "x1");
or
y = gdaReadByIndex("myfile.gda", 1);

If you want to read only a part of a matrix, array, string, or string array from a GDA, call gdaReadSome. Sparse matrices and structures may not be read in parts.

### 21.3.3 Updating Variables in GDA's

To overwrite an entire variable in a GDA, you may call gdaUpdate or gdaUpdateAndPack. If the new variable is not the same size as the variable that it is replacing, gdaUpdate will leave empty bytes in the file, while gdaUpdateAndPack will pack the file (from the location of the variable that is being replaced to the end of the file) to remove those empty bytes.
gdaUpdate is usually faster, since it does not move data in the file unnecessarily. However, calling gdaUpdate several times for one file may result in a file with a large number of empty bytes.

On the other hand, gdaUpdateAndPack uses disk space efficiently, but it may be slow for large files (especially if the variable to be updated is one of the first variables in the file).

If speed and disk space are both concerns and you are going to update several variables, it will be most efficient to use gdaUpdate to update the variables and then call gdaPack once at the end to pack the file.

The syntax is the same for both gdaUpdate and gdaUpdateAndPack:

```
ret = gdaUpdate("myfile.gda",rndn(1000,100),"x1");
ret = gdaUpdateAndPack("myfile.gda",rndn(1000,100),"x1");
```

To overwrite part of a variable in a GDA, call gdaWriteSome.

### 21.4 Matrix Files

GAUSS matrix files are files created by the save command.

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The save command takes a matrix in memory, adds a header that contains information on the number of rows and columns in the matrix, and stores it on disk. Numbers are stored in double precision just as they are in matrices in memory. These files have the extension .fmt.

Matrix files can be no larger than a single matrix. No variable names are associated with matrix files.

GAUSS matrix files can be load'ed into memory using the load or loadm command or they can be opened with the open command and read with the readr command. With the readr command, a subset of the rows can be read. With the load command, the entire matrix is load'ed.

GAUSS matrix files can be open'ed for read, but not for append, or for update.

If a matrix file has been opened and assigned a file handle, rowsf and colsf can be used to determine how many rows and columns it has without actually reading it into memory. seekr and readr can be used to jump to particular rows and to read them into memory. This is useful when only a subset of rows is needed at any time. This procedure will save memory and be much faster than load'ing the entire matrix into memory.

### 21.5 File Formats

This section discusses the GAUSS binary file formats.

There are four currently supported matrix file formats:
Version Extension Support
Small Matrix v89 .fmt Obsolete, use v96.

Extended Matrix v89 .fmt Obsolete, use v96.
Matrix v92 .fmt Obsolete, use v96.
Universal Matrix v96 .fmt Supported for read/write.

There are four currently supported string file formats:

| Version | Extension | Support |
| :--- | :--- | :--- |
|  |  |  |
| Small String v89 | . fst | Obsolete, use v96. |
| Extended String v89 | . fst | Obsolete, use v96. |
| String v92 | . fst | Obsolete, use v96. |
| Universal String v96 | .fst | Supported for read/write. |

There are four currently supported data set formats:

| Version | Extension | Support |
| :---: | :---: | :---: |
| Small Data Set v89 | .dat, <br> .dht | Obsolete, use v96. |
| Extended Data Set v89 | .dat, <br> .dht | Obsolete, use v96. |
| Data Set v92 | .dat | Obsolete, use v96. |
| Universal Data Set v96 | . dat | Supported for read/write. |

### 21.5.1 Small Matrix v89 (Obsolete)

Matrix files are binary files, and cannot be read with a text editor. They are created with save. Matrix files with up to 8190 elements have a .fmt extension and a 16-byte header formatted as follows:

## Offset Description

0-1 DDDD hex, identification flag
2-3 rows, unsigned 2-byte integer
4-5 columns, unsigned 2-byte integer
6-7 size of file minus 16-byte header, unsigned 2-byte integer
8-9 type of file, 0086 hex for real matrices, 8086 hex for complex matrices
10-15 reserved, all 0's
The body of the file starts at offset 16 and consists of IEEE format double precision floating point numbers or character elements of up to 8 characters. Character elements take up 8 bytes and are padded on the right with zeros. The size of the body of the file is $8 *$ rows $*$ cols rounded up to the next 16-byte paragraph boundary. Numbers are stored row by row. A $2 \times 3$ real matrix will be
stored on disk in the following way, from the lowest addressed element to the highest addressed element:

$$
[1,1] \quad[1,2] \quad[1,3] \quad[2,1] \quad[2,2] \quad[2,3]
$$

For complex matrices, the size of the body of the file is $16 *$ rows*cols. The entire real part of the matrix is stored first, then the entire imaginary part. A $2 \times 3$ complex matrix will be stored on disk in the following way, from the lowest addressed element to the highest addressed element:

| (real part) | $[1,1]$ | $[1,2]$ | $[1,3]$ | $[2,1]$ | $[2,2]$ | $[2,3]$ |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| (imaginary part) | $[1,1]$ | $[1,2]$ | $[1,3]$ | $[2,1]$ | $[2,2]$ | $[2,3]$ |

### 21.5.2 Extended Matrix v89 (Obsolete)

Matrices with more than 8190 elements are saved in an extended format. These files have a 16-byte header formatted as follows:

## Offset Description

0-1 EEDD hex, identification flag
2-3 type of file, 0086 hex for real matrices, 8086 hex for complex matrices
4-7 rows, unsigned 4-byte integer
8-11 columns, unsigned 4-byte integer
12-15 size of file minus 16-byte header, unsigned 4-byte integer
The size of the body of an extended matrix file is $8 *$ rows* cols (not rounded up to a paragraph boundary). Aside from this, the body is the same as the small matrix v89 file.

### 21.5.3 Small String v89 (Obsolete)

String files are created with save. String files with up to 65519 characters have a 16-byte header formatted as follows:

## Offset Description

0-1 DFDF hex, identification flag
2-3 1, unsigned 2-byte integer
4-5 length of string plus null byte, unsigned 2-byte integer
6-7 size of file minus 16-byte header, unsigned 2-byte integer
8-9 001D hex, type of file
10-15 reserved, all 0's
The body of the file starts at offset 16. It consists of the string terminated with a null byte. The size of the file is the 16 -byte header plus the length of the string and null byte rounded up to the next 16-byte paragraph boundary.

### 21.5.4 Extended String v89 (Obsolete)

Strings with more than 65519 characters are saved in an extended format. These files have a 16-byte header formatted as follows:

## Offset Description

0-1 EEDF hex, identification flag
2-3 001D hex, type of file
4-7 1, unsigned 4-byte integer
8-11 length of string plus null byte, unsigned 4-byte integer
12-15 size of file minus 16-byte header, unsigned 4-byte integer
The body of the file starts at offset 16. It consists of the string terminated with a null byte. The size of the file is the 16 -byte header plus the length of the string and null byte rounded up to the next 8 -byte boundary.

### 21.5.5 Small Data Set v89 (Obsolete)

All data sets are created with create. v89 data sets consist of two files; one . dht contains the header information; the second (.dat) contains the binary data. The data will be one of three types:

8-byte IEEE floating point<br>4-byte IEEE floating point<br>2-byte signed binary integer, twos complement

Numbers are stored row by row.
The .dht file is used in conjunction with the . dat file as a descriptor file and as a place to store names for the columns in the . dat file. Data sets with up to 8175 columns have a . dht file formatted as follows:

## Offset Description

0-1 DADA hex, identification flag
2-5 reserved, all 0's
6-7 columns, unsigned 2-byte integer
8-9 row size in bytes, unsigned 2-byte integer
10-11 header size in bytes, unsigned 2-byte integer
12-13 data type in . dat file (2 48 ), unsigned 2-byte integer
14-17 reserved, all 0's
18-21 reserved, all 0's
22-23 control flags, unsigned 2-byte integer
24-127 reserved, all 0's
Column names begin at offset 128 and are stored 8 bytes each in ASCII format. Names with less than 8 characters are padded on the right with bytes of 0 .

The number of rows in the . dat file is calculated in GAUSS using the file size, columns, and data type. This means that users can modify the . dat file by adding or deleting rows with other software without updating the header information.

Names for the columns should be lowercase for character data, to be able to distinguish them from numeric data with vartype.

GAUSS currently examines only the 4 's bit of the control flags. This bit is set to 0 for real data sets, 1 for complex data sets. All other bits are 0 .

Data sets are always stored a row at a time. A real data set with 2 rows and 3 columns will be stored on disk in the following way, from the lowest addressed element to the highest addressed
element:

| $[1,1]$ | $[1,2]$ | $[1,3]$ |
| :--- | :--- | :--- |
| $[2,1]$ | $[2,2]$ | $[2,3]$ |

The rows of a complex data set are stored with the real and imaginary parts interleaved, element by element. A $2 \times 3$ complex data set, then, will be stored on disk in the following way, from the lowest addressed element to the highest addressed element:

| $[1,1] r$ | $[1,1] i$ | $[1,2] r$ | $[1,2] i$ | $[1,3] r$ | $[1,3] i$ |
| :--- | :--- | :--- | :--- | :--- | :--- |
| $[2,1] r$ | $[2,1] i$ | $[2,2] r$ | $[2,2] i$ | $[2,3] r$ | $[2,3] i$ |

### 21.5.6 Extended Data Set v89 (Obsolete)

Data sets with more than 8175 columns are saved in an extended format that cannot be read by the 16-bit version. These files have a . dht descriptor file formatted as follows:

## Offset Description

0-1 EEDA hex, identification flag
2-3 data type in . dat file (2 48 ), unsigned 2-byte integer
4-7 reserved, all 0's
8-11 columns, unsigned 4-byte integer
12-15 row size in bytes, unsigned 4-byte integer
16-19 header size in bytes, unsigned 4-byte integer
20-23 reserved, all 0's
24-27 reserved, all 0's
28-29 control flags, unsigned 2-byte integer
30-127 reserved, all 0's
Aside from the differences in the descriptor file and the number of columns allowed in the data file, extended data sets conform to the v89 data set description specified above.

### 21.5.7 Matrix v92 (Obsolete)

## Offset Description

| $0-3$ | always 0 |
| :--- | :--- |
| $4-7$ | always 0xEECDCDCD |
| $8-11$ | reserved |
| $12-15$ | reserved |
| $16-19$ | reserved |
| $20-23$ | $0-$ real matrix, 1 - complex matrix |
| $24-27$ | number of dimensions <br>  <br>  <br>  <br>  <br>  <br> $1-$ - scalar vector <br> 2 - column vector, matrix |
| $28-31$ | header size, $128+$ number of dimensions $* 4$, padded to 8 -byte boundary <br> $32-127$ |
|  | reserved |

If the data is a scalar, the data will directly follow the header.
If the data is a row vector, an unsigned integer equaling the number of columns in the vector will precede the data, along with 4 padding bytes.

If the data is a column vector or a matrix, there will be two unsigned integers preceding the data. The first will represent the number of rows in the matrix and the second will represent the number of columns.

The data area always begins on an even 8-byte boundary. Numbers are stored in double precision ( 8 bytes per element, 16 if complex). For complex matrices, all of the real parts are stored first, followed by all the imaginary parts.

### 21.5.8 String v92 (Obsolete)

## Offset Description

0-3 always 0
4-7 always 0xEECFCFCF

## Offset Description

8-11 reserved
12-15 reserved
16-19 reserved
20-23 size of string in units of 8 bytes
24-27 length of string plus null terminator in bytes
28-127 reserved

The size of the data area is always divisible by 8 , and is padded with nulls if the length of the string is not evenly divisible by 8 . If the length of the string is evenly divisible by 8 , the data area will be the length of the string plus 8 . The data area follows immediately after the 128-byte header.

### 21.5.9 Data Set v92 (Obsolete)

## Offset Description

0-3 always 0
4-7 always 0xEECACACA
8-11 reserved
12-15 reserved
16-19 reserved
20-23 rows in data set
24-27 columns in data set
28-31 0 - real data set, 1 - complex data set
32-35 type of data in data set, 2,4 , or 8
36-39 header size in bytes is $128+$ columns * 9
40-127 reserved

The variable names begin at offset 128 and are stored 8 bytes each in ASCII format. Each name corresponds to one column of data. Names less than 8 characters are padded on the right with bytes of zero.

The variable type flags immediately follow the variable names. They are 1-byte binary integers, one per column, padded to an even 8 -byte boundary. A 1 indicates a numeric variable and a 0 indicates a character variable.

The contents of the data set follow the header and start on an 8-byte boundary. Data is either 2-byte signed integer, 4-byte single precision floating point or 8-byte double precision floating point.

### 21.5.10 Matrix v96

## Offset Description

0-3 always 0xFFFFFFFF
4-7 always 0
8-11 always 0xFFFFFFFF
12-15 always 0
16-19 always 0xFFFFFFFF
20-23 0xFFFFFFFF for forward byte order, 0 for backward byte order
24-27 0xFFFFFFFF for forward bit order, 0 for backward bit order
28-31 always 0xABCDEF01
32-35 currently 1
36-39 reserved
40-43 floating point type, 1 for IEEE 754
44-47 1008 (double precision data)
48-51 8, the size in bytes of a double matrix
52-55 0 - real matrix, 1 - complex matrix
56-59 $\quad 1$ - imaginary part of matrix follows real part (standard GAUSS style)
2 - imaginary part of each element immediately follows real part
(FORTRAN style)
60-63 number of dimensions
0 - scalar
1-row vector
2 - column vector or matrix
64-67 1-row major ordering of elements, 2 - column major
68-71 always 0
72-75 header size, 128 + dimensions * 4, padded to 8-byte boundary
76-127 reserved

If the data is a scalar, the data will directly follow the header.
If the data is a row vector, an unsigned integer equaling the number of columns in the vector will
precede the data, along with 4 padding bytes.

If the data is a column vector or a matrix, there will be two unsigned integers preceding the data. The first will represent the number of rows in the matrix and the second will represent the number of columns.

The data area always begins on an even 8-byte boundary. Numbers are stored in double precision ( 8 bytes per element, 16 if complex). For complex matrices, all of the real parts are stored first, followed by all the imaginary parts.

### 21.5.11 Data Set v96

Offset Description
0-3 always 0xFFFFFFFF
4-7 always 0
8-11 always 0xFFFFFFFF
12-15 always 0
16-19 always 0xFFFFFFFF
20-23 0xFFFFFFFF for forward byte order, 0 for backward byte order
24-27 0xFFFFFFFF for forward bit order, 0 for backward bit order
28-31 0xABCDEF02
32-35 version, currently 1
36-39 reserved
40-43 floating point type, 1 for IEEE 754
44-47 $\quad 12$ - signed 2-byte integer
1004 - single precision floating point
1008 - double precision float
48-51 2,4 , or 8 , the size of an element in bytes
52-55 0 - real matrix, 1 - complex matrix
56-59 $\quad 1$ - imaginary part of matrix follows real part (standard GAUSS style)
2 - imaginary part of each element immediately follows real part (FORTRAN style)
60-63 always 2
64-67 1 for row major ordering of elements, 2 for column major

## Offset Description

68-71 always 0
72-75 header size, $128+$ columns * 33, padded to 8 -byte boundary
76-79 reserved
80-83 rows in data set
84-87 columns in data set
88-127 reserved

The variable names begin at offset 128 and are stored 32 bytes each in ASCII format. Each name corresponds to one column of data. Names less than 32 characters are padded on the right with bytes of zero.

The variable type flags immediately follow the variable names. They are 1-byte binary integers, one per column, padded to an even 8 -byte boundary. A 1 indicates a numeric variable and a 0 indicates a character variable.

Contents of the data set follow the header and start on an 8-byte boundary. Data is either 2-byte signed integer, 4-byte single precision floating point or 8-byte double precision floating point.

### 21.5.12 GAUSS Data Archive

A GAUSS Data Archive consists of a header, followed by the variable data and, finally, an array of variable descriptors containing information about each variable.

## Header

The header for a GAUSS Data Archive is laid out as follows:
Offset Type Description

0-3 32-bit unsigned integer always 0xFFFFFFFF
4-7 32-bit unsigned integer always 0
8-11 32-bit unsigned integer always 0xFFFFFFFF

| Offset | Type | Description |
| :--- | :--- | :--- |
| 12-15 | 32-bit unsigned integer | always 0 |
| $16-19$ | 32-bit unsigned integer | always 0xFFFFFFFF |
| $20-23$ | 32-bit unsigned integer | 0xFFFFFFFF for forward byte order, |
|  |  | 0 for backward byte order |
| $24-27$ | 32-bit unsigned integer | always 0 |
| $28-31$ | 32-bit unsigned integer | always 0xABCDEF08 |
| $32-35$ | 32-bit unsigned integer | version, currently 1 |
| $36-39$ | 32-bit unsigned integer | reserved |
| $40-43$ | 32-bit unsigned integer | floating point type, 1 for IEEE 754 |
| $44-55$ | 32-bit unsigned integers | reserved |
| $56-63$ | 64-bit unsigned integer | number of variables |
| $64-67$ | 32-bit unsigned integer | header size, 128 |
| $68-95$ | 32-bit unsigned integers | reserved |
| $96-103$ | 64-bit unsigned integer | offset of variable descriptor table from end of |
|  |  | header |
| $104-127$ | 64-bit unsigned integers | reserved |

## Variable Data

After the header comes the variable data. Matrices are laid out in row-major order, and strings are written with a null-terminating byte.

