

# SAS/STAT® 13.2 User's Guide Shared Concepts and Topics



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# Chapter 19 Shared Concepts and Topics

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This chapter introduces a number of concepts that are common to two or more SAS/STAT procedures. Most sections display a listing of the procedures for which the shared topic is relevant.

# **Levelization of Classification Variables**

A classification variable is a variable that enters the statistical analysis or model not through its values, but through its levels. The process of associating values of a variable with levels is termed *levelization*.

This section covers in particular procedures that support a CLASS statement for specifying classification variables. Some of the concepts discussed also apply to procedures that use different syntax to request levelization of variables (for example, the CLASS() transformation in the TRANSREG procedure).

During the process of levelization, observations that share the same value are assigned to the same level. The manner in which values are grouped can be affected by the inclusion of formats. The sort order of the levels can be determined with the ORDER= option in the procedure statement. With the GENMOD, GLMSELECT, and LOGISTIC procedures, you can also control the sort order separately for each variable in the CLASS statement.

Consider the data on nine observations in Table 19.1. The variable A is integer valued, and the variable X is a continuous variable with a missing value for the fourth observations. The fourth and fifth columns of Table 19.1 apply two different formats to the variable X.

<b>Table 19.1</b>	Example D	ata for	Levelization
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Obs	A	X	FORMAT x 3.0	FORMAT x 3.1
1	2	1.09	1	1.1
2	2	1.13	1	1.1
3	2	1.27	1	1.3
4	3			
5	3	2.26	2	2.3
6	3	2.48	2	2.5
7	4	3.34	3	3.3
8	4	3.34	3	3.3
9	4	3.14	3	3.1

By default, levelization of the variables groups observations by the formatted value of the variable, except for numerical variables for which no explicit format is provided. Numerical variables for which no explicit format is provided are sorted by their internal value. The levelization of the four columns in table Table 19.1 leads to the level assignment in Table 19.2.

**Table 19.2** Values and Levels

Obs	A Obs Value Level		Value	X Value Level		RMAT x 3.0 ue Level	FORMAT x 3.1 Value Level		
1	2	1	1.09	1	1	1	1.1	1	
2	2	1	1.13	2	1	1	1.1	1	
3	2	1	1.27	3	1	1	1.3	2	
4	3	2							
5	3	2	2.26	4	2	2	2.3	3	
6	3	2	2.48	5	2	2	2.5	4	
7	4	3	3.34	7	3	3	3.3	6	
8	4	3	3.34	7	3	3	3.3	6	
9	4	3	3.14	6	3	3	3.1	5	

The ORDER= option in the PROC statement specifies the sort order for the levels of CLASS variables. When ORDER=FORMATTED (which is the default) is in effect for numeric variables for which you have supplied no explicit format, the levels are ordered by their internal values. To order numeric class levels with no explicit format by their BEST12. formatted values, you can specify the BEST12. format explicitly for the CLASS variables.

The following table shows how values of the ORDER= option are interpreted.

Value of ORDER=	Levels Sorted By
DATA	Order of appearance in the input data set
FORMATTED	External formatted value, except for numeric variables with no explicit format, which are sorted by their unformatted (internal) value
FREQ	Descending frequency count; levels with the most observa- tions come first in the order
INTERNAL	Unformatted value

For FORMATTED and INTERNAL values, the sort order is machine dependent. For more information about sort order, see the chapter on the SORT procedure in the Base SAS Procedures Guide and the discussion of BY-group processing in SAS Language Reference: Concepts.

The GLMSELECT, LOGISTIC, and GENMOD procedures support a MISSING option in the CLASS statement. When this option is in effect, missing values (. for a numeric variable and blanks for a character variable) are included in the levelization and are assigned a level. Table 19.3 displays the results of levelizing the values in Table 19.1 when the MISSING option is in effect.