For string arrays, an array of rows $\times$ columns struct satable's is written out first, followed by the string array data in row-major order with each element null terminated. A struct satable consists of two members:

| Member | Type | Description |
| :--- | :--- | :--- |
|  |  |  |
| off | size_t | offset of element data from beginning of string array data |
| len | size_t | length of element data, including null-terminating byte |

On a 32-bit machine, a size_t is 4 bytes. On a 64-bit machine, it is 8 bytes.
Arrays are written with the orders (sizes) of each dimension followed by the array data. For example, the following $2 \times 3 \times 4$ array:

```
[1,1,1] through [1,3,4] =
    1
5
9
[2,1,1] through [2,3,4]=
```



```
    17}1818\quad19 20 
    21
```

would be written out like this:

123456789101112131415161718192021222324

## Variable Structures

The variable data is followed by an array of variable descriptors. For each variable in the GDA, there is a corresponding variable descriptor in this array. A variable descriptor is laid out as follows:

## Offset Type <br> Description

0-3 32-bit unsigned integer variable type
4-7 32-bit unsigned integer data type, 10 for 8 byte floating point
8-11 32-bit unsigned integer dimensions, used only for arrays
12-15 32-bit unsigned integer complex flag, 1 for real data, 0 for complex
16-19 32-bit unsigned integer size of pointer, indicates whether the variable was written on a 32 -bit or 64 -bit platform

| Offset | Type | Description |
| :--- | :--- | :--- |
| 20-23 | 32-bit unsigned integer | huge flag, indicates whether the variable is larger than <br> INT_MAX |
| $24-31$ | 64-bit unsigned integer  <br> 60ws for matrices and string arrays  <br> 32-39 64-bit unsigned integer <br> columns for matrices and string arrays, length for strings,  <br> including null-terminating byte  |  |
| $40-47$ | 64-bit unsigned integer | index of the variable in the GDA |
| $48-55$ | 64-bit unsigned integer | offset of variable data from end of header <br> $56-63$ |
| 64-bit unsigned integer | length of variable data in bytes <br> name of variable, null-terminated |  |

The variable type (bytes 0-3) may be any of the following:

| $\mathbf{2 0}$ | array |
| :--- | :--- |
| $\mathbf{3 0}$ | matrix |
| $\mathbf{4 0}$ | string |
| $\mathbf{5 0}$ | string array |

The size of pointer element (bytes 16-19) is the size of a pointer on the machine on which the variable was written to the GDA. It will be set to 4 on 32 -bit machines and 8 on 64 -bit machines. This element is used only for string array variables. If a GDA containing string arrays is created on a 32-bit machine and then read on a 64-bit machine, or vice versa, then the size of pointer element indicates how the members of the struct satable's must be converted in order to be read on the current machine.

The huge flag (bytes 20-23) is set to 1 if the variable size is greater than INT_MAX, which is defined as 2147483647 . A variable for which the huge flag is set to 1 may not be read into GAUSS on a 32 -bit machine.

The variable index element (bytes 40-47) contains the index of the variable in the GDA. Although the variable data is not necessarily ordered by index (see gdaUpdate), the variable descriptors are. Therefore, the indices are always in ascending order.


## Foreign Language Interface



The Foreign Language Interface (FLI) allows users to create functions written in C, FORTRAN, or other languages, and call them from a GAUSS program. The functions are placed in dynamic libraries (DLLs, also known as shared libraries or shared objects) and linked in at run-time as needed. The FLI functions are:
dlibrary Link and unlink dynamic libraries at run-time.
dllcall Call functions located in dynamic libraries.

GAUSS recognizes a default dynamic library directory, a directory where it will look for your dynamic-link libraries when you call dlibrary. You can specify the default directory in gauss.cfg by setting dlib_path. As it is shipped, gauss.cfg specifies \$(GAUSSDIR)/dlib as the default directory.

### 22.1 Writing FLI Functions

Your FLI functions should be written to the following specifications:

1. Take 0 or more pointers to doubles as arguments.

This does not mean you cannot pass strings to an FLI function. Just recast the double pointer to a char pointer inside the function.
2. Take those arguments either in a list or a vector.
3. Return an integer.

In C syntax, then, your functions would take one of the following forms:

1. int func(void);
2. int func (double *arg1 $\mathbb{I}$, double *arg2,...】);
3. int func(double *arg[]);

Functions can be written to take a list of up to 100 arguments, or a vector (in C terms, a 1-dimensional array) of up to 1000 arguments. This does not affect how the function is called from GAUSS; the dllcall statement will always appear to pass the arguments in a list. That is, the dllcall statement will always look as follows:
dllcall func(a,b,c,d\|,e...\|);

For details on calling your function, passing arguments to it, getting data back, and what the return value means, see dllcall in the GAUSS Language Reference.

### 22.2 Creating Dynamic Libraries

The following describes how to build a dynamic library called hyp. dll (on Windows) or libhyp. so (on UNIX/Linux) from the source file hyp.c.

As mentioned in the previous section, your FLI functions may take only pointers to doubles as arguments. Therefore, you should define your FLI functions to be merely wrapper functions that cast their arguments as necessary and then call the functions that actually do the work. This is demonstrated in the source file hyp.c:

```
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
/* This code is not meant to be efficient. It is meant
** to demonstrate the use of the FLI.
*/
/* this does all the work, not exported */
static int hypo(double *x, double *y, double *h, int r, int c)
{
    double *wx;
    double *wy;
    double *dp;
    double *sp1;
    double *sp2;
    int i, elems;
    elems = r*c;
    /* malloc work arrays */
    if ((wx = (double *)malloc(elems*sizeof(double))) =\,= NULL)
        return 30; /* out of memory */
    if ((wy = (double *)malloc(elems*sizeof(double))) =\,= NULL)
    {
```

```
    free(wx);
    return 30; /* out of memory */
}
dp = wx;
sp1 = x;
/* square x into work area wx */
for (i=0; i<elems; i++)
{
    *dp = *sp1 * *sp1;
    ++sp1;
    ++dp;
}
dp = wy;
sp2 = y;
/* square y into work area wy */
for (i=0; i<elems; i++)
{
    *dp = *sp2 * *sp2;
    ++sp2;
    ++dp;
}
dp = h;
sp1 = wx;
sp2 = wy;
/* compute hypotenuse into h which was allocated by GAUSS */
for (i=0; i<elems; i++)
{
    *dp = sqrt(*sp1 + *sp2);
    ++sp1;
    ++sp2;
    ++dp;
```

```
    }
    /* free whatever you malloc */
    free(wx);
    free(wy);
    return 0;
}
/* exported wrapper, all double * arguments, calls the real
** function with whatever data types it expects
*/
int hypotenuse(double *x, double *y, double *h,
            double *r, double *c)
{
    return hypo( x, y, h, (int)*r, (int)*c );
}
```

The following Makefiles contain the compile and link commands you would use to build the dynamic library on various platforms. For explanations of the various flags used, see the documentation for your compiler and linker.

## Windows

```
hyp.dll: hyp.obj
link /dll /out:hyp.dll hyp.obj
hyp.obj: hyp.c
cl -c -MD -GX hyp.c
```


## Solaris

\$(CCOPTS) indicates any optional compilation flags you might add.

```
CCOPTIONS = -g -xsb -xarch=v9 -KPIC
    \(C C=c c\)
```

libhyp.so: hyp.c
\$(CC) -G \$(CCOPTIONS) -o \$@ hyp.c -lm

## Linux

\$(CCOPTS) indicates any optional compilation flags you might add.

$$
\begin{aligned}
& \text { CCOPTIONS }=-\mathrm{g}-02-\mathrm{lm} \text {-lc -shared } \\
& \text { CC }=\text { gcc } \\
& \text { libhyp. so: hyp.cpp } \\
& \$(C C) \$(C C O P T I O N S) \text {-o } \$ @ \text { hyp.c }
\end{aligned}
$$

For details on linking your dynamic library, see dlibrary in the GAUSS Language Reference.

## Data Transformations

## 23

GAUSS allows expressions that directly reference variables (columns) of a data set. This is done within the context of a data loop:

```
dataloop infile outfile;
    drop wagefac wqlec shordelt foobly;
    csed = ln(sqrt(csed));
    select csed > 0.35 and married $=\,= "y";
    make chfac = hcfac + wcfac;
    keep csed chfac stid recsum voom;
endata;
```

GAUSS translates the data loop into a procedure that performs the required operations, and then calls the procedure automatically at the location (in your program) of the data loop. It does this by translating your main program file into a temporary file and then executing the temporary file.

A data loop may be placed only in the main program file. Data loops in files that are \#include'd or autoloaded are not recognized.

### 23.1 Data Loop Statements

A data loop begins with a dataloop statement and ends with an endata statement. Inside a data loop, the following statements are supported:

| code | Create variable based on a set of logical expressions. |
| :--- | :--- |
| delete | Delete rows (observations) based on a logical expression. |
| drop | Specify variables NOT to be written to data set. |
| extern | Allow access to matrices and strings in memory. |
| keep | Specify variables to be written to output data set. |
| lag | Lag variables a number of periods. |
| listwise | Control deletion of missing values. |
| make | Create new variable. |
| outtyp | Specify output file precision. |
| recode | Change variable based on a set of logical expressions. |
| select | Select rows (observations) based on a logical expression. |
| vector | Create new variable from a scalar returning expression. |

In any expression inside a data loop, all text symbols not immediately followed by a left parenthesis ' (' are assumed to be data set variable (column) names. Text symbols followed by a left parenthesis are assumed to be procedure names. Any symbol listed in an extern statement is assumed to be a matrix or string already in memory.

### 23.2 Using Other Statements

All program statements in the main file and not inside a data loop are passed through to the temporary file without modification. Program statements within a data loop that are preceded by a '\#' are passed through to the temporary file without modification. The user familiar with the code generated in the temporary file can use this to do out-of-the-ordinary operations inside the data loop.

### 23.3 Debugging Data Loops

The translator that processes data loops can be turned on and off. When the translator is on, there are three distinct phases in running a program:

Translation Translation of main program file to temporary file.
Compilation Compilation of temporary file.
Execution Execution of compiled code.

### 23.3.1 Translation Phase

In the translation phase, the main program file is translated into a temporary file. Each data loop is translated into a procedure and a call to this procedure is placed in the temporary file at the same location as the original data loop. The data loop itself is commented out in the temporary file. All the data loop procedures are placed at the end of the temporary file.

Depending upon the status of line number tracking, error messages encountered in this phase will be printed with the file name and line numbers corresponding to the main file.

### 23.3.2 Compilation Phase

In the compilation phase, the temporary file is compiled. Depending upon the status of line number tracking, error messages encountered in this phase will be printed with the file name and
line numbers corresponding to both the main file and the temporary file.

### 23.3.3 Execution Phase

In the execution phase, the compiled program is executed. Depending on the status of line number tracking, error messages will include line number references from both the main file and the temporary file.

### 23.4 Reserved Variables

The following local variables are created by the translator and used in the produced code:

| x_cv | x_iptr | x_ncol | x_plag |
| :--- | :--- | :--- | :--- |
| x_drop | x_keep | x_nlag | x_ptrim |
| x_fpin | x_lval | x_nrow | x_shft |
| x_fpout | x_lvar | x_ntrim | x_tname |
| x_i | x_n | x_out | x_vname |
| x_in | x_name | x_outtyp | x_x |

These variables are reserved, and should not be used within a dataloop... endata section.

## The GAUSS Profiler



GAUSS now includes a profiler, which enables you to determine exactly how much time your programs are spending on each line and in each called procedure, thereby providing you with the information you need to increase the efficiency of your programs. The GAUSS Profiler and tcollect are both run from a command prompt window, not at a GAUSS prompt.

### 24.1 Using the GAUSS Profiler

There are two steps to using the GAUSS Profiler: collection and analysis.

### 24.1.1 Collection

To collect profiling information, you must run your GAUSS program in tcollect, an executable shipped with GAUSS that is identical to tgauss except that it generates a file containing profiling information each time it is run:

```
tcollect -b myfile.e
```

The output displayed by tcollect includes the name of the output file containing the profiling information. tcollect output files have a gaussprof prefix and a . gco extension.

Note that running tcollect on long programs may generate a very large .gco output file. Thus you may want to delete the . gco files on your machine regularly.

### 24.1.2 Analysis

To analyze the information stored in the tcollect output file, you must run the gaussprof executable, which is also shipped with GAUSS, on that file. gaussprof produces an organized report, displaying the time usage by procedure and by line.

Assuming that running myfile.e in tcollect produced an output file called gaussprof_001.gco, you could analyze the results in that file as follows:
gaussprof gaussprof_001.gco

The syntax for gaussprof is:
gaussprof [flags] profile_data_file ...
where [flags] may be any of the following:
-p profile procedure calls
-1 profile line numbers
-h suppress headers
-sp order procedure call sort order where order contains one or more of the folllowing:

| $\mathbf{e}$ | exclusive time |
| :--- | :--- |
| $\mathbf{t}$ | total time |
| $\mathbf{c}$ | number of times called |
| $\mathbf{p}$ | procedure name |
| $\mathbf{a}$ | ascending order |
| $\mathbf{d}$ | descending order (default) |

Columns are sorted all ascending or all descending.
-sl order line number sort order where order contains one or more of the following:
$t \quad$ time spent on line
c number of times line was executed
f file name
1 line number
a ascending order
d descending order (default)
Columns are sorted all ascending or all descending.

The default, with no flags, is: -pl -sp dep -sl dtf.


## Publication Quality Graphics

## 25

GAUSS Publication Quality Graphics (PQG) is a set of routines built on the graphics functions in GraphiC by Scientific Endeavors Corporation.

The main graphics routines include $x y$, $x y z$, surface, polar and $\log$ plots, as well as histograms, bar, and box graphs. Users can enhance their graphs by adding legends, changing fonts, and adding extra lines, arrows, symbols and messages.

The user can create a single full size graph, inset a smaller graph into a larger one, tile a window with several equally sized graphs or place several overlapping graphs in the window. Graphic panel size and location are all completely under the user's control.

### 25.1 General Design

GAUSS PQG consists of a set of main graphing procedures and several additional procedures and global variables for customizing the output.

All of the actual output to the window happens during the call to these main routines:

| bar | Bar graphs. |
| :--- | :--- |
| box | Box plots. |
| contour | Contour plots. |
| draw | Draw graphs using only global variables. |
| hist | Histogram. |
| histp | Percentage histogram. |
| histf | Histogram from a vector of frequencies. |
| loglog | Log scaling on both axes. |
| logx | Log scaling on X axis. |
| logy | Log scaling on Y axis. |
| polar | Polar plots. |
| surface | 3-D surface with hidden line removal. |
| xy | Cartesian graph. |
| xyz | 3-D Cartesian graph. |

### 25.2 Using Publication Quality Graphics

### 25.2.1 Getting Started

There are four basic parts to a graphics program. These elements should be in any program that uses graphics routines. The four parts are the header, data setup, graphics format setup, and graphics call.

## Header

In order to use the graphics procedures, the pgraph library must be activated. This is done in the library statement at the top of your program or command file. The next line in your program will typically be a command to reset the graphics global variables to their default state. For example:

```
library mylib, pgraph;
graphset;
```


## Data Setup

The data to be graphed must be in matrices. For example:

$$
\begin{aligned}
& x=\operatorname{seq}(1,1,50) ; \\
& y=\sin (x)
\end{aligned}
$$

## Graphics Format Setup

Most of the graphics elements contain defaults that allow the user to generate a plot without modification. These defaults, however, may be overridden by the user through the use of global variables and graphics procedures. Some of the elements that may be configured by the user are axes numbering, labeling, cropping, scaling, line and symbol sizes and types, legends, and colors.

## Calling Graphics Routines

The graphics routines take as input the user data and global variables that have previously been set. It is in these routines where the graphics file is created and displayed.

Following are three PQG examples. The first two programs are different versions of the same graph. The variables that begin with _p are the global control variables used by the graphics
routines. (For a detailed description of these variables, see Global Control Variables, Section 25.6.

Example 1 The routine being called here is a simple XY plot. The entire window will be used. Four sets of data will be plotted with the line and symbol attributes automatically selected. This graph will include a legend, title, and a time/date stamp (time stamp is on by default):

```
library pgraph;
graphset; /* reset global variables */
x = seqa(.1,.1,100); /* generate data */
y = sin(x);
y = y ~ y*.8 ~ y*.6 ~ y*.4; /* 4 curves plotted against x */
_plegctl = 1; /* legend on */
title("Example xy Graph"); /* Main title */
xy(x,y);
/* activate PGRAPH library */
/* reset global variables */
/* generate data */
/* Call to main routine */
```

Example 2 Here is the same graph with more of the graphics format controlled by the user. The first two data sets will be plotted using symbols at data points only (observed data); the data points in the second two sets will be connected with lines (predicted results):

```
library pgraph;
graphset;
x = seqa(.1,.1,100);
y = sin(x);
y = y ~ y*.8 ~ y*.6 ~ y*.4;
_pdate = "";
_plctrl = { 1, 1, 0, @ };
_pltype = { 1, 2, 6, 6 };
_pstype = { 1, 2, 0, @ };
_plegctl= { 2, 3, 1.7, 4.5 };
_plegstr= "Sin wave 1.\0"\
    "Sin wave . 8\0"\
    "Sin wave .6\0"\
    "Sin wave .4";
ylabel("Amplitude"); /* Y axis label */
```

```
xlabel("X Axis"); /* X axis label */
title("Example xy Graph"); /* main title */
xy(x,y); /* call to main routine */
```

Example 3 In this example, two graphics panels are drawn. The first is a full-sized surface representation, and the second is a half-sized inset containing a contour of the same data located in the lower left corner of the window:
library pgraph; /* activate pgraph library */

```
/* Generate data for surface and contour plots */
x = seqa(-10,0.1,71)'; /* note x is a row vector */
y = seqa(-10,0.1,71); /* note y is a column vector */
z = cos(5*sin(x) - y); /* z is a 71x71 matrix */
```

begwind; /* initialize graphics windows */
makewind $(9,6.855,0,0,0)$; /* first window full size */
makewind(9/2,6.855/2,1,1,0); /* second window inset to first */

```
setwind(1); /* activate first window */
    graphset; /* reset global variables */
    _pzclr = { 1, 2, 3, 4 }; /* set Z level colors */
    title("cos(5*sin(x) - y)"); /* set main title */
    xlabel("X Axis"); /* set X axis label */
    ylabel("Y Axis"); /* set Y axis label */
    scale3d(miss(0,0),miss(0,0),-5|5); /* scale Z axis */
    surface(x,y,z); /* call surface routine */
```

nextwind;
graphset;
_pzclr = \{ 1, 2, 3, 4 \};
_pbox = 15;
contour ( $\mathrm{x}, \mathrm{y}, \mathrm{z}$ );
endwind;
/* activate second window. */
/* reset global variables */
/* set Z level colors */
/* white border */
/* call contour routine */
/* Display windows */

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While the structure has changed somewhat, the four basic elements of the graphics program are all here. The additional routines begwind, endwind, makewind, nextwind, and setwind are all used to control the graphic panels.

As Example 3 illustrates, the code between graphic panel functions (that is, setwind or nextwind) may include assignments to global variables, a call to graphset, or may set up new data to be passed to the main graphics routines.

You are encouraged to run the example programs supplied with GAUSS. Analyzing these programs is perhaps the best way to learn how to use the PQG system. The example programs are located on the examples subdirectory.