Obs	A S Value Level		X Value Level			RMAT x 3.0 ue Level	FORMAT x 3.1 Value Level	
1	2	1	1.09	2	1	2	1.1	2
2	2	1	1.13	3	1	2	1.1	2
3	2	1	1.27	4	1	2	1.3	3
4	3	2		1	•	1		1
5	3	2	2.26	5	2	3	2.3	4
6	3	2	2.48	6	2	3	2.5	5
7	4	3	3.34	8	3	4	3.3	7
8	4	3	3.34	8	3	4	3.3	7
9	4	3	3.14	7	3	4	3.1	6

Table 19.3 Values and Levels with MISSING Option

When the MISSING option is not specified, or for procedures whose CLASS statement does not support this option, it is important to understand the implications of missing values for your statistical analysis. When a SAS/STAT procedure levelizes the CLASS variables, an observation for which a CLASS variable has a missing value is excluded from the analysis. This is true regardless of whether the variable is used to form the statistical model. Consider, for example, the case where some observations contain missing values for variable A but the records for these observations are otherwise complete with respect to all other variables in the statistical models. The analysis results from the following statements do not include any observations for which variable A contains missing values, even though A is not specified in the MODEL statement:

```
class A B;
model y = B \times B*x;
```

Many statistical procedures print a "Number of Observations" table that shows the number of observations read from the data set and the number of observations used in the analysis. Pay careful attention to this table—especially when your data set contains missing values—to ensure that no observations are unintentionally excluded from the analysis.

# **Parameterization of Model Effects**

The general form of a linear regression model is defined in Chapter 3, "Regression Models and Models with Classification Effects" as

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\epsilon}$$

This section describes how matrices of regressor effects such as **X** are constructed in SAS/STAT software. These constructions (*parameterization* rules) apply to regression models, models with classification effects, generalized linear models, and mixed models. The simplest and most general parameterization rules are the ones used in the GLM procedure, and they are discussed first. Several procedures also support alternate parameterizations of classification variables, including the CATMOD, GENMOD, GLMSELECT, LOGISTIC, PHREG, SURVEYLOGISTIC, and SURVEYPHREG procedures. These are discussed after the GLM parameterization of classification variables and model effects.

All modeling procedures that have a CLASS statement support classification variables and effects, and those procedures that additionally support the supplemental parameterizations have a PARAM= option in the CLASS statement.

# **GLM Parameterization of Classification Variables and Effects**

This section applies to the following procedures:

GAM, GENMOD, GLIMMIX, GLM, GLMPOWER, GLMSELECT, LIFEREG, LOGISTIC, MI, MIXED, MULLTEST, ORTHOREG, PHREG, PLS, QUANTREG, ROBUSTREG, SURVEYLOGISTIC, and SURVEYPHREG.

# Intercept

By default, SAS/STAT linear models automatically include a column of 1s in **X** which corresponds to an intercept parameter. In many procedures you can use the NOINT option in the MODEL statement to suppress this intercept. For example, the NOINT option is useful when the MODEL statement contains a classification effect and you want the parameter estimates to be in terms of the mean response for each level of that effect.

# **Regression Effects**

Numeric variables or polynomial terms that involve them can be included in the model as regression effects (covariates). The actual values of such terms are included as columns of the relevant model matrices. You can use the bar operator with a regression effect to generate polynomial effects. For example, XIXIX expands to X X\*X X\*X\*X, which is a cubic model.

# **Main Effects**

If a classification variable has m levels, the GLM parameterization generates m columns for its main effect in the model matrix. Each column is an indicator variable for a given level. The order of the columns is the sort order of the values of their levels and frequently can be controlled with the ORDER= option in the procedure or CLASS statement.

Table 19.4 is an example where  $\beta_0$  denotes the intercept and A and B are classification variables with two and three levels, respectively.

	Data	I	,	Ą	В
Α	В	$eta_0$	A1	A2	B1 B2 B3
1	1	1	1	0	1 0 0
1	2	1	1	0	0 1 0
1	3	1	1	0	0 0 1
2	1	1	0	1	1 0 0
2	2	1	0	1	0 1 0
2	3	1	0	1	0 0 1

**Table 19.4** Example of Main Effects

Typically, there are more columns for these effects than there are degrees of freedom to estimate them. In other words, the GLM parameterization of main effects is singular.