### 25.2.2 Graphics Coordinate System

PQG uses a $4190 \times 3120$ pixel resolution grid on a $9.0 \times 6.855$-inch printable area. There are three units of measure supported with most of the graphics global elements:

## Inch Coordinates

Inch coordinates are based on the dimensions of the full-size $9.0 \times 6.855$-inch output page. The origin is $(0,0)$ at the lower left corner of the page. If the picture is rotated, the origin is at the upper left. (For more information, see Inch Units in Graphic Panels, Section 25.3.5.)

## Plot Coordinates

Plot coordinates refer to the coordinate system of the graph in the units of the user's $\mathrm{X}, \mathrm{Y}$ and Z axes.

## Pixel Coordinates

Pixel coordinates refer to the $4096 \times 3120$ pixel coordinates of the full-size output page. The origin is $(0,0)$ at the lower left corner of the page. If the picture is rotated, the origin is at the upper left.

### 25.3 Graphic Panels

Multiple graphic panels for graphics are supported. These graphic panels allow the user to display multiple graphs on one window or page.

A graphic panel is any rectangular subsection of the window or page. Graphc panels may be any size and position on the window and may be tiled or overlapping, transparent or nontransparent.

### 25.3.1 Tiled Graphic Panels

Tiled graphic panels do not overlap. The window can easily be divided into any number of tiled graphic panels with the window command. window takes three parameters: number of rows, number of columns, and graphic panel attribute ( $1=$ transparent, $0=$ nontransparent).

This example will divide the window into six equally sized graphic panels. There will be two rows of three graphic panels-three graphic panels in the upper half of the window and three in the lower half. The attribute value of 0 is arbitrary since there are no other graphic panels beneath them.

```
window(nrows,ncols,attr);
window (2,3,0);
```


### 25.3.2 Overlapping Graphic Panels

Overlapping graphic panels are laid on top of one another as they are created, much as if you were using the cut and paste method to place several graphs together on one page. An overlapping graphic panel is created with the makewind command.

In this example, makewind will create an overlapping graphic panel that is 4 inches wide by 2.5 inches tall, positioned 1 inch from the left edge of the page and 1.5 inches from the bottom of the page. It will be nontransparent:

```
makewind(hsize,vsize,hpos,vpos,attr);
```

```
window(2,3,0);
makewind(4,2.5,1,1.5,0);
```


### 25.3.3 Nontransparent Graphic Panels

A nontransparent graphic panel is one that is blanked before graphics information is written to it. Therefore, information in any previously drawn graphic panels that lie under it will not be visible.

### 25.3.4 Transparent Graphic Panels

A transparent graphic panel is one that is not blanked, allowing the graphic panel beneath it to "show through". Lines, symbols, arrows, error bars, and other graphics objects may extend from one graphic panel to the next by using transparent graphic panels. First, create the desired graphic panel configuration. Then create a full-window, transparent graphic panel using the makewind or window command. Set the appropriate global variables to position the desired object on the transparent graphic panel. Use the draw procedure to draw it. This graphic panel will act as a transparent "overlay" on top of the other graphic panels. Transparent graphic panels can be used to add text or to superimpose one graphic panel on top of another.

### 25.3.5 Using Graphic Panel Functions

The following is a summary of the graphic panel functions:

| begwind | Graphic panel initialization procedure. |
| :--- | :--- |
| endwind | End graphic panel manipulations, display graphs. |
| window | Partition window into tiled graphic panels. |
| makewind | Create graphic panel with specified size and position. |
| setwind | Set to specified graphic panel number. |

nextwind $\quad$ Set to next available graphic panel number.
getwind Get current graphic panel number.
savewind $\quad$ Save graphic panel configuration to a file.
loadwind Load graphic panel configuration from a file.

This example creates four tiled graphic panels and one graphic panel that overlaps the other four:

```
library pgraph;
graphset;
begwind;
window(2,2,0); /* Create four tiled graphic panels
    (2 rows, 2 columns) */
xsize = 9/2; /* Create graphic panel that overlaps the
    tiled graphic panels */
ysize = 6.855/2;
makewind(xsize,ysize,xsize/2,ysize/2,0);
x = seqa(1,1,1000); /* Create X data */
y = (sin}(\textrm{x})+1) * 10.; /* Create Y data */
setwind(1); /* Graph #1, upper left corner */
    xy(x,y);
nextwind; /* Graph #2, upper right corner */
    logx(x,y);
nextwind; /* Graph #3, lower left corner */
    logy(x,y);
nextwind; /* Graph #4, lower right corner */
    loglog(x,y);
nextwind; /* Graph #5, center, overlayed */
    bar(x,y);
endwind; /* End graphic panel processing,
                                display graph */
```


### 25.3.6 Inch Units in Graphic Panels

Some global variables allow coordinates to be input in inches. If a coordinate value is in inches and is being used in a graphic panel, that value will be scaled to "graphic panel inches" and positioned relative to the lower left corner of the graphic panel. A "graphic panel inch" is a true inch in size only if the graphic panel is scaled to the full window, otherwise X coordinates will be scaled relative to the horizontal graphic panel size and Y coordinates will be scaled relative to the vertical graphic panel size.

### 25.3.7 Saving Graphic Panel Configurations

The functions savewind and loadwind allow the user to save graphic panel configurations. Once graphic panels are created (using makewind and window), savewind may be called. This will save to disk the global variables containing information about the current graphic panel configuration. To load this configuration again, call loadwind. (See loadwind in the GAUSS Language Reference.

### 25.4 Graphics Text Elements

Graphics text elements, such as titles, messages, axes labels, axes numbering, and legends, can be modified and enhanced by changing fonts and by adding superscripting, subscripting, and special mathematical symbols.

To make these modifications and enhancements, the user can embed "escape codes" in the text strings that are passed to title, xlabel, ylabel and asclabel or assigned to _pmsgstr and _plegstr.

The escape codes used for graphics text are:
$\backslash 000$ String termination character (null byte).
[ Enter superscript mode, leave subscript mode.
] Enter subscript mode, leave superscript mode.
@ Interpret next character as literal.
$\backslash 20 n$ Select font number $n$. (see Selecting Fonts, following).

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The escape code $\backslash \mathbf{L}$ (or $\backslash \mathbf{l}$ ) can be embedded into title strings to create a multiple line title:

```
title("This is the first line\lthis is the second line");
```

A null byte $\backslash 000$ is used to separate strings in _plegstr and _pmsgstr:

```
_pmsgstr = "First string\000Second string\000Third string";
```

or

```
_plegstr = "Curve 1\000Curve 2";
```

Use [..] to create the expression $M(t)=E\left(e^{t x}\right)$ :
_pmsgstr = "M(t) = E(e[tx])";

Use @ to generate [ and ] in an X axis label:

```
xlabel("Data used for x is: data@[.,1 2 3@]");
```


### 25.4.1 Selecting Fonts

Four fonts are supplied with the Publication Quality Graphics system. They are Simplex, Complex, Simgrma, and Microb. (For a list of the characters available in each font, see Appendix A.)

Fonts are loaded by passing to the fonts procedure a string containing the names of all fonts to be loaded. For example, this statement will load all four fonts:

```
fonts("simplex complex microb simgrma");
```

The fonts command must be called before any of the fonts can be used in text strings. A font can then be selected by embedding an escape code of the form " $\backslash 20 n$ " in the string that is to be written in the new font. The $n$ will be $1,2,3$ or 4 , depending on the order in which the fonts were loaded in fonts. If the fonts were loaded as in the previous example, the escape characters for each would be:
$\backslash 201$ Simplex
$\backslash 202$ Complex
$\backslash 203$ Microb
$\backslash 204$ Simgrma
The following example demonstrates how to select a font for use in a string:

```
title("\201This is the title using Simplex font");
xlabel("\202This is the label for X using Complex font");
ylabel("\203This is the label for Y using Microb font");
```

Once a font is selected, all succeeding text will use that font until another font is selected. If no fonts are selected by the user, a default font (Simplex) is loaded and selected automatically for all text work.

### 25.4.2 Greek and Mathematical Symbols

The following examples illustrate the use of the Simgrma font; they assume that Simgrma was the fourth font loaded. (For the available Simgrma characters and their numbers, see Appendix A.) The Simgrma characters are specified by either:

1. The character number, preceeded by a " $\backslash$ ".
2. The regular text character with the same number.

For example, to get an integral sign " $\int$ " in Simgrma, embed either a " $\backslash \mathbf{0 4 4}$ " or a "," in a string that has been set to use the Simgrma font.

To produce the title $f(x)=\sin ^{2}(\pi x)$, the following title string should be used:

$$
\text { title("\201f(x) }=\sin [2](\backslash 204 p \backslash 201 x) ") ;
$$

The " $p$ " (character 112) corresponds to " $\pi$ " in Simgrma.
To number the major X axis tick marks with multiples of $\pi / 4$, the following could be passed to asclabel:

```
lab = "\2010 \204p\201/4 \204p\201/2 3\204p\201/4 \204p";
asclabel(lab,0);
xtics(0,pi,pi/4,1);
```

xtics is used to make sure that major tick marks are placed in the appropriate places.
This example will number the X axis tick marks with the labels $\mu^{-2}, \mu^{-1}, 1, \mu$, and $\mu^{2}$ :

```
lab = "\204m\201[-2] \204m\201[-1] 1 \204m m\201[2]";
asclabel(lab,0);
```

This example illustrates the use of several of the special Simgrma symbols:

$$
\text { _pmsgstr }=\text { "\2041\2011/2\204p ,\201e[-\204m[\2012]\201/2]d\204m"; }
$$

This produces:

$$
\sqrt{ } 1 / 2 \pi \int e^{-\mu^{2} / 2} d \mu
$$

### 25.5 Colors

| 0 | Black | 8 | Dark Grey |
| :--- | :--- | ---: | :--- |
| 1 | Blue | 9 | Light Blue |
| 2 | Green | 10 | Light Green |
| 3 | Cyan | 11 | Light Cyan |
| 4 | Red | 12 | Light Red |
| 5 | Magenta | 13 | Light Magenta |
| 6 | Brown | 14 | Yellow |
| 7 | Grey | 15 | White |

### 25.6 Global Control Variables

The following global variables are used to control various graphics elements. Default values are provided. Any or all of these variables can be set before calling one of the main graphing routines. The default values can be modified by changing the declarations in pgraph.dec and the statements in the procedure graphset in pgraph.src. graphset can be called whenever the user wants to reset these variables to their default values.
_pageshf $2 \times 1$ vector, the graph will be shifted to the right and up if this is not 0 . If this is 0 , the graph will be centered on the output page. Default is 0 .
Note: Used internally. (For the same functionality, see makewind in the GAUSS Language Reference.) This is used by the graphic panel routines. The user must not set this when using the graphic panel procedures.
_pagesiz $2 \times 1$ vector, size of the graph in inches on the printer output. Maximum size is $9.0 \times 6.855$ inches (unrotated) or $6.855 \times 9.0$ inches (rotated). If this is 0 , the maximum size will be used. Default is 0 .

Note: Used internally. (For the same functionality, see makewind in the GAUSS Language Reference). This is used by the graphic panel routines. The user must not set this when using the graphic panel procedures.
_parrow $\quad \mathrm{M} \times 11$ matrix, draws one arrow per row of the input matrix (for total of M arrows). If scalar zero, no arrows will be drawn.
[M,1] x starting point.
[M,2] y starting point.
[M,3] x ending point.
[M,4] y ending point.
[M,5] ratio of the length of the arrow head to half its width.
[M,6] size of arrow head in inches.
[M,7] type and location of arrow heads. This integer number will be interpreted as a decimal expansion $m n$, for example: if 10 , then $m=1, n=$ 0 .
$m$, type of arrow head:
0 solid
1 empty
2 open
3 closed
$n$, location of arrow head:
0 none
1 at the final end
2 at both ends
[M,8] color of arrow, see Colors, Section 25.5.
[ $\mathbf{M}, 9]$ coordinate units for location:
$1 \mathrm{x}, \mathrm{y}$ starting and ending locations in plot coordinates
$2 \mathrm{x}, \mathrm{y}$ starting and ending locations in inches
$3 \mathrm{x}, \mathrm{y}$ starting and ending locations in pixels
[M,10] line type:
1 dashed
2 dotted
3 short dashes
4 closely spaced dots
5 dots and dashes
6 solid
[ $\mathbf{M}, \mathbf{1 1}$ ] controls thickness of lines used to draw arrow. This value may be zero or greater. A value of zero is normal line width.

To create two single-headed arrows, located using inches, use

$$
\text { _parrow }=\left\{\begin{array}{lllllllllll} 
& 1 & 1 & 2 & 2 & 3 & 0.2 & 11 & 10 & 2 & 6 \\
0 & 0, \\
3 & 4 & 2 & 2 & 3 & 0.2 & 11 & 10 & 2 & 6 & 0
\end{array}\right\} ;
$$

_parrow3 $\mathrm{M} \times 12$ matrix, draws one 3-D arrow per row of the input matrix (for a total of M arrows). If scalar zero, no arrows will be drawn.
[ $\mathbf{M}, \mathbf{1}] \times$ starting point in 3-D plot coordinates.
[ $\mathbf{M}, 2]$ y starting point in 3-D plot coordinates.
[ $\mathbf{M}, 3] \mathrm{z}$ starting point in 3-D plot coordinates.
[M,4] x ending point in 3-D plot coordinates.
[M,5] y ending point in 3-D plot coordinates.
[M,6] z ending point in 3-D plot coordinates.
$[\mathbf{M}, 7]$ ratio of the length of the arrow head to half its width.
[ $\mathbf{M}, 8]$ size of arrow head in inches.
[ $\mathbf{M}, 9]$ type and location of arrow heads. This integer number will be interpreted as a decimal expansion $m n$. For example: if 10 , then $m=1, n$ $=0$.
$m$, type of arrow head:
0 solid
1 empty
2 open
3 closed
$n$, location of arrow head:
0 none
1 at the final end
2 at both ends
$[\mathbf{M , 1 0 ]}$ color of arrow, see Colors, Section 25.5.
[M,11] line type:
1 dashed
2 dotted
3 short dashes
4 closely spaced dots
5 dots and dashes
6 solid
[ $\mathbf{M}, \mathbf{1 2}$ ] controls thickness of lines used to draw arrow. This value may be zero or greater. A value of zero is normal line width.

To create two single-headed arrows, located using plot coordinates, use

$$
\text { _parrow3 }=\left\{\begin{array}{llllllllllll}
1 & 1 & 1 & 2 & 2 & 2 & 3 & 0.2 & 11 & 10 & 6 & 0, \\
3 & 4 & 5 & 2 & 2 & 2 & 3 & 0.2 & 11 & 10 & 6 & 0
\end{array}\right\} ;
$$

## _paxes

_paxht scalar, size of axes labels in inches. If 0 , a default size will be computed. Default is 0 .
_pbartyp $1 \times 2$ or $\mathrm{K} \times 2$ matrix. Controls bar shading and colors in bar graphs and histograms.

The first column controls the bar shading:

0 no shading
1 dots

2 vertical cross-hatch
3 diagonal lines with positive slope
4 diagonal lines with negative slope
5 diagonal cross-hatch
6 solid
The second column controls the bar color, see Colors, Section 25.5.
_pbarwid scalar, width of bars in bar graphs and histograms. The valid range is $0-1$. If 0 , the bars will be a single pixel wide. If 1 , the bars will touch each other. Default is 0.5 , so the bars take up about half the space open to them.
_pbox scalar, draws a box (border) around the entire graph. Set to desired color of box to be drawn. Use 0 if no box is desired. Default is 0 .
_pboxctl $5 \times 1$ vector, controls box plot style, width, and color. Used by procedure box only.
[1] box width between 0 and 1 . If 0 , the box plot is drawn as two vertical lines representing the quartile ranges with a filled circle representing the $50^{\text {th }}$ percentile.
[2] box color, see Colors, Section 25.5. If 0, the colors may be individually controlled using global variable _pcolor.
[3] $\mathrm{min} / \mathrm{max}$ style for the box symbol. One of the following:
1 minimum and maximum taken from the actual limits of the data. Elements 4 and 5 are ignored.
2 statistical standard with the minimum and maximum calculated according to interquartile range as follows:

$$
\begin{aligned}
\text { intqrange } & =75^{t^{\text {th }}-25^{\text {th }}} \\
\min & =25^{\text {th }}-1.5 \text { intqrange } \\
\max & =75^{\text {th }}+1.5 \text { intqrange }
\end{aligned}
$$

Elements 4 and 5 are ignored.
3 minimum and maximum percentiles taken from elements 4 and 5 .
[4] minimum percentile value ( $0-100$ ) if _pboxctl[3] $=3$.
[5] maximum percentile value ( $0-100$ ) if _pboxctl[3] $=3$.

## _pboxlim $\quad 5 \times \mathrm{M}$ output matrix containing computed percentile results from procedure box.

 M corresponds to each column of input $y$ data.[1,M] minimum whisker limit according to _pboxctl[3].
[2,M] 25th percentile (bottom of box).
[3,M] 50th percentile (median).
[4,M] 75th percentile (top of box).
[5,M] maximum whisker limit according to _pboxctl [3].
_pcolor scalar or $K \times 1$ vector, colors for main curves in $\mathbf{x y}$, $\mathbf{x y z}$ and $\log$ graphs. To use a single color set for all curves set this to a scalar color value. If 0 , use default colors. Default is 0 .
The default colors come from a global vector called _pcsel. This vector can be changed by editing pgraph. dec to change the default colors, see Colors, Section 25.5 (_pcsel is not documented elsewhere).
_pcrop scalar or $1 \times 5$ vector, allows plot cropping for different graphic elements to be individually controlled. Valid values are 0 (disabled) or 1 (enabled). If cropping is enabled, any graphical data sent outside the axes area will not be drawn. If this is a scalar, it is expanded to a $1 \times 5$ vector using the given value for all elements. All cropping is enabled by default.
[1] crop main curves/symbols.
[2] crop lines generated using _pline.
[3] crop arrows generated using _parrow.
[4] crop circles/arcs generated using _pline.
[5] crop symbols generated using _psym.