# Interaction Effects

Often a model includes interaction (crossed) effects to account for how the effect of a variable changes with the values of other variables. With an interaction, the terms are first reordered to correspond to the order of the variables in the CLASS statement. Thus, B\*A becomes A\*B if A precedes B in the CLASS statement. Then, the GLM parameterization generates columns for all combinations of levels that occur in the data. The order of the columns is such that the rightmost variables in the interaction change faster than the leftmost variables (Table 19.5). In the MIXED and GLIMMIX procedures, which support both fixedand random-effects models, empty columns (that is, columns that would contain all 0s) are not generated for fixed effects, but they are generated for random effects.

Da	ata	I	,	4		В				A;	*B		
Α	В	$\beta_0$	A1	A2	B1	B2	В3	A1B1	A1B2	A1B3	A2B1	A2B2	A2B3
1	1	1	1	0	1	0	0	1	0	0	0	0	0
1	2	1	1	0	0	1	0	0	1	0	0	0	0
1	3	1	1	0	0	0	1	0	0	1	0	0	0
2	1	1	0	1	1	0	0	0	0	0	1	0	0
2	2	1	0	1	0	1	0	0	0	0	0	1	0
2	3	1	0	1	0	0	1	0	0	0	0	0	1

Table 19.5 Example of Interaction Effects

In the preceding matrix, main-effects columns are not linearly independent of crossed-effects columns; in fact, the column space for the crossed effects contains the space of the main effect.

When your model contains many interaction effects, you might be able to code them more parsimoniously by using the bar operator (1). The bar operator generates all possible interaction effects. For example, AIBIC expands to A B A\*B C A\*C B\*C A\*B\*C. To eliminate higher-order interaction effects, use the at sign (@) in conjunction with the bar operator. For instance, AlBICID@2 expands to A B A\*B C A\*C B\*C D A\*D B\*D C\*D.

# **Nested Effects**

Nested effects are generated in the same manner as crossed effects. Hence, the design columns generated by the following two statements are the same (but the ordering of the columns is different):

model Y=A B(A); model Y=A A\*B;

The nesting operator in SAS/STAT software is more of a notational convenience than an operation distinct from crossing. Nested effects are typically characterized by the property that the nested variables never appear as main effects. The order of the variables within nesting parentheses is made to correspond to the order of these variables in the CLASS statement. The order of the columns is such that variables outside the parentheses index faster than those inside the parentheses, and the rightmost nested variables index faster than the leftmost variables (Table 19.6).

	Data	I	,	4			В	(A)		
Α	В	$\beta_0$	A1	A2	B1A1	B2A1	B3A1	B1A2	B2A2	B3A2
1	1	1	1	0	1	0	0	0	0	0
1	2	1	1	0	0	1	0	0	0	0
1	3	1	1	0	0	0	1	0	0	0
2	1	1	0	1	0	0	0	1	0	0
2	2	1	0	1	0	0	0	0	1	0
2	3	1	0	1	0	0	0	0	0	1

**Table 19.6** Example of Nested Effects

# **Continuous-Nesting-Class Effects**

When a continuous variable nests or crosses with a classification variable, the design columns are constructed by multiplying the continuous values into the design columns for the classification effect (Table 19.7).

Data		I	I A		X(A)		
Х	Α	$eta_0$	A1	A2	X(A1)	X(A2)	
21	1	1	1	0	21	0	
24	1	1	1	0	24	0	
22	1	1	1	0	22	0	
28	2	1	0	1	0	28	
19	2	1	0	1	0	19	
23	2	1	0	1	0	23	

**Table 19.7** Example of Continuous-Nesting-Class Effects

This model estimates a separate intercept and a separate slope for X within each level of A.

# **Continuous-by-Class Effects**

Continuous-by-class effects generate the same design columns as continuous-nesting-class effects. Table 19.8 shows the construction of the X\*A effect. The two columns for this effect are the same as the columns for the X(A) effect in Table 19.7.

Ι	Data	I	X	Α	Χ,	*A
X	Α	$\beta_0$	X	A1 A2	X*A1	X*A2
21	1	1	21	1 0	21	0
24	1	1	24	1 0	24	0
22	1	1	22	1 0	22	0
28	2	1	28	0 1	0	28
19	2	1	19	0 1	0	19
23	2	1	23	0 1	0	23

**Table 19.8** Example of Continuous-by-Class Effects

You can use continuous-by-class effects together with pure continuous effects to test for homogeneity of slopes.

# **General Effects**

An example that combines all the effects is X1\*X2\*A\*B\*C(D E). The continuous list comes first, followed by the crossed list, followed by the nested list in parentheses. You should be aware of the sequencing of parameters when you use statements that depend on the ordering of parameters. Such statements include CONTRAST and ESTIMATE statements, which are used in a number of procedures to estimate and test functions of the parameters.

Effects might be renamed by the procedure to correspond to ordering rules. For example, B\*A(E D) might be renamed A\*B(D E) to satisfy the following:

- Classification variables that occur outside parentheses (crossed effects) are sorted in the order in which they appear in the CLASS statement.
- Variables within parentheses (nested effects) are sorted in the order in which they appear in the CLASS statement.

The sequencing of the parameters generated by an effect can be described by which variables have their levels indexed faster:

- Variables in the crossed list index faster than variables in the nested list.
- Within a crossed or nested list, variables to the right index faster than variables to the left.

For example, suppose a model includes four effects—A, B, C, and D—each having two levels, 1 and 2. If the CLASS statement is

# class A B C D;

then the order of the parameters for the effect  $B*A(C\ D)$ , which is renamed  $A*B(C\ D)$ , is

Note that first the crossed effects B and A are sorted in the order in which they appear in the CLASS statement so that A precedes B in the parameter list. Then, for each combination of the nested effects in turn, combinations of A and B appear. The B effect changes fastest because it is rightmost in the cross list. Then A changes next fastest, and D changes next fastest. The C effect changes most slowly because it is leftmost in the nested list.

# **Other Parameterizations**

This section applies to the following procedures: CATMOD, GENMOD, GLMSELECT, LOGISTIC, PHREG, and SURVEYPHREG.

Some SAS/STAT procedures, including GENMOD, GLMSELECT, and LOGISTIC, support nonsingular parameterizations for classification effects. A variety of these nonsingular parameterizations are available. In most of these procedures you use the PARAM= option in the CLASS statement to specify the parameterization.

Consider a model with one CLASS variable A that has four levels, 1, 2, 5, and 7. Details of the possible choices for the PARAM= option follow.

# **EFFECT**

Three columns are created to indicate group membership of the nonreference levels. For the reference level, all three dummy variables have a value of -1. For example, if the reference level is 7 (REF=7), the design matrix columns for A are as follows.

Effect Coding				
	Design Matrix			
A	A1	<b>A2</b>	A5	
1	1	0	0	
2	0	1	0	
5	0	0	1	
7	-1	-1	-1	

Parameter estimates of CLASS main effects that use the effect coding scheme estimate the difference in the effect of each nonreference level compared to the average effect over all four levels.

The EFFECT parameterization is the default parameterization in the CATMOD procedure. See the section "Generation of the Design Matrix" on page 1939, in Chapter 32, "The CATMOD Procedure," for further details about parameterization of model effects with the CATMOD procedure.

**GLM** 

As in the GLM procedure, four columns are created to indicate group membership. The design matrix columns for A are as follows.

GLM Coding					
	Design Matrix				
A	A1	<b>A2</b>	A5	A7	
1	1	0	0	0	
2	0	1	0	0	
5	0	0	1	0	
7	0	0	0	1	

Parameter estimates of CLASS main effects that use the GLM coding scheme estimate the difference in the effects of each level compared to the last level. See the previous section for details about the GLM parameterization of model effects.

ORDINAL | THERMOMETER Three columns are created to indicate group membership of the higher levels of the effect. For the first level of the effect (which for A is 1), all three dummy variables have a value of 0. The design matrix columns for A are as follows.