This example will crop main curves, and lines and circles drawn by _pline.

```
_pcrop = {111010 };
```

_pcross scalar. If 1, the axes will intersect at the ( 0,0 ) X-Y location if it is visible. Default is 0 , meaning the axes will be at the lowest end of the $\mathrm{X}-\mathrm{Y}$ coordinates.
_pdate date string. If this contains characters, the date will be appended and printed. The default is set as follows (the first character is a font selection escape code):
_pdate = "\201GAUSS ";

If this is set to a null string, no date will be printed. (For more information on using fonts within strings, see Graphics Text Elements, Section 25.4.
_perrbar $\quad \mathrm{M} \times 9$ matrix, draws one error bar per row of the input matrix. If scalar 0 , no error bars will be drawn. Location values are in plot coordinates.
[M,1] x location.
[M,2] left end of error bar.
[M,3] right end of error bar.
[M,4] y location.
[M,5] bottom of error bar.
[M,6] top of error bar.
[M,7] line type:
1 dashed
2 dotted
3 short dashes
4 closely spaced dots
5 dots and dashes
6 solid
[M,8] color, see Colors, Section 25.5.
[M,9] line thickness.. This value may be 0 or greater. A value of 0 is normal line width.

To create one error bar using solid lines, use

$$
\text { _perrbar }=\left\{\begin{array}{lllllllllll}
1 & 0 & 2 & 2 & 1 & 3 & 6 & 2 & 0 & \} ;
\end{array}\right.
$$

_pframe $\quad 2 \times 1$ vector, controls frame around axes area. On 3-D plots this is a cube surrounding the 3-D workspace.
[1] 1 frame on
0 frame off
[2] 1 tick marks on frame
0 no tick marks
The default is a frame with tick marks.
_pgrid $2 \times 1$ vector to control grid.
[1] grid through tick marks:
0 no grid
1 dotted grid
2 fine dotted grid
3 solid grid
[2] grid subdivisions between major tick marks:
0 no subdivisions
1 dotted lines at subdivisions
2 tick marks only at subdivisions
The default is no grid and tick marks at subdivisions.
_plctrl scalar or $K \times 1$ vector to control whether lines and/or symbols will be displayed for the main curves. This also controls the frequency of symbols on main curves. The number of rows $(\mathrm{K})$ is equal to the number of individual curves to be plotted in the graph. Default is 0 .

0 draw line only.
>0 draw line and symbols every _plctrl points.
<0 draw symbols only every _plctrl points.
-1 all of the data points will be plotted with no connecting lines.
This example draws a line for the first curve, draws a line and plots a symbol every 10 data points for the second curve, and plots symbols only every 5 data points for the third curve:

$$
\text { _plctrl = \{ 0, 10, -5 \}; }
$$

_plegctl scalar or $1 \times 4$ vector, legend control variable.
If scalar 0 , no legend is drawn (default). If nonzero scalar, create legend in the default location in the lower right of the page.

If $1 \times 4$ vector, set as follows:
[1] legend position coordinate units:
1 coordinates are in plot coordinates
2 coordinates are in inches
3 coordinates are in pixel
[2] legend text font size, where $1<=$ size $<=9$. Default is 5 .
[3] $x$ coordinate of lower left corner of legend box.
[4] y coordinate of lower left corner of legend box.
This example puts a legend in the lower right corner:
_plegctl = 1;

This example creates a smaller legend and positions it 2.5 inches from the left and 1 inch from the bottom.

$$
\text { _plegctl }=\left\{\begin{array}{llll}
2 & 2.5 & 1
\end{array}\right\} ;
$$

_plegstr string, legend entry text. Text for multiple curves is separated by a null byte (" $\backslash 000$ ").
For example:

```
_plegstr = "Curve 1\000Curve 2\000Curve 3";
```

_plev $\quad \mathrm{M} \times 1$ vector, user-defined contour levels for contour. Default is 0 . (See contour in the GAUSS Language Reference.)
_pline
$\mathrm{M} \times 9$ matrix, to draw lines, circles, or radii. Each row controls one item to be drawn. If this is a scalar zero, nothing will be drawn. Default is 0 .
[M,1] item type and coordinate system:

1 line in plot coordinates
2 line in inch coordinates
3 line in pixel coordinates
4 circle in plot coordinates
5 circle in inch coordinates
6 radius in plot coordinates
7 radius in inch coordinates
[M,2] line type:
1 dashed
2 dotted
3 short dashes
4 closely spaced dots
5 dots and dashes
6 solid
[ $\mathbf{M}, \mathbf{3 - 7}$ ] coordinates and dimensions:
if item type is line ( $1<==$ pline $[\mathrm{M}, 1]<=3$ ):
[ $\mathbf{M}, 3] \times$ starting point.
[ $\mathbf{M}, 4]$ y starting point.
[ $\mathbf{M}, 5]$ x ending point.
[ $\mathbf{M}, 6]$ y ending point.
[ $\mathbf{M}, 7] 0$ if this is a continuation of a curve, 1 if this begins a new curve.
if item type is circle (_pline[M, 1] = 4 or _pline $[\mathrm{M}, 1]=5$ ):
[ $\mathbf{M}, 3] \times$ center of circle.
[M,4] y center of circle.
[ $\mathbf{M}, 5$ ] radius.
$[\mathbf{M , 6 ]}$ starting point of arc in radians.
[ $\mathbf{M}, 7]$ ending point of arc in radians.
if item type is radius (_pline[ $\mathrm{M}, 1]=6$ or _pline $[\mathrm{M}, 1]=7$ ):
[ $\mathbf{M}, 3$ ] x center of circle.
[M,4] y center of circle.
[ $\mathbf{M}, 5$ ] beginning point of radius, 0 is the center of the circle.
[M,6] ending point of radius.
[M,7] angle in radians.
[M,8] color, see Colors, Section 25.5.
[M,9] controls line thickness. This value may be zero or greater. A value of zero is normal line width.
_pline3d $\quad \mathrm{M} \times 9$ matrix. Allows extra lines to be added to an xyz or surface graph in 3-D plot coordinates.
[M,1] x starting point.
[M,2] y starting point.
[M,3] z starting point.
[M,4] x ending point.
[M,5] y ending point.
[M,6] z ending point.
[M,7] color.
[M,8] line type:
1 dashed
2 dotted
3 short dashes
4 closely spaced dots
5 dots and dashes
6 solid
[M,9] line thickness, $0=$ normal width.
[M,10] hidden line flag, $1=$ obscured by surface, $0=$ not obscured.
_plotshf $2 \times 1$ vector, distance of plot from lower left corner of output page in inches.
[1] $x$ distance.
[2] y distance.
If scalar 0 , there will be no shift. Default is 0 .
Note: Used internally. (For the same functionality, see axmargin in the
GAUSS Language Reference.) This is used by the graphic panel routines. The user must not set this when using the graphic panel procedures.
_pltype scalar or $K \times 1$ vector, line type for the main curves. If this is a nonzero scalar, all lines will be this type. If scalar 0 , line types will be default styles. Default is 0 .

1 dashed
2 dotted
3 short dashes
4 closely spaced dots
5 dots and dashes
6 solid

The default line types come from a global vector called _plsel. This vector can be changed by editing pgraph.dec to change the default line types (_plsel is not documented elsewhere.)
_plwidth scalar or $\mathrm{K} \times 1$ vector, line thickness for main curves. This value may be zero or greater. A value of zero is normal (single pixel) line width. Default is 0 .
_pmcolor $9 \times 1$ vector, color values to use for plot, see Colors, Section 25.5.
[1] axes.
[2] axes numbers.
[3] X axis label.
[4] Y axis label.
[5] Z axis label.
[6] title.
[7] box.
[8] date.
[9] background.

If this is scalar, it will be expanded to a $9 \times 1$ vector.
_pmsgctl $\mathrm{L} \times 7$ matrix of control information for printing the strings contained in _pmsgstr.
[L,1] horizontal location of lower left corner of string.
[L,2] vertical location of lower left corner of string.
[ $\mathbf{L}, \mathbf{3}]$ character height in inches.
[L,4] angle in degrees to print string. This may be -180 to 180 relative to the positive X axis.
[L,5] location coordinate system.
1 location of string in plot coordinates
2 location of string in inches
[L,6] color.
[L,7] font thickness, may be 0 or greater. If 0 use normal line width.
_pmsgstr
_pnotify scalar, controls window output during the creation of the graph. Default is 1 .
0 no activity to the window while writing .tkf file 1 display progress as fonts are loaded, and .tkf file is being generated
_pnum
scalar, $2 \times 1$ or $3 \times 1$ vector for independent control for axes numbering. The first element controls the X axis numbers, the second controls the Y axis numbers, and the third (if set) controls the Z axis numbers. Default is 1 .

If this value is scalar, it will be expanded to a vector.
0 no axes numbers displayed
1 axes numbers displayed, vertically oriented on axis

## 2 axes numbers displayed, horizontally oriented on axis

For example:

```
_pnum = { 0, 2 }; /* no X axis numbers, */
    /* horizontal on Y axis */
```

_pnumht scalar, size of axes numbers in inches. If 0 , a size of .13 will be used. Default is 0 .
_protate
_pscreen
_psilent scalar. If 0 , a beep will sound when the graph is finished drawing to the window. Default is 1 (no beep).
_pstype scalar or $K \times 1$ vector, controls symbol used at data points. To use a single symbol type for all points, set this to one of the following scalar values:

| $\mathbf{1}$ | circle | $\mathbf{8}$ | solid circle |
| :--- | :--- | :--- | :--- |
| $\mathbf{2}$ | square | $\mathbf{9}$ | solid square |
| $\mathbf{3}$ | triangle | $\mathbf{1 0}$ | solid triangle |
| $\mathbf{4}$ | plus | $\mathbf{1 1}$ | solid plus |
| $\mathbf{5}$ | diamond | $\mathbf{1 2}$ | solid diamond |
| $\mathbf{6}$ | inverted triangle | $\mathbf{1 3}$ | solid inverted triangle |
| $\mathbf{7}$ | star | $\mathbf{1 4}$ | solid star |

If this is a vector, each line will have a different symbol. Symbols will repeat if there are more lines than symbol types. Default is 0 (no symbols are shown).
_psurf $2 \times 1$ vector, controls 3-D surface characteristics.
[1] if 1 , show hidden lines. Default is 0 .
[2] color for base, see Colors, Section 25.5. The base is an outline of the X-Y plane with a line connecting each corner to the surface. If 0 , no base is drawn. Default is 7 .

## _psym $\quad \mathrm{M} \times 7$ matrix, M extra symbols will be plotted.

[M,1] x location.
[M,2] y location.
[M,3] symbol type, see _pstype earlier.
[M,4] symbol height. If this is 0 , a default height of 5.0 will be used.
[M,5] symbol color, see Colors, Section 25.5.
[M,6] type of coordinates:
1 plot coordinates
2 inch coordinates
[M,7] line thickness. A value of zero is normal line width.
_psym3d $\quad \mathrm{M} \times 7$ matrix for plotting extra symbols on a 3-D (surface or xyz) graph.
[M,1] x location in plot coordinates.
[ $\mathbf{M}, 2]$ y location in plot coordinates.
[ $\mathbf{M}, 3]$ z location in plot coordinates.
[M,4] symbol type, see _pstype earlier.
[M,5] symbol height. If this is 0 , a default height of 5.0 will be used.
[M,6] symbol color, see Colors, Section 25.5.
[M,7] line thickness. A value of 0 is normal line width.
Use _psym for plotting extra symbols in inch coordinates.
_psymsiz scalar or $K \times 1$ vector, symbol size for the symbols on the main curves. This is NOT related to _psym. If 0 , a default size of 5.0 is used.
_ptek string, name of Tektronix format graphics file. This must have a .tkf extension. If this is set to a null string, the graphics file will be suppressed. The default is graphic.tkf.
_pticout scalar. If 1, tick marks point outward on graphs. Default is 0 .
_ptitlht scalar, the height of the title characters in inches. If this is 0 , a default height of approx. 0.13 inch will be used.
_pversno string, the graphics version number.
_pxpmax scalar, the maximum number of places to the right of the decimal point for the X axis numbers. Default is 12 .
_pxsci scalar, the threshold in digits above which the data for the X axis will be scaled and a power of 10 scaling factor displayed. Default is 4 .
_pypmax scalar, the maximum number of places to the right of the decimal point for the Y axis numbers. Default is 12 .
_pysci scalar, the threshold in digits above which the data for the Y axis will be scaled and a power of 10 scaling factor displayed. Default is 4 .
_pzclr scalar, row vector, or $K \times 2$ matrix, $Z$ level color control for procedures surface and contour. (See surface in the GAUSS Language Reference.)
_pzoom
$1 \times 3$ row vector, magnifies the graphics display for zooming in on detailed areas of the graph. If scalar 0 , no magnification is performed. Default is 0 .
[1] magnification value. 1 is normal size.
[2] horizontal center of zoomed plot (0-100).
[3] vertical center of zoomed plot (0-100).
To see the upper left quarter of the screen magnified 2 times use:

$$
\text { _pzoom }=\{22575\} ;
$$

_pzpmax scalar, the maximum number of places to the right of the decimal point for the Z axis numbers. Default is 3 .
_pzsci scalar, the threshold in digits above which the data for the Z axis will be scaled and a power of 10 scaling factor displayed. Default is 4 .


## Graphics Editor

### 26.1 Introduction to the Graphics Editor

The GAUSS graphics editor is a utility for composing pages containing GAUSS graphics files. Its primary purpose is to provide the user with a toolbox for creating and annotating graphs created by GAUSS using all of the fonts available on your Windows system. It is not meant to be a full-featured publishing tool but rather a supplemental utility for dynamically importing and easily arranging multiple graphics files on a single page.

### 26.1.1 Overview

The graphics editor allows the user to interactively create any number of graphical objects for composing documents. It is launched by selecting Tools from the GAUSS menu bar, then Graphics Editor, or by clicking on the Graphics Editor icon on the GAUSS toolbar.

Once the document has been created, it may be saved for later modification. All of the objects and their respective properties contained in the document are preserved. The document may also be exported to other formats.

### 26.2 Graphics Editor Workspace

The graphics editor workspace is a window allowing access to a single page with tools for composing the document. The page is defined by user-defined properties such as page orientation and margin settings.

It provides a dialog bar for user-selection of the current pen and brush properties.
Zoom capability is provided for detailed accuracy and accomodating a wide-variety of computer display resolutions.


Figure 26.1: Graphics Editor Workspace

### 26.2.1 Toolbar

The toolbar is displayed across the top of the application window, below the menu bar. The toolbar provides quick mouse access to many tools used in the graphics editor.

To hide or display the toolbar, choose Toolbar from the View menu (ALT, V, T).


Figure 26.2: Graphics Editor Toolbar

### 26.2.2 Status Bar

The status bar is displayed at the bottom of the graphics editor window. To display or hide the status bar, use the Status Bar command in the View menu.

The left area of the status bar describes actions of menu items as you use the arrow keys to navigate through menus. This area similarly shows messages that describe the actions of toolbar buttons as you depress them, before releasing them. If after viewing the description of the toolbar command you wish not to execute the command, then release the mouse button while the pointer is off the toolbar button.

In addition, the status bar provides short hints while using the graphical interface such as defining, sizing, and moving objects.


Figure 26.3: Graphics Editor Status Bar

## Indicator Description

The remaining status bar panes indicate the following:

The $x / y$ mouse position on the page (in units specified in the View Properties menu).
CAPS indicates the CAPS LOCK key is latched down.
NUM indicates the NUM LOCK key is latched down.
SCRL indicates the SCROLL LOCK key is latched down.

### 26.2.3 File menu commands

The File menu offers the following commands:
\(\left.$$
\begin{array}{ll}\text { New } & \text { Opens a new, untitled document (CTRL+N). } \\
\text { Open } & \text { Opens an existing document in a new window (CTRL+O). } \\
\text { Import } & \begin{array}{l}\text { Imports a file of another format. Currently only the GAUSS graphics format } \\
\text {.tkf is supported. }\end{array} \\
\text { Save } & \begin{array}{l}\text { Saves the active document to its current name and directory. When you save } \\
\text { a document for the first time, the graphics editor displays the Save As dialog } \\
\text { box so you can name your document (CTRL+S). If you want to change the } \\
\text { name and directory of an existing document before saving it, choose the }\end{array}
$$ <br>

Save As command.\end{array} \quad $$
\begin{array}{l}\text { Saves and names the active document. The graphics editor displays the Save }\end{array}
$$\right\}\)| As dialog box so you can name your document. |
| :--- |


| Print Setup | Allows you to select a printer and printer connection. This command <br> presents a Print Setup dialog box where you specify the printer and its <br> connection. |
| :--- | :--- |
| Exit | Ends the graphics editor session. The graphics editor prompts you to save <br> documents with unsaved changes (ALT+F4). |

### 26.2.4 Edit menu commands

The Edit menu offers the following commands:

| Undo | Reverses the last editing action, if possible (CTRL+Z or <br> ALT+BACKSPACE). The name of the command changes depending on <br> what the last action was. The Undo command changes to Can't Undo on the <br> menu if the last action cannot be reversed. |
| :--- | :--- |
| Cut | Removes the currently selected data from the document and put it on the <br> clipboard (CTRL+X). This command is unavailable if there is no data <br> currently selected. Cutting data to the clipboard replaces the contents <br> previously stored there. |
| Copy | Copies currently selected data onto the clipboard (CTRL+C). This command <br> is unavailable if there is no data currently selected. Copying data to the <br> clipboard replaces the contents previously stored there. |
| Paste | Inserts a copy of the clipboard contents at the insertion point (CTRL+V). <br> This command is unavailable if the clipboard is empty. |

### 26.2.5 View menu commands

The View menu offers the following commands:

Toolbar Displays and hides the Toolbar, which includes buttons for some of the most common commands such as File Open. A check mark appears next to the
menu item when the Toolbar is displayed. See Toolbar, Section ??, for help
on using the toolbar.

Status Bar $\quad$| Displays and hides the Status Bar, which describes the action to be executed |
| :--- |
| by the selected menu or depressed toolbar button and keyboard latch state. A |
| check mark appears next to the menu item when the Status Bar is displayed. |
| See Status Bar, Section ??, for help on using the status bar. |

Properties

Zoom | Allows you to change user-defined page/view settings. See Setting the |
| :--- |
| Page/View Properties, Section 26.2 .11 , for more information. |

Allows you to change user-defined zoom control. See Using the Zoom
Feature, Section 26.2 .11, for more information.

### 26.2.6 Draw menu commands

The Draw menu offers the following commands; see Graphical Objects, Section 26.2.13, for more detailed information on each.

Select Puts the editor into object selection state.
TKF Graphics Opens a GAUSS graphics window.
Window

Line Draws a line.

Arrow Draws an arrow.

Rectangle Draws a rectangle.
Ellipse Draws an ellipse.

Text Allows you to enter text.