_0	<b>Ordinal Coding</b>				
	Design Matrix				
A	<b>A2</b>	<b>A5</b>	<b>A7</b>		
1	0	0	0		
2	1	0	0		
5	1	1	0		
7	1	1	1		

The first level of the effect is a control or baseline level. Parameter estimates of CLASS main effects, using the ORDINAL coding scheme, estimate the differences between effects of successive levels. When the parameters have the same sign, the effect is monotonic across the levels.

**POLYNOMIAL** | **POLY** Three columns are created. The first represents the linear term (x), the second represents the quadratic term  $(x^2)$ , and the third represents the cubic term  $(x^3)$ , where x is the level value. If the CLASS levels are not numeric, they are translated into 1, 2, 3, ... according to their sort order. The design matrix columns for A are as follows.

	Polynomial Coding				
Design Matrix					
A	APOLY1	APOLY2	APOLY3		
1	1	1	1		
2	2	4	8		
5	5	25	125		
7	7	49	343		

# REFERENCE | REF

Three columns are created to indicate group membership of the nonreference levels. For the reference level, all three dummy variables have a value of 0. For example, if the reference level is 7 (REF=7), the design matrix columns for A are as follows.

<b>Reference Coding</b>				
	Design Matrix			
A	A1	<b>A2</b>	A5	
1	1	0	0	
2	0	1	0	
5	0	0	1	
7	0	0	0	

Parameter estimates of CLASS main effects that use the reference coding scheme estimate the difference in the effect of each nonreference level compared to the effect of the reference level.

The REFERENCE parameterization is also available through the MODEL statement in the CATMOD procedure. See the section "Generation of the Design Matrix" on page 1939, in Chapter 32, "The CATMOD Procedure," for further details about parameterization of model effects with the CATMOD procedure.

# **ORTHEFFECT**

The columns are obtained by applying the Gram-Schmidt orthogonalization to the columns for PARAM=EFFECT. The design matrix columns for A are as follows.

	Orthogonal Effect Coding				
	Design Matrix				
A	AOEFF1	AOEFF2	AOEFF3		
1	1.41421	-0.81650	-0.57735		
2	0	1.63299	-0.57735		
5	0	0	1.73205		
7	-1.41421	-0.81649	-0.57735		

**ORTHORDINAL** | **ORTHOTHERM** The columns are obtained by applying the Gram-Schmidt orthogonalization to the columns for PARAM=ORDINAL. The design matrix columns for A are as follows.

	Orthogonal Ordinal Coding				
	Design Matrix				
A	AOORD1	AOORD2	AOORD3		
1	-1.73205	0	0		
2	0.57735	-1.63299	0		
5	0.57735	0.81650	-1.41421		
7	0.57735	0.81650	1.41421		

# **ORTHPOLY**

The columns are obtained by applying the Gram-Schmidt orthogonalization to the columns for PARAM=POLY. The design matrix columns for A are as follows.

	Orthogonal Polynomial Coding				
	Design Matrix				
A	AOPOLY1	AOPOLY2	AOPOLY5		
1	-1.15311	0.90712	-0.92058		
2	-0.73380	-0.54041	1.47292		
5	0.52414	-1.37034	-0.92058		
7	1.36277	1.00363	0.36823		

# **ORTHREF**

The columns are obtained by applying the Gram-Schmidt orthogonalization to the columns for PARAM=REFERENCE. The design matrix columns for A are as follows.

	Orthogonal Reference Coding				
	Design Matrix				
A	AOREF1	AOREF2	AOREF3		
1	1.73205	0	0		
2	-0.57735	1.63299	0		
5	-0.57735	-0.81650	1.41421		
7	-0.57735	-0.81650	-1.41421		

# **CODE Statement**

This statement documentation applies to the following procedures: GENMOD, GLIMMIX, GLM, GLMSELECT, LOGISTIC, MIXED, PLM, and REG. It also applies to the HPLOGISTIC and HPREG procedures in SAS High-Performance Analytics software.