### 26.2.7 Export menu commands

The Export menu offers the following commands; see File Management, Section 26.3, for more detailed information on each.

| Encapsulated <br> Postscript | Writes an Encapsulated Postscript file. |
| :--- | :--- |
| JPEG Image | Writes a JPEG compressed image file. |
| Windows <br> Metafile | Writes a Windows Enhanced Metafile. |

### 26.2.8 Help menu commands

The Help menu offers the following commands, which provide you assistance with this application:

$$
\begin{array}{ll}
\text { Help Topics } & \begin{array}{l}
\text { Displays the opening screen of Help. From the opening screen, you can } \\
\text { jump to step-by-step instructions for using the graphics editor and various } \\
\text { types of reference information. Once you open Help, you can click the }
\end{array} \\
\text { Contents button whenever you want to return to the opening screen. }
\end{array}
$$

About Displays the copyright notice and version number of this application.

### 26.2.9 Object Action Context Menu

Once an object has been selected, its action context menu may be displayed by right-clicking inside the object.

The following actions may be selected from this menu:

Refresh Redraws the object.

Cut | Removes the currently selected data from the document and put it on the |
| :--- |
| clipboard. This command is unavailable if there is no data currently selected. |
| Cutting data to the clipboard replaces the contents previously stored there. |
| Copy |
| Copies currently selected data onto the clipboard. This command is |
| unavailable if there is no data currently selected. |
| Deletes currently selected data. This command is unavailable if there is no |
| data currently selected. |
| Z-Order |
| Changes the objects position in the z-order of the document's list. The |
| z-order allows the user to control in what order the object is drawn on the |
| page. To change, select the Z-Order menu item from the action context menu |
| and select one of the following: |
| Move to Top - Moves the object to the top of the list. |
| Move to Bottom - Moves the object to the bottom of the list. |

Edit $\quad$\begin{tabular}{l}
Allows you to modify the object. This menu item currently applies to text <br>
objects only. <br>
Deselect

 

De-selects the object.
\end{tabular}

Oroperties

### 26.2.10 Page Context Menu

The Page Context menu is displayed by pressing the right mouse button when no object is selected.
The following actions may be selected from this menu:

| Paste | Copies an object from the clipboard to the page if one is available. |
| :--- | :--- |
| Retain Aspect | Check or uncheck the aspect ratio state. When checked, this forces the <br> object to retain its aspect ratio while sizing it. |
| Ratio | a |

See Modifying the Graphical Objects, Section 26.2.14, for more information.

### 26.2.11 Setting the Page/View Properties

The following describes how to set various page and view properties, including how to set the page orientation and margins, use the zoom feature, and set the color options.

## Setting the Page Orientation and Margins

Page Orientation The document page orientation of landscape or portrait is set from the Properties dialog under the View menu.

Measure Units Allows all coordinates and measurements to be in inches or centimeters.
$\begin{array}{ll}\text { Reference } & \text { A reference margin indicating the document's current orientation and } \\ \text { Margin } & \text { margin settings is also available. This is useful for customizing your page to }\end{array}$ be compatible with the printer currently in use. Because printer margins vary from one printer to another, it is useful to be able to set your page to the margins that most accurately match your printer.

The reference margin settings are available in the Properties dialog under the View menu.

Reset Colors Pushing this button will reset the available colors to the initial IBM 16-color scheme. Each color may be set to a custom color; see Pen/Fill Properties, Section 26.2.12 for more information.

## Using the Zoom Feature

To set the zoom, click the zoom drop-down control on the toolbar or select Zoom in the View menu.

## Setting the Color Options

You may reset the colors to the original IBM 16-color scheme by pressing the Reset Colors to Initial button.

### 26.2.12 Setting the Pen/Fill Properties

All drawing is done with a currently selected pen and brush. The current pen has a width attribute and color attribute.

The current brush (for painting object backgrounds) has a color attribute.

## Setting the Pen Color

Left-click in one of the color boxes in the dialog bar to set the current color. The dialog bar is located to the left of the drawing area. The color is immediately displayed in the sample box at the top of the color box area.

Double-clicking in the color box will allow you you to customize that particular color.

## Setting the Pen Width

Left-click in one of the width boxes in the dialog bar shown below the color boxes.

## Setting the Fill (brush) Color

Right-click in one of the color boxes in the dialog bar to set the current fill color. The dialog bar is located to the left of the drawing area. The fill color is immediately displayed in the sample box at the top of the color box area.

Transparent Fill To set a transparent fill color, right-click in the sample box at the top of the color boxes. This will cause the drawing object to have no fill associated with it and allow objects beneath it to show through.
$\begin{array}{ll}\text { Customizing } & \begin{array}{l}\text { Double-clicking in the color box will allow you to customize that particular } \\ \text { color. }\end{array}\end{array}$

### 26.2.13 Graphical Objects

The graphics editor allows the user to interactively create any number of the following graphical objects for composing your document:

## Creating a TKF Graphics window

A TKF Graphics Window is a window object containing a GAUSS-generated graphics file.
To create a TKF graphics window, select the Graphics Window menu item from the Draw menu or press the Create TKF window icon on the toobar.

A properties dialog is presented which allows the selection of a GAUSS-generated TKF graphics file. There are two ways to do this. If one or more graphics files are currently being displayed in GAUSS, those filenames will appear in the Active Graphs drop-down control and may be selected. Otherwise, pressing the Browse button will present a common open file dialog for selecting the file.

The graphics window border and fill colors may be set using their respective Color buttons.

Once the file has been selected and the user presses OK, the graphics window is created in a default size located at the top-left corner of the page.

At this time, the window object may be modified.

## Creating a Text Object

To create a text object, select the Text menu item from the Draw menu or press the Draw text button on the toolbar.

Next, position the mouse where you want the top-left corner of your text then press and hold the left mouse button. (You may also move the object by pressing and holding the right mouse button while keeping the left button depressed). Drag the mouse to the bottom-left corner and let up on the mouse button.

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The text region will be redrawn in the current background fill color and contain a text cursor inside. At this point you may enter your text at the cursor using the last selected text font.

If the text requires more lines than the current bounding box allows, the box will be resized as needed.

To save your text when you have finished typing, press the OK icon on the text toolbar indicated by a green check mark or press the SHIFT+ENTER key. Clicking the mouse button outside the text window will also save the text and complete the operation.

The box will be redrawn with the proper font background, border color and margin settings.

To cancel out of the text and lose changes, press the Cancel icon on the text toolbar indicated by a red X or press the ESCAPE key.

Note: The text object may be rotated at any angle from the Object Properties menu.

## Creating a Line

To create a line, select the Line menu item from the Draw menu or press the Draw line button on the toolbar. This puts the editor into the draw line state indicated by a crosshair cursor.

Next, position the mouse where you want the first end point of the line then press and hold the left mouse button. (You may also move the line by pressing and holding the right mouse button while keeping the left button depressed). Drag the mouse to the location for the second end point and release the mouse button.

Note: Pressing the CTRL key while defining a line or arrow forces the line to be vertical or horizontal.

The line will be redrawn in the current pen color.
At any time after the above process you may modify the line object.

## Creating an Arrow

To create an arrow, select the Arrow menu item from the Draw menu or press the Draw arrow button on the toolbar. This puts the editor into the draw arrow state indicated by a crosshair cursor. Arrow style controls for defining the arrow appear on the dialog bar to the left of the drawing area.

Defining the endpoints are identical to the steps for defining a line above. However, the arrow head size and shape may be set using the additional arrow style controls in the dialog bar.

Once the second endpoint is defined, the arrow will be redrawn in the current pen color.
At any time after the above process you may modify the arrow object.

## Arrow Styles

There are a combination of two styles of arrows: Open/Closed, and Single/Double-headed arrows.
A closed arrow is one whose arrowhead is filled in with the current pen color. An open one has no fill.

A single-headed arrow is a line with an arrowhead on one end. A double-headed arrow has an arrowhead at both ends.

## Creating a Rectangle

To create a rectangle object, select the Rectangle menu item from the Draw menu or press the Draw rectangle button on the toolbar. This puts the editor into the draw rectangle state indicated by a crosshair cursor.

Next, position the mouse where you want the top-left corner then press and hold the left mouse button. (You may also move the object by pressing and holding the right mouse button while keeping the left button depressed). Drag the mouse to the bottom-left corner and release the mouse button.

The rectangle will be redrawn in the current background fill and border color.

At any time after the above process you may modify the rectangle object.

## Creating an Ellipse

Note: a circle is first created when defining an ellipse. After the circle has been defined it may be dynamically reshaped into an ellipse of the desired size using the mouse.

To create the circle, select the Ellipse menu item from the Draw menu or press the Draw ellipse button on the toolbar. This puts the editor into the draw ellipse state indicated by a crosshair cursor.

Next, position the mouse where you want the center of the circle then press and hold the left mouse button. (You may also move the object by pressing and holding the right mouse button while keeping the left button depressed). Drag the mouse to the desired radius and release the mouse button.

The circle will be redrawn in the current background fill and border color.

At any time after the above process you may modify the circle to any other elliptical size and shape.

### 26.2.14 Modifying the Graphical Objects

First, ensure you are in selection mode by pressing the Select toolbar button or choosing the Select item in the Draw menu. Selection mode is indicated with an arrow cursor.

Next, select the object you want to modify by left-clicking anywhere inside or on the object. It will then become highlighted.

Once selected, it may be sized, moved, or modified with one of the actions listed in the object's action context menu.

## Aspect Ratio

You can force an object's aspect ratio to be retained while sizing it by checking this menu item from the Page Context menu.

When this is checked, all the sides of the object are sized by the same amount as the side being moved, eliminating the need to resize all sides independently.

If the object is being sized by a corner point, this feature is ignored, eliminating the need for the user to check/uncheck the aspect ratio menu item needlessly.

The aspect ratio feature has no effect for lines and arrows.

## Object Properties

The properties dialog box allows you to modify various attributes of the object depending on its type.

See Graphical Objects, Section 26.2.13, for more information about object properties.

## Sizing an Object

First select the object.
Next, click and hold the left mouse button in one of the object's highlight points. Depending on the type of object, the new size is defined by how you move the mouse. Lifting the mouse button sets the new size and causes it to be redrawn.

If the object is a TKF graphics window, text object, rectangle, or ellipse, then the aspect ratio may be retained depending on the selection state of the aspect ratio menu item state when grabbing one of the four sides.

Grabbing the corner of an object allows you to size it in any direction ignoring the state of the aspect ratio menu.

If the object is a line or arrow, then it may be moved during the sizing operation by pressing and holding the right mouse button while still depressing the left mouse button.

Note: A rotated text object may not be sized. It may only be sized in a non-rotated state.

## Moving an Object

First select the object.

Next, click and hold the left mouse button somewhere inside the object. Drag the object to the new location and lift the mouse button. The object is then redrawn in the new location.

If the object is a line or arrow, then it may also be moved during the sizing operation.

### 26.3 File Management

The graphics editor stores the document as a list of vector-based graphical objects. These are binary files and cannot be edited by hand. It uses a default extension of .pge.

See the File menu command for available file operations.

You may export your document to other formats for inclusion in web pages, word-processors, and publishing applications.

### 26.3.1 Exporting Files

The Export menu enables you to easily export graphic files to some of the most frequently used graphic formats.

## Writing an Encapsulated Postscript Image

To write an Encapsulated Postscript file, select the Encapsulated Postscript menu item from the Export menu. This displays the entire file dialog.

| Filename | Enter or browse to the desired output filename. The default extension is <br> .eps. |
| :--- | :--- |
| Convert lines <br> to Black | Check this item if you want to convert all colors in the image to black. |
| Scale Factor | Enter a scale factor if necessary. By default, the graphics editor uses a very <br> high internal resolution for the best possible quality. However, some <br> applications are unable to correctly scale the data when importing. This may <br> be worked around by scaling the data during the export stage. |
| Minimum line | Enter the minimum line width value if you want to darken the lines in <br> the exported file. |

## Note About Fonts

Although the fonts you select for your text box may appear fine in the graphics editor, it is possible the target application importing it may not interpret them correctly. Every application has its own EPS interpreter, and the availability of your font depends on it. If you are having problems of this type, try using the Enhanced Metafile format conversion. This format has no such font limitations.

## Writing a JPEG Image

A JPEG image file is a widely used bitmap format for inclusion in web pages due to its compression characteristics.

To write a JPEG file, select the JPEG Image File menu item from the Export menu.
Note: Because JPEG is a bitmap format, the image written to the file is exactly as seen on the display. Thus you may be requried to zoom out on some lower-resolution displays to obtain an image of the entire document.

## Writing a TIFF Image

A TIFF (Tag Image File Format) image file is an older but widely used bitmap format.
To write a TIFF file, select the TIFF Image File menu item from the Export menu.
Note: Because TIFF is a bitmap format, the image written to the file is exactly as seen on the display. Thus you may be required to zoom out on some lower-resolution displays to obtain an image of the entire document.

## Windows Metafile

An enhanced metafile is a vector-based file and is considered the best method for export/import on the Microsoft Windows platform.

To create a Windows Enhanced Metafile, select the Windows Metafile menu item from the Export menu.

## Autoscale

Checking this option forces the translated to automatically scale the Enhanced Metafile. This is the best setting for most applications for importing. However, some applications require a more precise format. If the importing application has trouble, uncheck this option.

## Time and Date

## 27

GAUSS offers a comprehensive set of time and date functions. These functions afford the user the ability to return the current time and date, to carry out most related calculations and format the results for output. GAUSS also allows the user to perform timed iterations.

In the year 1 AD the calendar in general use was the Julian calendar. The Gregorian calendar that we use today was not invented until the late 1500 's. This new calendar changed the method of calculating leap years on century marks. With the Julian system simply every fourth year was a leap year. The Gregorian system made every fourth year a leap year with the exception of century marks which are only leap years if divisible by 400 . The British adoption of this calendar, which the GAUSS date functions are based on, did not happen until the year 1752. In that year eleven days were removed; September 2, 1752 was followed by September 14, 1752.
dtvnormal and utctodtv are accurate back to 1 AD. The rest of the GAUSS date functions assume a normal Gregorian system regardless of year. Thus, they will not account for the days taken out in September of 1752, nor will they account for all century marks being leap years before the adoption of the Gregorian system in 1752.

The time is given by your operating system, daylight savings time is not automatically accounted for by GAUSS in calculations.

### 27.1 Time and Date Formats

The Time and Date formats in GAUSS fall into one of two major categories, matrix/vector and string. The matrix/vector formats can be used for either calculations or if desired for output. The string formats are, however, mostly for use as ouput. Some manipulation of strings is possible with the use of the stof function.

A $4 \times 1$ vector is returned by both the date and time functions.

```
d = date;
d;
    1997.00 /* Year */
    5.00000 /* Month */
    29.0000 /* Day */
    56.4700 /* Hundredths of a second since midnight */
    t = time;
    t;
    10.00 /* Hours since midnight */
    17.00 /* Minutes */
    33.00 /* Seconds */
    13.81 /* Hundredths of a second */
```

These vectors can be written to a string of the desired form by passing them through the corresponding function.

```
d = { 1997, 5, 29, 56.47 };
datestr(d);
```

5/29/97
datestrymd(d);

```
19970529
t = { 10, 17, 33, 13.81};
timestr(t);
```

10:17:33

A list and brief description of these, and other related functions is provided in the table in section 27.2.

Another major matrix/vector format is the dtv, or date and time vector. The dtv vector is a $1 \times 8$ vector used with the dtvnormal and utctodtv functions. The format for the dtv vector is:

| Year | Month | Day | Hour | Min | Sec | DoW | DiY |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| 1955 | 4 | 21 | 4 | 16 | 0 | 4 | 110 |

Where:

Year Year, four digit integer.
Month 1-12, Month in year.
Day 1-31, Day of month.
Hour 0-23, Hours since midnight.
Min 0-59, Minutes.
Sec 0-59, Seconds.
DoW 0-6, Day of week, $0=$ Sunday.
DiY 0-365, Days since Jan 1 of current year.
dtvnormal normalizes a date. The last two elements are ignored for input, as shown in the following example. They are set to the correct values on output. The input can be $1 \times 8$ or $\mathrm{N} \times 8$.

```
dtv = { 1954 3 17 4 16 0 0 0 };
dtv = dtvnormal(dtv);
```

```
    1954
    dtv[3] = dtv[3] + 400;
    print dtv;
1954
    dtv = dtvnormal(dtv);
    print dtv;
1955}4
```


### 27.2 Time and Date Functions

Following is a partial listing of the time and date functions available in GAUSS.

| datestr | Formats a Date vector to a string (mo/dy/yr). |
| :--- | :--- |
| datestrymd | Formats a Date vector to an eight character string of the type <br> yyyymmdd. |
| dayinyr | Returns day number in the year of a given date. |
| _daypryr | Returns the number of days in the years given as input. |
| dtvnormal | Normalizes a $1 \times 8$ dtv vector. |
| etdays | Computes the difference in days between two dates. |
| ethsec | Computes the difference between two times in hundredths of a <br> second. |
| etstr | Formats a time difference measured in hundreths of a second to a <br> string. |


| _isleap | Returns a vector of ones and zeros, 1 if leap year 0 if not. |
| :--- | :--- |
| timestr | Formats a Time vector to a string hr:mn:sc. |
| timeutc | Universal time coordinate, number of seconds since January 1, 1970 <br> Greenwich Mean Time. |
| utctodtv | Converts a scalar, number of seconds since, or before, Jan 1 1970 <br> Greenwich mean time, to a dtv vector. |

Below is an example of two ways to calculate a time difference.

```
        d1 = { 1996, 12, 19, 82 };
        d2 = { 1997, 4, 28, 4248879.3 };
        dif = ethsec(d1,d2);
        ds = etstr(dif);
\[
d i f=1.1274488 e+09
\]
\[
d s=130 \text { days } 11 \text { hours } 48 \text { minutes } 7.97 \text { seconds }
\]
```

If only the number of days is needed use etdays.

```
    d1 = { 1996, 12, 19, 82 };
    d2 = { 1997, 4, 28, 4248879.3 };
    dif = etdays(d1,d2);
dif = 130.00000
```

The last element of $d l$ is optional when used as an input for etdays.
_isleap returns a matrix of ones and zeros, ones when the corresponding year is a leap year.

```
x = seqa(1970,1,20);
y = _isleap(x);
delif(x,abs(y-1));
1972.0000 /* Vector containing all leap years
1976.0000 between 1970 - 1989 */
1980.0000
1984.0000
1988.0000
```

To calculate the days of a number of consecutive years:

```
x = seqa(1983,1,3);
y = _daypryr(x);
sumc (y);
```

1096.0000

To add a portion of the following year:

```
g = { 1986, 2, 23, 0 };
dy = dayinyr(g);
sumc(y)+dy;
```

1150.0000

For more information on any of these functions see their respective pages in the command reference.

### 27.2.1 Timed Iterations

Iterations of a program can be timed with the use of the hsec function in the following manner.

```
et = hsec; /* Start timer */
/* Segment of code to be timed */
et = (hsec-et)/100; /* Stop timer, convert to seconds */
```

In the case of a program running from one day into the next you would need to replace the hsec function with the date function. The ethsec function should be used to compute the time difference; a straight subtraction as in the previous example will not give the desired result.

```
dstart = date; /* Start timer */
/* Segment of code to be timed */
dend = date; /* Stop timer */
dif = ethsec(dstart,dend)/100; /* Convert time difference to seconds */
```



## ATOG

28

ATOG is a stand-alone conversion utility that converts ASCII files into GAUSS data sets. ATOG can convert delimited and packed ASCII files into GAUSS data sets. ATOG can be run from a batch file or the command line; it is not run from a GAUSS prompt but rather from a command prompt window.

The syntax is:
atog cmdfile
where cmdfile is the name of the command file. If no extension is given, . cmd will be assumed. If no command file is specified, a command summary will be displayed.

### 28.1 Command Summary

The following commands are supported in ATOG:

| append | Append data to an existing file. |
| :--- | :--- |
| complex | Treat data as complex variables. |
| input | The name of the ASCII input file. |
| invar | Input file variables (column names). |
| msym | Specify missing value character. |
| nocheck | Don't check data type or record length. |
| output | The name of the GAUSS data set to be created. |
| outtyp | Output data type. |
| outvar | List of variables to be included in output file. |
| preservecase | Preserve case of variable names in output file. |

The principle commands for converting an ASCII file that is delimited with spaces or commas are given in the following example:

```
input agex.asc;
output agex;
invar $ race # age pay $ sex region;
outvar region age sex pay;
outtyp d;
```

In this example, a delimited ASCII file agex. asc is converted to a double precision GAUSS data file agex.dat. The input file has five variables. The file will be interpreted as having five columns:

| column | name | data type |
| :---: | :--- | :--- |
| 1 | race | character |
| 2 | AGE | numeric |
| 3 | PAY | numeric |
| 4 | sex | character |
| 5 | region | character |

The output file will have four columns since the first column of the input file (race) is not included in the output variables. The columns of the output file are:

| column | name | data type |
| :---: | :--- | :--- |
| 1 | region | character |
| 2 | AGE | numeric |
| 3 | sex | character |
| 4 | PAY | numeric |

The variable names are saved in the file header. Unless preservecase has been specified, the names of character variables will be saved in lowercase, and the names of numeric variables will be saved in uppercase. The $\$$ in the invar statement specifies that the variables that follow are character type. The \# specifies numeric. If \$ and \# are not used in an invar statement, the default is numeric.

Comments in command files must be enclosed between ' $@$ ' characters.

### 28.2 Commands

A detailed explanation of each command follows.

## append

Instructs ATOG to append the converted data to an existing data set:
append;

No assumptions are made regarding the format of the existing file. Make certain that the number, order, and type of data converted match the existing file. ATOG creates v96 format data files, so will only append to v96 format data files.

## complex

Instructs ATOG to convert the ASCII file into a complex GAUSS data set:

```
complex;
```

Complex GAUSS data sets are stored by rows, with the real and imaginary parts interleaved, element by element. ATOG assumes the same structure for the ASCII input file, and will thus read TWO numbers out for EACH variable specified.
complex cannot be used with packed ASCII files.

## input

Specifies the file name of the ASCII file to be converted. The full path name can be used in the file specification.

For example, the command:

```
input data.raw;
```

will expect an ASCII data file in the current working directory.
The command:
input /research/data/myfile.asc;
specifies a file to be located in the /research/data subdirectory.

28-4

## invar

Soft Delimited ASCII Files Soft delimited files may have spaces, commas, or cr/lf as delimiters between elements. Two or more consecutive delimiters with no data between them are treated as one delimiter. For example:

```
invar age $ name sex # pay var[1:10] x[005];
```

The invar command above specifies the following variables:

| column | name | data type |
| :---: | :--- | :--- |
| 1 | AGE | numeric |
| 2 | name | character |
| 3 | sex | character |
| 4 | PAY | numeric |
| 5 | VAR01 | numeric |
| 6 | VAR02 | numeric |
| 7 | VAR03 | numeric |
| 8 | VAR04 | numeric |
| 9 | VAR05 | numeric |
| 10 | VAR06 | numeric |
| 11 | VAR07 | numeric |
| 12 | VAR08 | numeric |
| 13 | VAR09 | numeric |
| 14 | VAR10 | numeric |
| 15 | X001 | numeric |
| 16 | X002 | numeric |
| 17 | X003 | numeric |
| 18 | X004 | numeric |
| 19 | X005 | numeric |

As the input file is translated, the first 19 elements will be interpreted as the first row (observation), the next 19 will be interpreted as the second row, and so on. If the number of elements in the file is not evenly divisible by 19 , the final incomplete row will be dropped and a warning message will be given.

Hard Delimited ASCII Files Hard delimited files have a printable character as a delimiter between elements. Two delimiters without intervening data between them will be interpreted as a missing. If $\backslash \mathbf{n}$ is specified as a delimiter, the file should have one element per line and blank lines will be considered missings. Otherwise, delimiters must be printable characters. The dot '.' is illegal and will always be interpreted as a missing value. To specify the backslash as a delimiter, use $\backslash \backslash$. If $\backslash \mathbf{r}$ is specified as a delimiter, the file will be assumed to contain one case or record per line with commas between elements and no comma at the end of the line.

For hard delimited files the delimit subcommand is used with the invar command. The delimit subcommand has two optional parameters. The first parameter is the delimiter. The default is a comma. The second parameter is an ' $N$ '. If the second parameter is present, ATOG will expect N delimiters. If it is not present, ATOG will expect $\mathrm{N}-1$ delimiters.

This example:

```
invar delimit(, N) $ name # var[5];
```

will expect a file like this:

| BILL, | 222.3, | 123.2, | 456.4, | 345.2, | 533.2, |
| :--- | ---: | ---: | ---: | ---: | ---: |
| STEVE, | 624.3, | 340.3, | , | 624.3, | 639.5, |
| TOM, | 244.2, | 834.3, | 602.3, | 333.4, | 822.5, |

while

```
invar delimit(,) $ name # var[5];
```

or

```
invar delimit $ name # var[5];
```

will expect a file like this:

```
BILL , 222.3, 123.2, 456.4, 345.2, 533.2,
STEVE, 624.3, 340.3, , 624.3, 639.5,
TOM , 244.2, 834.3, 602.3, 333.4, 822.5
```

The difference between specifying N or $\mathrm{N}-1$ delimiters can be seen here:

$$
\begin{array}{rll}
456.4, & 345.2, & 533.2, \\
, & 624.3, & 639.5, \\
602.3, & 333.4, &
\end{array}
$$

If the invar statement specified three variables and $\mathrm{N}-1$ delimiters, this file would be interpreted as having three rows containing a missing in the $[2,1]$ element and the $[3,3]$ element like this:

```
456.4 345.2 533.2
    624.3 639.5
602.3 333.4
```

If N delimiters had been specified, this file would be interpreted as having two rows, and a final incomplete row that is dropped:
$\begin{array}{lll}456.4 & 345.2 & 533.2\end{array}$
$624.3 \quad 639.5$

The spaces were shown only for clarity and are not significant in delimited files so:

```
BILL,222.3,123.2,456.4,345.2,533.2,
STEVE,624.3,340.3, ,624.3,639.5,
TOM,244.2,834.3,602.3,333.4,822.5
```

would work just as well.
Linefeeds are significant only if $\backslash \boldsymbol{n}$ is specified as the delimiter, or when using $\backslash \mathbf{r}$. This example:

```
invar delimit(\r) $ name # var[5];
```

will expect a file with no comma after the final element in each row:

| BILL , | 222.3, | 123.2, | 456.4, | 345.2, | 533.2 |
| :--- | :--- | :--- | :--- | :--- | :--- |
| STEVE, | 624.3, | 340.3, | 245.3, | 624.3, | 639.5 |
| TOM , 244.2, | 834.3, | 602.3, | 333.4, | 822.5 |  |

Packed ASCII Files Packed ASCII files must have fixed length records. The record subcommand is used to specify the record length, and variables are specified by giving their type, starting position, length, and the position of an implicit decimal point if necessary.
outvar is not used with packed ASCII files. Instead, invar is used to specify only those variables to be included in the output file.

For packed ASCII files the syntax of the invar command is as follows:
invar record $=$ reclen (format) variables (format) variables;
where,
reclen the total record length in bytes, including the final carriage return/line feed if applicable. Records must be fixed length.
format (start,length.prec) where:
start starting position of the field in the record, 1 is the first position. The default is 1 .
length the length of the field in bytes. The default is 8 .
prec optional; a decimal point will be inserted automatically prec places in from the RIGHT edge of the field.

If several variables are listed after a format definition, each succeeding field will be assumed to start immediately after the preceding field. If an asterisk is used to specify the starting position, the current logical default will be assumed. An asterisk in the length position will select the current default for both length and prec. This is illegal: (3,8.*).

The type change characters \$ and \# are used to toggle between character and numeric data type.
Any data in the record that is not defined in a format is ignored.
The examples below assume a 32-byte record with a carriage return/line feed occupying the last 2 bytes of each record. The data below can be interpreted in different ways using different invar statements:


This example:

```
invar record=32 $(1,3) group dept #(11,4.2) x[3] (*,5) y;
```

will result in:

| variable | value | type |
| :--- | :--- | :--- |
| group | ABC | character |
| dept | DEF | character |
| X1 | 12.34 | numeric |
| X2 | 56.78 | numeric |
| X3 | 90.12 | numeric |
| Y | 34567 | numeric |

This example:

```
invar record=32 $ dept (*,2) id # (*,5) wage (*,2) area
```

will result in:

| variable | value | type |
| :--- | :--- | :--- |
| dept | ABCDEFGH | character |
| id | IJ | character |
| WAGE | 12345 | numeric |
| AREA | 67 | numeric |

## msym

Specifies the character in the input file that is to be interpreted as a missing value. This example:

```
msym &;
```

defines the character '\&' as the missing value character. The default '.' (dot) will always be interpreted as a missing value unless it is part of a numeric value.

## nocheck

Optional; suppresses automatic checking of packed ASCII record length and output data type. The default is to increase the record length by 2 bytes if the second record in a packed file starts with $\mathrm{cr} / \mathrm{lf}$, and any files that have explicitly defined character data will be output in double precision regardless of the type specified.

## output

The name of the GAUSS data set. A file will be created with the extension . dat. For example:
output /gauss/dat/test;
creates the file test. dat on the /gauss/dat directory.

## outtyp

Selects the numerical accuracy of the output file. Use of this command should be dictated by the accuracy of the input data and storage space limitations. The format is:

```
outtyp fmt;
```

where fmt is:
D or 8 double precision
F or 4 single precision (default)
I or 2 integer
The ranges of the different formats are:

| bytes | data type | significant <br> digits | range |
| :--- | :--- | :--- | ---: |
|  |  | 4 | $-32768<=X<=32767$ |
| 2 | integer | 4 | $8.43 \times 10^{-37}<=\|X\|<=3.37 \times 10^{+38}$ |
| 4 | single precision | $6-7$ | $4.19 \times 10^{-307}<=\|X\|<=1.67 \times 10^{+308}$ |
| 8 | double precision | $15-16$ |  |

If the output type is integer, the input numbers will be truncated to integers. If your data has more than 6 or 7 significant digits, specify outtyp as double.

Character data require outtyp d. ATOG automatically selects double precision when character data is specified in the invar statement, unless you have specified nocheck.

The precision of the storage selected does not affect the accuracy of GAUSS calculations using the data. GAUSS converts all data to double precision when the file is read.

## outvar

Selects the variables to be placed in the GAUSS data set. The outvar command needs only the list of variables to be included in the output data set. They can be in any order. In this example:

```
invar $name #age pay $sex #var[1:10] x[005];
outvar sex age x001 x003 var[1:8];
```

the outvar statement selects the following variables:

| column | name | data type |
| :---: | :--- | :--- |
| 1 | sex | character |
| 2 | AGE | numeric |
| 3 | X001 | numeric |
| 4 | X003 | numeric |
| 5 | VAR01 | numeric |
| 6 | VAR02 | numeric |
| 7 | VAR03 | numeric |
| 8 | VAR04 | numeric |
| 9 | VAR05 | numeric |
| 10 | VAR06 | numeric |
| 11 | VAR07 | numeric |
| 12 | VAR08 | numeric |

outvar is not used with packed ASCII files.

## preservecase

Optional; preserves the case of variable names. The default is nopreservcase, which will force variable names for numeric variables to upper case and character variables to lower case.

### 28.3 Examples

Example 1 The first example is a soft delimited ASCII file called agex1.asc. The file contains seven columns of ASCII data:

```
Jan 167.3 822.4 6.34E06 yes 84.3 100.4
Feb 165.8 987.3 5.63E06 no 22.4 65.6
Mar 165.3 842.3 7.34E06 yes 65.4 78.3
```

The ATOG command file is agex1.cmd:

```
input /gauss/agex1.asc;
output agex1;
invar $month #temp pres vol $true var[02];
outvar month true temp pres vol;
```

The output data set will contain the following information:

| name | month | true | TEMP | PRES | VOL |
| :--- | :--- | :--- | ---: | ---: | ---: |
| case 1 | Jan | yes | 167.3 | 822.4 | $6.34 \mathrm{e}+6$ |
| case 2 | Feb | no | 165.8 | 987.3 | $5.63 \mathrm{e}+6$ |
| case 3 | Mar | yes | 165.3 | 842.3 | $7.34 \mathrm{e}+6$ |
| type | char | char | numeric | numeric | numeric |

The data set is double precision since character data is explicitly specified.
Example 2 The second example is a packed ASCII file xlod.asc The file contains 32-character records:


The ATOG command file is xlod.cmd:

```
input /gauss/dat/xlod.asc;
output xlod2;
invar record=32 \$(1,3) client[2] zone (*,1) reg \#(20,5) zip;
```

The output data set will contain the following information:

| name | client1 | client2 | zone | reg | ZIP |
| :--- | :--- | :--- | :--- | :--- | :--- |
| case 1 | AEG | DRF | CST | y | 60631 |
| case 2 | EDJ | TAJ | PST | n | 98064 |
| case 3 | GWD | NAD | MST | y | 59725 |
| type | char | char | char | char | numeric |

The data set is double precision since character data is explicitly specified.
Example 3 The third example is a hard delimited ASCII file called cplx. asc. The file contains six columns of ASCII data:

$$
\begin{array}{rrrrrr}
456.4, & 345.2, & 533.2, & -345.5, & 524.5, & 935.3, \\
-257.6, & 624.3, & 639.5, & 826.5, & 331.4, & 376.4, \\
602.3, & -333.4, & 342.1, & 816.7, & -452.6, & -690.8
\end{array}
$$

The ATOG command file is cplx.cmd:

```
input /gauss/cplx.asc;
output cplx;
invar delimit #cvar[3];
complex;
```

The output data set will contain the following information:

| name | cvar1 | cvar2 | cvar3 |
| :---: | ---: | ---: | ---: |
| case 1 | $456.4+345.2 \mathrm{i}$ | $533.2-345.5 \mathrm{i}$ | $524.5+935.3 \mathrm{i}$ |
| case 2 | $-257.6+624.3 \mathrm{i}$ | $639.5+826.5 \mathrm{i}$ | $331.4+376.4 \mathrm{i}$ |
| case 3 | $602.3-333.4 \mathrm{i}$ | $342.1+816.7 \mathrm{i}$ | $-452.6-690.8 \mathrm{i}$ |
| type | numeric | numeric | numeric |

The data set defaults to single precision, since no character data is present, and no outtyp command is specified.

### 28.4 Error Messages

atog - Can't find input file
The ASCII input file could not be opened.
atog - Can't open output file
The output file could not be opened.
atog - Can't open temporary file
Notify Aptech Systems.
atog - Can't read temporary file
Notify Aptech Systems.

## atog - Character data in output file Setting output file to double precision

The output file contains character data. The type was set to double precision automatically.

## atog - Character data longer than 8 bytes were truncated

The input file contained character elements longer than 8 bytes. The conversion continued and the character elements were truncated to 8 bytes.
atog - Disk Full
The output disk is full. The output file is incomplete.
atog - Found character data in numeric field

This is a warning that character data was found in a variable that was specified as numeric. The conversion will continue.

## atog - Illegal command

An unrecognizable command was found in a command file.

## atog - Internal error

Notify Aptech Systems.

## atog - Invalid delimiter

The delimiter following the backslash is not supported.

## atog - Invalid output type

Output type must be I, F, or D.

## atog - Missing value symbol not found

No missing value was specified in an msym statement.

## atog - No Input file

No ASCII input file was specified. The input command may be missing.

## atog - No input variables

No input variable names were specified. The invar statement may be missing.
atog - No output file
No output file was specified. The output command may be missing.
atog - output type d required for character data Character data in output file will be lost

Output file contains character data and is not double precision.

## atog - Open comment

The command file has a comment that is not closed. Comments must be enclosed in @'s:
@ comment @

```
atog - Out of memory
```

Notify Aptech Systems.

```
atog - read error
```

A read error has occurred while converting a packed ASCII file.

## atog - Record length must be 1-16384 bytes

The record subcommand has an out of range record length.

## atog - Statement too long

Command file statements must be less than 16384 bytes.

## atog - Syntax error at:

There is unrecognizable syntax in a command file.

## atog - Too many input variables

More input variables were specified than available memory permitted.
atog - Too many output variables
More output variables were specified than available memory permitted.

## atog - Too many variables

More variables were specified than available memory permitted.

## atog - Undefined variable

A variable requested in an outvar statement was not listed in an invar statement.

```
atog WARNING: missing ')' at:
```

The parentheses in the delimit subcommand were not closed.

## atog WARNING: some records begin with cr/lf

A packed ASCII file has some records that begin with a carriage return/linefeed. The record length may be wrong.
atog - complex illegal for packed ASCII file.
A complex command was encountered following an invar command with record specified.
atog - Cannot read packed ASCII. (complex specified)
An invar command with record specified was encountered following a complex command.

## Error Messages

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The following is a list of error messages intrinsic to the GAUSS programming language. Error messages generated by library functions are not included here.

## G0002 File too large

| load | Input file too large. |
| :--- | :--- |
| getf | Input file too large. |

G0003 Indexing a matrix as a vector
A single index can be used only on vectors. Vectors have only one row or only one column.

## G0004 Compiler stack overflow - too complex

An expression is too complex. Break it into smaller pieces. Notify Aptech Systems.

G0005 File is already compiled

## G0006 Statement too long

Statement longer than 4000 characters.

## G0007 End of file encountered

## G0008 Syntax error

| Compiler | Unrecognizable or incorrect syntax. Semicolon missing on previous <br> statement. |
| :--- | :--- |
| create | Unrecognizable statement in command file, or numvar or outvar <br> statement error. |

G0009 Compiler pass out of memory
Compiler pass has run out of memory. Notify Aptech Systems.

G0010 Can't open output file

G0011 Compiled file must have correct extension
GAUSS requires a .gcg extension.

G0012 Invalid drive specifier

G0013 Invalid filename

G0014 File not found

G0015 Directory full

## G0016 Too many \#include's

\#include'd files are nested too deep.

## G0017 WARNING: local outside of procedure

A local statement has been found outside a procedure definition. The local statement will be ignored.

## G0018 Read error in program file

## G0019 Can't edit .gcg file

## G0020 Not implemented yet

Command not supported in this implementation.

## G0021 use must be at the beginning of a program

## G0022 User keyword cannot be used in expression

G0023 Illegal attempt to redefine symbol to an index variable

G0024 Invalid use of ->, probably should be .
G0025 Undefined symbol
A symbol has been referenced that has not been given a definition.

## G0026 Too many symbols

The global symbol table is full. (To set the limit, see new in the GAUSS Language Reference.)

## G0027 Invalid directory

## G0028 Can't open configuration file

GAUSS cannot find the configuration file.

## G0029 Missing left parenthesis

## G0030 Insufficient workspace memory

The space used to store and manipulate matrices and strings is not large enough for the operations attempted. (To make the main program space smaller and reclaim enough space to continue, see new in the GAUSS Language Reference.)

G0031 Execution stack too deep - expression too complex
An expression is too complex. Break it into smaller pieces. Notify Aptech Systems.

## G0032 fn function too large

## G0033 Missing right index bracket

## G0034 Missing arguments

## G0035 Argument too large

## G0036 Matrices are not conformable

For a description of the function or operator being used and conformability rules, see Matrix Operators, Section 11.2, or the GAUSS Language Reference.

## G0037 Result too large

The size of the result of an expression is greater than the limit for a single matrix.

G0038 Not all the eigenvalues can be computed

G0039 Matrix must be square to invert

G0040 Not all the singular values can be computed

## G0041 Argument must be scalar

A matrix argument was passed to a function that requires a scalar.

## G0042 Matrix must be square to compute determinant

## G0043 Not implemented for complex matrices

## G0044 Matrix must be real

G0045 Attempt to write complex data to real data set
Data sets, unlike matrices, cannot change from real to complex after they are created.
Use create complex to create a complex data set.

## G0046 Columns don't match

The matrices must have the same number of columns.

## G0047 Rows don't match

The matrices must have the same number of rows.

## G0048 Matrix singular

The matrix is singular using the current tolerance.

## G0049 Target matrix not complex

## G0050 Out of memory for program

The main program area is full. (To increase the main program space, see new in the GAUSS Language Reference.)

## G0051 Program too large

The main program area is full. (To increase the main program space, see new in the GAUSS Language Reference.)

## G0052 No square root - negative element

## G0053 Illegal index

An illegal value has been passed in as a matrix index.

## G0054 Index overflow

An illegal value has been passed in as a matrix index.

## G0055 retp outside of procedure

A retp statement has been encountered outside a procedure definition.

## G0056 Too many active locals

The execution stack is full. There are too many local variables active. Restructure your program. Notify Aptech Systems.

## G0057 Procedure stack overflow - expression too complex

The execution stack is full. There are too many nested levels of procedure calls. Restructure your program. Notify Aptech Systems.

## G0058 Index out of range

You have referenced a matrix element that is out of bounds for the matrix being referenced.

G0059 exec command string too long

G0060 Nonscalar index

G0061 Cholesky downdate failed

## G0062 Zero pivot encountered

crout The Crout algorithm has encountered a diagonal element equal to 0 . Use croutp instead.

## G0063 Operator missing

An expression contains two consecutive operands with no intervening operator.

## G0064 Operand missing

An expression contains two consecutive operators with no intervening operand.

## G0065 Division by zero!

## G0066 Must be recompiled under current version

You are attempting to use compiled code from a previous version of GAUSS. Recompile the source code under the current version.

## G0068 Program compiled under GAUSS-386 real version

## G0069 Program compiled under GAUSS-386i complex version

## G0070 Procedure calls too deep

You may have a runaway recursive procedure.

## G0071 Type mismatch

You are using an argument of the wrong data type (e.g., inputting a matrix when a string is called for).

## G0072 Too many files open

The limit on simultaneously open files is 10 .

## G0073 Redefinition of

| declare | An attempt has been made to initialize a variable that is already <br> initialized. This is an error when declare $:=$ is used. declare $!=$ <br> or declare $?=$ may be a better choice for your application. |
| :--- | :--- |

declare An attempt has been made to redefine a string as a matrix or procedure, or vice versa. delete the symbol and try again. If this happens in the context of a single program, you have a programming error. If this is a conflict between different programs, use a new statement before running the second program.
let A string is being forced to type matrix. Use an external matrix symbol; statement before the let statement.

## G0074 Can't run program compiled under GAUSS Light

## G0075 gscroll input vector the wrong size

## G0076 Call Aptech Systems Technical Support

## G0077 New size cannot be zero

You cannot reshape a matrix to a size of zero.
G0078 vargetl outside of procedure
G0079 varputl outside of procedure
G0080 File handle must be an integer
G0081 Error renaming file
G0082 Error reading file
G0083 Error creating temporary file
G0084 Too many locals
A procedure has too many local variables.

## G0085 Invalid file type

You cannot use this kind of file in this way.

## G0086 Error deleting file

## G0087 Couldn't open

The auxiliary output file could not be opened. Check the file name and make sure there is room on the disk.

## G0088 Not enough memory to convert the whole string

## G0089 WARNING: duplicate definition of local

## G0090 Label undefined

Label referenced has no definition.

## G0091 Symbol too long

Symbols can be no longer than 32 characters.

## G0092 Open comment

A comment was never closed.

G0093 Locate off screen

G0094 Argument out of range

G0095 Seed out of range

## G0096 Error parsing string

parse encountered a token that was too long.

## G0097 String not closed

A string must have double quotes at both ends.

## G0098 Invalid character for imaginary part of complex number

## G0099 Illegal redefinition of user keyword

## G0100 Internal E R R 0 R \#\#\#

Notify Aptech Systems.

## G0101 Argument cannot be zero

The argument to $\ln$ or $\mathbf{l o g}$ cannot be zero.

G0102 Subroutine calls too deep
Too many levels of gosub. Restructure your program.

G0103 return without gosub
You have encountered a subroutine without executing a gosub.

G0104 Argument must be positive

G0105 Bad expression or missing arguments
Check the expression in question, or you forgot an argument.

G0106 Factorial overflow

## G0107 Nesting too deep

Break the expression into smaller statements.

G0108 Missing left bracket [

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G0109 Not enough data items
You omitted data in a let statement
G0110 Found ) expected ] -
G0111 Found ] expected ) -
G0112 Matrix multiplication overflow
G0113 Unclosed (
G0114 Unclosed [
G0115 Illegal redefinition of functionYou are attempting to turn a function into a matrix or string. If this is a name conflict,delete the function.
G0116 sysstate: invalid case
G0117 Invalid argument
G0118 Argument must be integer
File handles must be integral.
G0120 Illegal type for save
G0121 Matrix not positive definite

The matrix is either not positive definite, or singular using the current tolerance.

## G0122 Bad file handle

The file handle does not refer to an open file or is not in the valid range for file handles.

## G0123 File handle not open

The file handle does not refer to an open file.
G0124 readr call too largeYou are attempting to read too much in one call.
G0125 Read past end of fileYou have already reached the end of the file.
G0126 Error closing file
G0127 File not open for write
G0128 File already open
G0129 File not open for read
G0130 No output variables specified
G0131 Can't create file, too many variables
G0132 Can't write, disk probably full
G0133 Function too long
G0134 Can't seekr in this type of file
G0135 Can't seek to negative row
G0136 Too many arguments or misplaced assignment operatorYou have an assignment operator (=) where you want a comparison operator (==), oryou have too many arguments.
G0137 Negative argument - erf or erfc
G0138 User keyword must have one argument
G0139 Negative parameter - Incomplete Beta
G0140 Invalid second parameter - Incomplete Beta
G0141 Invalid third parameter - Incomplete Beta
G0142 Nonpositive parameter - gamma
G0143 NaN or missing value - cdfchic
G0144 Negative parameter - cdfchic
G0145 Second parameter < 1.0 - cdfchic
G0146 Parameter too large - Incomplete Beta
G0147 Bad argument to trig function
G0148 Angle too large to trig function
G0149 Matrices not conformableFor a description of the function or operator being used and conformability rules, seeMatrix Operators, Section 11.2, or the GAUSS Language Reference.
G0150 Matrix not square
G0151 Sort failure

## G0152 Variable not initialized

You have referenced a variable that has not been initialized to any value.

G0153 Unsuccessful close on auxiliary output
The disk may be full.

## G0154 Illegal redefinition of string

## G0155 Nested procedure definition

A proc statement was encountered inside a procedure definition.

## G0156 Illegal redefinition of procedure

You are attempting to turn a procedure into a matrix or string. If this is a name conflict, delete the procedure.

## G0157 Illegal redefinition of matrix

G0158 endp without proc
You are attempting to end a procedure that you never started.

## G0159 Wrong number of parameters

You called a procedure with the wrong number of arguments.

## G0160 Expected string variable

## G0161 User keywords return nothing

## G0162 Can't save proc/keyword/fn with global references

Remove the global references or leave this in source code form for the autoloader to handle. (See library in the GAUSS Language Reference.)

## G0163 Wrong size format matrix

## G0164 Bad mask matrix

G0165 Type mismatch or missing arguments

## G0166 Character element too long

The maximum length for character elements is 8 characters.

G0167 Argument must be column vector

## G0168 Wrong number of returns

The procedure was defined to return a different number of items.

## G0169 Invalid pointer

You are attempting to call a local procedure using an invalid procedure pointer.

G0170 Invalid use of ampersand

## G0171 Called symbol is wrong type

You are attempting to call a local procedure using a pointer to something else.

G0172 Can't resize temporary file

G0173 varindx failed during open
The global symbol table is full.

G0174 '،.'’ and "، '’ operators must be inside [ ] brackets These operators are for indexing matrices.
G0175 String too long to compare
G0176 Argument out of range
G0177 Invalid format string
G0178 Invalid mode for getf
G0179 Insufficient heap space
G0180 Trim too much
You are attempting to trim more rows than the matrix has.
G0181 Illegal assignment - type mismatch
G0182 2nd and 3rd arguments different order
G0274 Invalid parameter for conv
G0275 Parameter is NaN (Not A Number)The argument is a NaN (see Special Data Types, Section 10.6.9).
G0276 Illegal use of reserved word
G0277 Null string illegal here
G0278 proc without endp
You must terminate a procedure definition with an endp statement.
G0286 Multiple assign out of memory
G0287 Seed not updated

The seed argument to rndns and rndus must be a simple local or global variable reference. It cannot be an expression or constant. These functions are obsolete, please use rndlen and rndlcu

## G0288 Found break not in do loop

## G0289 Found continue not in do loop

## G0290 Library not found

The specified library cannot be found on the lib_path path. Make sure installation was correct.

G0291 Compiler pass out of memory
Notify Aptech Systems.

G0292 File listed in library not found
A file listed in a library could not be opened.

## G0293 Procedure has no definition

The procedure was not initialized. Define it.

## G0294 Error opening temporary file

One of the temporary files could not be opened. The directory may be full.

## G0295 Error writing temporary file

One of the temporary files could not be written to. The disk may be full.

## G0296 Can't raise negative number to nonintegral power

## G0300 File handle must be a scalar

## G0301 Syntax error in library

G0302 File has been truncated or corrupted
getname File header cannot be read.
load Cannot read input file, or file header cannot be read.open $\quad$ File size does not match header specifications, or file header cannotbe read.
G0317 Can't open temp file
G0336 Disk full
G0339 Can't debug compiled program
G0341 File too big
G0347 Can't allocate that many globals
G0351 Warning: Not reinitializing : declare ?=The symbol is already initialized. It will be left as is.
G0352 Warning: Reinitializing : declare != The symbol is already initialized. It will be reset.
G0355 Wrong size line matrix
G0360 Write error
G0364 Paging error
G0365 Unsupported executable file type

G0368 Unable to allocate translation space
G0369 Unable to allocate buffer

G0370 Syntax Error in code statement

G0371 Syntax Error in recode statement

G0372 Token verify error
Notify Aptech Systems.

G0373 Procedure definition not allowed
A procedure name appears on the left side of an assignment operator.
G0374 Invalid make statement

G0375 make Variable is a Number

G0376 make Variable is Procedure

G0377 Cannot make Existing Variable

G0378 Cannot make External Variable

G0379 Cannot make String Constant

G0380 Invalid vector statement

G0381 vector Variable is a Number

G0382 vector Variable is Procedure
G0383 Cannot vector Existing Variable
G0384 Cannot vector External Variable
G0385 Cannot vector String Constant
G0386 Invalid extern statement
G0387 Cannot extern number
G0388 Procedures always external
A procedure name has been declared in an extern statement. This is a warning only.
G0389 extern variable already localA variable declared in an extern statement has already been assigned local status.
G0390 String constant cannot be external
G0391 Invalid code statement
G0392 code Variable is a Number
G0393 code Variable is Procedure
G0394 Cannot code Existing Variable
G0395 Cannot code External Variable
G0396 Cannot code String Constant
G0397 Invalid recode statement
G0398 recode Variable is a Number
G0399 recode Variable is Procedure
G0400 Cannot recode External Variable
G0401 Cannot recode String Constant
G0402 Invalid keep statement
G0403 Invalid drop statement
G0404 Cannot define Number
G0405 Cannot define String
G0406 Invalid select statement
G0407 Invalid delete statement
G0408 Invalid outtyp statement
G0409 outtyp already defaulted to 8Character data has been found in the output data set before an outtyp 2 or outtyp 4 statement. This is a warning only.
G0410 outtyp must equal 2,4 , or 8
G0411 outtyp override...precision set to 8Character data has been found in the output data set after an outtyp 2 or outtyp 4statement. This is a warning only.
G0412 default not allowed in recode statement default allowed only in code statement.
G0413 Missing file name in dataloop statement
G0414 Invalid listwise statement
G0415 Invalid lag statement
G0416 lag variable is a number
G0417 lag variable is a procedure
G0418 Cannot lag External Variable
G0419 Cannot lag String Constant
G0421 Command not supported in Run-Time Module
G0428 Cannot use debug command inside program
G0429 Invalid number of subdiagonals
G0431 Error closing dynamic library
G0432 Error opening dynamic library
G0433 Cannot find DLL function
G0434 Error opening default dynamic library
G0435 Invalid mode
G0436 Matrix is empty
G0437 loadexe not supported; use dlibrary instead
G0438 callexe not supported; use dllcall instead
G0439 File has wrong bit order
G0440 File has wrong byte order
G0441 Type vector malloc failed
G0442 No type vector in gfblock
G0445 Illegal left-hand side reference in procedure
G0446 Argument is the wrong size
G0447 vfor called with illegal loop level
G0454 Failure opening printer for output
G0456 Failure buffering output for printerG0457 Cannot take log of a negative number
G0458 Attempt to index proc/fn/keyword as a matrix
G0459 Missing right brace
G0460 Unexpected end of statement
G0461 Too many data items
G0462 Negative trim value
G0463 Failure generating graph
G0465 Redefinition of structure, number of elements
G0466 Redefinition of structure, type mismatch
G0467 Redefinition of structure, unrecognized member
G0468 Structure definition inside procedure definition
G0469 Cannot create translator temp file
G0470 Symbol not found
G0472 Invalid name
G0473 String not terminated with null byte
G0477 FOR loops nested too deep
G0486 Character argument too long
G0487 License expired
G0490 License manager initialization error
G0491 License manager error
G0492 Licensing failure
G0497 Missing right parenthesis
G0500 Cannot create temporary filename
G0503 Cannot assign matrix to scalar member
G0504 Invalid structure member
G0505 Invalid structure redefinition
G0506 Structure assignment mismatch
G0507 Undefined structure
G0508 Structure argument mismatch
G0509 Too many structure members
G0510 Duplicate name for structure member
G0514 Not supported for structures
G0515 Too many values in locator
G0516 Too many dimensions in result
G0517 Too many dimensions in argument
G0518 Not implemented for complex

## G0519 Illegal dereference of structure array

G0520 Arguments not conformable

G0521 Argument must be real

G0522 Illegal indexing of dereferenced structure

G0523 Numeric argument must be integer

G0524 Found comma, expecting index

G0525 Argument contains NaNs

G0526 Argument must be compact format

G0529 Array orders must be >= 1

G0531 Two trailing dimensions of argument must be the same size

G0532 Both dimensions of argument must be the same size

G0533 1-dimensional argument must contain only 1 element

G0534 Cannot create file

G0538 Zero illegal in for loop increment

G0541 Illegal assignment to FOR loop counter

G0542 Object too large for 32 -bit version
G0543 Array has too many dimensions for matrix assign
G0547 Array not conformable for indexing
G0548 Array not conformable for boolean operation
G0549 Global structure pointer cannot point to local structure
G0550 Invalid use of *
G0551 Feature not authorized
G0553 Path too long


## Maximizing Performance

These hints will help you maximize the performance of your new GAUSS System.

### 30.1 Library System

Some temporary files are created during the autoloading process. If you have a tmp_path configuration variable or a tmp environment string that defines a path on a RAM disk, the temporary files will be placed on the RAM disk.

For example:

```
set tmp=f:\tmp
```

tmp_path takes precedence over the tmp environment variable.
A disk cache will also help, as well as having your frequently used files in the first path in the src_path.

You can optimize your library . lcg files by putting the correct drive and path on each file name listed in the library. The lib command will do this for you.

Use the compile command to precompile your large frequently used programs. This will completely eliminate compile time when the programs are rerun.

### 30.2 Loops

The use of the built-in matrix operators and functions rather than do loops will ensure that you are utilizing the potential of GAUSS.

Here is an example:
Given the vector $\mathbf{x}$ with 8000 normal random numbers,

```
x = rndn(8000,1);
```

you could get a count of the elements with an absolute value greater than 1 with a do loop, like this:

```
c = 0;
i = 1;
do while i <= rows(x);
    if abs(x[i]) > 1;
        c = c+1;
    endif;
    i = i+1;
endo;
print c;
```

Or, you could use:

```
c = sumc(abs(x) .> 1);
print c;
```

The do loop takes over 40 times longer.

### 30.3 Memory Usage

Computers today can have large amounts of RAM. This doesn't mean that large data sets should be read entirely into memory. Many GAUSS procedures and applications are written to allow for data sets to be read in sections rather than all at once. Even if you have enough RAM to store the data set completely, you should consider taking advantage of this feature. The speed-ups using this feature can be significant. For example, ols is called using a data set stored in a matrix versus stored on the disk in a GAUSS data set. The computer is a 2.8 Megahertz computer with Windows XP.

```
y = rndn(250000,1);
x = rndn(250000,100);
xlbl = 0$+"X"+ftocv(seqa(1,1,100),1,0);
lbl = "Y" | xlbl;
call saved(y`x,"test",lbl);
__output = 0;
t0 = date;
call ols("",y,x);
t1 = date;
t2 = date;
call ols("test","Y",xlbl);
t3 = date;
print ethsec(t2,t3)/100 " seconds;
print;
print ethsec(t0,t1)/100 " seconds";
```

25.750000 seconds
9.6720000 seconds

This represents more than a $50 \%$ speedup by leaving the data on the disk.

## maxvec,maxbytes

maxvec is a GAUSS procedure that returns the value of the global variable __maxvec that determines the amount of data to be read in at a time from a GAUSS data set. This value can be modified for a particular run by setting __maxvec in your command file to some other value. The value returned by a call to maxvec can be permanently modified by editing system. dec and changing the value of __maxvec. The value returned when running GAUSS Light is always 8192.
maxbytes is a GAUSS procedure that returns the value of a scalar global __maxbytes that sets the amount of available RAM. This value can be modified for a particular run by setting __maxbytes in your command file to some other value. The value returned by a call to maxbytes can be permanently modified by editing system. dec and changing the value of __maxbytes.

If you wish to force GAUSS procedures and applications to read a GAUSS data set in its entirety, set __maxvec and __maxbytes to very large values.

### 30.3.1 Hard Disk Maintenance

The hard disk used for the swap file should be optimized occasionally with a disk optimizer. Use a disk maintenance program to ensure that the disk media is in good shape.

### 30.3.2 CPU Cache

There is a line for cache size in the gauss.cfg file. Set it to the size of the CPU data cache for your computer.

This affects the choice of algorithms used for matrix multiply functions.
This will not change the results you get, but it can radically affect performance for large matrices.

## Fonts

There are four fonts available in the Publication Quality Graphics System:

| Simplex | standard sans serif font |
| :--- | :--- |
| Simgrma | Simplex greek, math |
| Microb | bold and boxy |
| complex | standard font with serif |

The following tables show the characters available in each font and their ASCII values. (For details on selecting fonts for your graph, see Selecting Fonts, Section 25.4.1.

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## A. 1 Simplex

| 33 | ! | 61 | $=$ | 89 | Y | 117 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 34 | " | 62 | $>$ | 90 | Z | 118 |
| 35 | \# | 63 | ? | 91 | [ | 119 |
| 36 | \$ | 64 | @ | 92 | $\lambda$ | 120 |
| 37 | $\%$ | 65 | A | 93 | ] | 121 |
| 38 | \& | 66 | B | 94 | - | 122 |
| 39 | , | 67 | C | 95 | - | 123 |
| 40 | ( | 68 | D | 96 |  | 124 |
| 41 | ) | 69 | E | 97 | a | 125 |
| 42 | * | 70 | F | 98 | b | 126 |
| 43 | + | 71 | G | 99 | c |  |
| 44 | , | 72 | H | 100 | d |  |
| 45 | - | 73 | I | 101 | e |  |
| 46 | . | 74 | $\checkmark$ | 102 | f |  |
| 47 | / | 75 | K | 103 | g |  |
| 48 | 0 | 76 | L | 104 | h |  |
| 49 | 1 | 77 | M | 105 | i |  |
| 50 | 2 | 78 | N | 106 | j |  |
| 51 | 3 | 79 | 0 | 107 | k |  |
| 52 | 4 | 80 | P | 108 | 1 |  |
| 53 | 5 | 81 | Q | 109 | m |  |
| 54 | 6 | 82 | R | 110 | n |  |
| 55 | 7 | 83 | S | 111 | - |  |
| 56 | 8 | 84 | T | 112 | p |  |
| 57 | 9 | 85 | U | 113 | q |  |
| 58 | : | 86 | V | 114 | $r$ |  |
| 59 | ; | 87 | W | 115 | s |  |
| 60 | < | 88 | X | 116 | t |  |

## A. 2 Simgrma

| 33 | $\epsilon$ | 61 | $\neq$ | 89 | $\psi$ | 117 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 34 | ( | 62 | $\geqq$ | 90 | z | 118 |
| 35 | $\equiv$ | 63 | $\simeq$ | 91 | [ | 119 |
| 36 | $\approx$ | 64 | $\cup$ | 92 | a | 120 |
| 37 | $\uparrow$ | 65 | $\frac{1}{2}$ | 93 | ] | 121 |
| 38 | $\checkmark$ | 66 | $\frac{1}{3}$ | 94 | $\bigcirc$ | 122 |
| 39 | , | 67 | H | 95 | $\downarrow$ | 123 |
| 40 | c | 68 | $\triangle$ | 96 | " | 124 |
| 41 | $\supset$ | 69 | $\frac{1}{8}$ | 97 | $\alpha$ | 125 |
| 42 | $\times$ | 70 | $\phi$ | 98 | $\beta$ | 126 |
| 43 | $\pm$ | 71 | $\Gamma$ | 99 | $\eta$ |  |
| 44 | $\int$ | 72 | X | 100 | $\delta$ |  |
| 45 | 干 | 73 | $\frac{2}{3}$ | 101 | $\varepsilon$ |  |
| 46 | - | 74 | $\perp$ | 102 | $\varphi$ |  |
| 47 | $\div$ | 75 | $\frac{3}{8}$ | 103 | $\gamma$ |  |
| 48 | $\nabla$ | 76 | $\wedge$ | 104 | $\chi$ |  |
| 49 | $\checkmark$ | 77 | $\frac{5}{8}$ | 105 | $\iota$ |  |
| 50 | $\oint$ | 78 | $\frac{7}{8}$ | 106 | $t$ |  |
| 51 |  | 79 | $\frac{1}{4}$ | 107 | $\kappa$ |  |
| 52 |  | 80 | $\square$ | 108 | $\lambda$ |  |
| 53 |  | 81 | $\bigcirc$ | 109 | $\mu$ |  |
| 54 | $\exists$ | 82 | P | 110 | $\nu$ |  |
| 55 | \\| | 83 | $\Sigma$ | 111 | $\bigcirc$ |  |
| 56 | $\infty$ | 84 | $\lesssim$ | 112 | $\pi$ |  |
| 57 | $\odot$ | 85 | $r$ | 113 | $v$ |  |
| 58 | $\rightarrow$ | 86 | $\leftrightarrow$ | 114 | $\rho$ |  |
| 59 | $\leftarrow$ | 87 | $\Omega$ | 115 | $\sigma$ |  |
| 60 | $\leqq$ | 88 | 三 | 116 | $\tau$ |  |

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## A. 3 Microb

| 33 | $!$ | 61 | $=$ | 89 | Y | 117 | $u$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 34 | " | 62 | > | 90 | Z | 118 | $\checkmark$ |
| 35 | \# | 63 | ? | 91 | [ | 119 | $w$ |
| 36 | \$ | 64 | @ | 92 | $\backslash$ | 120 | $\times$ |
| 37 | \% | 65 | A | 93 | 1 | 121 | y |
| 38 | \& | 66 | B | 94 | - | 122 | z |
| 39 | ' | 67 | C | 95 | - | 123 | \{ |
| 40 | ¢ | 68 | D | 96 | - | 124 | 1 |
| 41 | ] | 69 | E | 97 | a | 125 | \} |
| 42 | * | 70 | F | 98 | b | 126 | $\sim$ |
| 43 | + | 71 | G | 99 | c |  |  |
| 44 | , | 72 | H | 100 | d |  |  |
| 45 | - | 73 | 1 | 101 | e |  |  |
| 46 | . | 74 | J | 102 | $f$ |  |  |
| 47 | 1 | 75 | K | 103 | g |  |  |
| 48 | 0 | 76 | L | 104 | h |  |  |
| 49 | 1 | 77 | M | 105 | 1 |  |  |
| 50 | 2 | 78 | N | 106 | j |  |  |
| 51 | 3 | 79 | 0 | 107 | k |  |  |
| 52 | 4 | 80 | P | 108 | । |  |  |
| 53 | 5 | 81 | Q | 109 | m |  |  |
| 54 | 6 | 82 | R | 110 | n |  |  |
| 55 | 7 | 83 | S | 111 | $\bigcirc$ |  |  |
| 56 | 8 | 84 | T | 112 | p |  |  |
| 57 | 9 | 85 | U | 113 | q |  |  |
| 58 | : | 86 | v | 114 | $r$ |  |  |
| 59 | ; | 87 | W | 115 | s |  |  |
| 60 | $<$ | 88 | X | 116 | t |  |  |

## A. 4 Complex

| 33 | ! | 61 | $=$ | 89 | Y | 117 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 34 | " | 62 | > | 90 | Z | 118 |
| 35 | \# | 63 | ? | 91 | [ | 119 |
| 36 | \$ | 64 | @ | 92 | $\backslash$ | 120 |
| 37 | \% | 65 | A | 93 | ] | 121 |
| 38 | \& | 66 | B | 94 | - | 122 |
| 39 | , | 67 | C | 95 | - | 123 |
| 40 | ( | 68 | D | 96 |  | 124 |
| 41 | ) | 69 | E | 97 | a | 125 |
| 42 | * | 70 | F | 98 | b | 126 |
| 43 | + | 71 | G | 99 | c |  |
| 44 | , | 72 | H | 100 | d |  |
| 45 | - | 73 | I | 101 | e |  |
| 46 | . | 74 | J | 102 | f |  |
| 47 | / | 75 | K | 103 | g |  |
| 48 | 0 | 76 | L | 104 | h |  |
| 49 | 1 | 77 | M | 105 | , |  |
| 50 | 2 | 78 | N | 106 | j |  |
| 51 | 3 | 79 | 0 | 107 | k |  |
| 52 | 4 | 80 | P | 108 | 1 |  |
| 53 | 5 | 81 | Q | 109 | m |  |
| 54 | 6 | 82 | R | 110 | n |  |
| 55 | 7 | 83 | S | 111 | - |  |
| 56 | 8 | 84 | T | 112 | p |  |
| 57 | 9 | 85 | U | 113 | q |  |
| 58 | : | 86 | V | 114 | r |  |
| 59 | ; | 87 | W | 115 | s |  |
| 60 | $<$ | 88 | X | 116 | t |  |



## Reserved Words Appendix

The following words are used for GAUSS functions. You cannot use these names for variables or procedures in your programs:

A
abs
acf
aconcat
acos
aeye
amax
amean
AmericanBinomCall
AmericanBinomCall_Greeks
AmericanBinomCall_ImpVol
AmericanBinomPut
AmericanBinomPut_Greeks
AmericanBinomPut_ImpVol

## AmericanBSCall

AmericanBSCall_Greeks
AmericanBSCall_ImpVol
AmericanBSPut
AmericanBSPut_Greeks
AmericanBSPut_ImpVol
amin
amult
and
annualTradingDays
arccos
arcsin
arctan
$\arctan 2$
areshape
arrayalloc
arrayindex
arrayinit
arraytomat
asclabel
asin
asum
atan
atan2
atranspose
axmargin

B
balance
base10
band
bandchol
begwind
bandcholsol
besselj
bandltsol
bessely
bandrv
bandsolpd
box
bar
boxcox
break

C
calcbox
call
callexe
cdfbeta
cdfbvn
cdfbvn2
cdfbvn2e
cdfchic
cdfchii
cdfchinc
cdfn
cdffc
cdfffnc
cdfgam
cdfmvn
cdfn2
cdfnc
cdfni
cdftc
cdftci
cdftnc
cdftvn
cdir
ceil
cfft
cffti
changedir
chdir
checkinterrupt
chol
choldn
cholsol
cholup
chrs
cint
clear
clearg
close
closeall
cls
cmsplit
cmsplit2
code
color
cols
colsf
combinate
combinated
comlog
commandeerm
commandeersa
compile
complex
con
cond
conformed
conj
cons
continue
contour
conv
convertsatostr
convertstrtosa
coreleft
corrm
corrms
corrvc
corrx
corrxs
cos
cosh
counts
countwts
create
crossprd
crout
croutp
csrcol
csrlin
csrtype
cumprode
cumsumc
curve
cvtos
cvtosa
dataopen
datasave
date
datestr
datestring
datestrymd
dayinyr
day0fWeek
debug
declare
delete
deletefile
delif
denseSubmat
design
det
detl
dfft
dffti
dfree
diag
diagrv
digamma
disable
dlibrary
dllcall
do
dos
doswincloseall
doswinopen
dotfeq
dotfeqmt
dotfge
dotfgemt
dotfgt
dotfgtmt
dotfle
dotflemt
dotflt
dotfltmt
dotfne
dotfnemt
draw
dsCreate
dstat
dstatmt
dstatmtControlCreate
dtdate
dtday
dttime
dttodtv
dttostr
dttoutc
dtvnormal
dtvtodt
dtvtoutc
dummy
dummybr
dummydn
ed
eigcg2
edit
editm
eig
eigcg
eigch
eigch2
eigh
eighv

| eigrg | erfc |
| :--- | :--- |
| eigrg2 | error |
| eigrs | errorlog |
| eigrs2 | etdays |
| eigv | ethsec |
| elapsedTradingDays | etstr |
| else | EuropeanBinomCall |
| elseif | EuropeanBinomCall_Greeks |
| enable | EuropeanBinomCall_ImpVol |
| end | EuropeanBinomPut |
| endfor | EuropeanBinomPut_Greeks |
| endif | EuropeanBinomPut_ImpVol |
| endo | EuropeanBSCall |
| endp | EuropeanBSCall_Greeks |
| endwind | EuropeanBSCall_ImpVol |
| envget | EuropeanBSPut |
| eof | EuropeanBSPut_Greeks |
| eq | EuropeanBSPut_ImpVol |
| eqSolve | exctsmpl |
| eqSolvemt | exec |
| eqSolvemtControlCreate | execbg |
| eqSolvemtOutCreate | exp |
| eqSolveSet | expr |
| eqv | external |
| erf | eye |

## F

fcheckerr
fclearerr
feq
feqmt
fflush
fft
ffti
fftm
fftmi
fftn
fge
fgemt
fgets
fgetsa
fgetsat
fgetst
fgt
fgtmt
fileinfo
files
filesa
fix
fle
flemt
floor
flt
fltmt
fmod
fn
fne
fnemt
font
fontload
fonts
fontunload
fontunloadall
fopen
for
format
formatcv
formatnv
fputs
fputst
fseek
fstrerror
ftell
ftocv
ftos
ftostrc

G
gamma
gammaii
gausset
gdaappend
gdacreate
gdadstat
gdadstatmat
gdagetindex
gdagetname
gdagetnames
gdagetorders
gdagettype
gdagettypes
gdagetvarinfo
gdaiscplx
gdapack
gdaread
gdareadbyindex
gdareadsome
gdareportvarinfo
gdaupdate
gdaupdateandpack
gdawrite
gdawritesome
gdtfastcat
ge
getarray
getdims
getf
getmatrix
getmatrix4d
getname
getnamef gosub
getNextTradingDay ..... goto
getNextWeekDay gradMTgetnr
getnrmt
getorders
getpathgetPreviousTradingDaygetPreviousWeekDaygetscalar3dgetscalar4dgetwind
HhardcopyhessMTmhasimagheaderheadermthesshessMThessMTghessMTgw
gradMTm
gradp
graph
graphgpg
graphinit
graphprt
graphset
graphsev3gt
hessMTmw
hessMTw
hessp
hist histf histp hsec

## if

imag
indcv
indexcat
indices
indices2
indicesf
indicesfn
indnv
indsav
int
intgrat2
intgrat3
inthp
intHP1
intHP2
intHP3
intHP4
inthpControlCreate
intquad1
intquad2
intquad3
intrleav
intrleavsa
intrsect
intrsectsaintrsectsa
intsimp
inv
invpd
invswp
iscplx
iscplxf
isinfnanmiss
ismiss
isSparse
K
key
keyw
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## Singularity Tolerance Appendix

The tolerance used to determine whether or not a matrix is singular can be changed. The default value is $1.0 \mathrm{e}-14$ for both the LU and the Cholesky decompositions. The tolerance for each decomposition can be changed separately. The following operators are affected by a change in the tolerance:

## Crout LU Decomposition

```
crout(x)
croutp(x)
inv(x)
det(x)
y/x when neither }x\mathrm{ nor }y\mathrm{ is scalar and }x\mathrm{ is square.
```


## Cholesky Decomposition

```
chol(x)
invpd(x)
solpd(y,x)
```

$y / x \quad$ when neither $x$ nor $y$ is scalar and $x$ is not square.

## C. 1 Reading and Setting the Tolerance

The tolerance value may be read or set using the sysstate function, cases 13 and 14 .

## C. 2 Determining Singularity

There is no perfect tolerance for determining singularity. The default is $1.0 \mathrm{e}-14$. You can adjust this as necessary.

A numerically better method of determining singularity is to use cond to determine the condition number of the matrix. If the equation

$$
1 / \operatorname{cond}(x)+1 \text { eq } 1
$$

is true, then the matrix is usually considered singular to machine precision. (See LINPACK for a detailed discussion on the relationship between the matrix condition and the number of significant figures of accuracy to be expected in the result.)

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