The CODE statement enables you to write SAS DATA step code to a file or catalog entry for computing predicted values of the fitted model. This code can then be included in a DATA step to score new data. For example, in the following program, the CODE statement writes the code for predicting the outcome of a logistic model to the file *mycode.sas*. The file is subsequently included in a DATA step to score the sashelp.Bmt data.

```
proc logistic data=sashelp.Bmt;
   class Group;
   model Status=Group;
   code file='mycode.sas';
run;
data Score;
   set sashelp.Bmt;
   %include mycode;
run;
```

# **Syntax: CODE Statement**

CODE < options > ;

Table 19.9 summarizes the *options* you can specify in the CODE statement.

Table 19.9 CODE Statement Options

Option	Description
CATALOG=	Names the catalog entry where the generated code is saved
DUMMIES	Retains the dummy variables in the data set
ERROR	Computes the error function
FILE=	Names the file where the generated code is saved
FORMAT=	Specifies the numeric format for the regression coefficients
GROUP=	Specifies the group identifier for array names and statement labels
IMPUTE	Imputes predicted values for observations with missing or invalid covariates
LINESIZE=	Specifies the line size of the generated code
LOOKUP=	Specifies the algorithm for looking up CLASS levels
RESIDUAL	Computes residuals

You cannot specify both the FILE= and CATALOG= options. If you specify neither, the SAS scoring code is written to the SAS log. You can specify the following options in the CODE statement.

# CATALOG=library.catalog.entry.type

# CAT=library.catalog.entry.type

specifies where to write the generated code in the form of library.catalog.entry.type. The compound name can have from one to four levels. The default library is determined by the USER= SAS system option, which by default is WORK. The default entry is SASCODE, and the default type is SOURCE.

# **DUMMIES | NODUMMIES**

specifies whether to keep dummy variables that represent the CLASS levels in the data set. The default is NODUMMIES, which specifies that dummy variables not be retained.

# **ERROR | NOERROR**

specifies whether to generate code to compute the error function. The default is NOERROR, which specifies that the error function not be generated.

# FILE=filename

names the external file that saves the generated code. When enclosed in a quoted string (for example, FILE="c:\mydir\scorecode.sas"), this option specifies the path for writing the code to an external file. You can also specify unquoted SAS filenames of no more than eight characters for *filename*. If the *filename* is assigned as a *fileref* in a Base SAS FILENAME statement, the file specified in the FILENAME statement is opened. The special filerefs LOG and PRINT are always assigned. If the specified *filename* is not an assigned *fileref*, the specified value for *filename* is concatenated with a .txt extension before the file is opened. For example, if FOO is not an assigned fileref, FILE=FOO causes FOO.txt to be opened. If filename has more than eight characters, an error message is printed.

# **FORMAT**=format

specifies the format for the regression coefficients and other numerical values that do not have a format from the input data set. The default format is BEST20.

# GROUP=group-name

specifies the group identifier for group processing. The *group-name* should be a valid SAS name of no more than 16 characters. It is used to construct array names and statement labels in the generated code.

# **IMPUTE**

imputes the predicted values according to an intercept-only model for observations with missing or invalid covariate values. For a continuous response, the predicted value is the mean of the response variable; for a categorical response, the predicted values are the proportions of the response categories. When the IMPUTE option is specified, the scoring code also creates a variable named \_WARN\_ that contains one or more single-character codes that indicate problems in computing predicted values. The character codes used in \_WARN\_ go in the following positions:

**Table 19.10** \_WARN\_ Variable Codes

Code	Column	Meaning
M	1	Missing covariate value
U	2	Unrecognized covariate category

#### LINESIZE=value

# LS=value

specifies the line size for the generated code. The default is 78. The permissible range is 78 to 254.

# LOOKUP=lookup-method

specifies the algorithm for looking up CLASS levels. You can specify the following *lookup-methods*:

### **AUTO**

selects the LINEAR algorithm if a CLASS variable has fewer than five categories; otherwise, the BINARY algorithm is used. This is the default.

# **BINARY**

uses a binary search. This method is fast, but might produce incorrect results and the normalized category values might contain characters that collate in different orders in ASCII and EBCDIC, if you generate the code on an ASCII machine and execute the code on an EBCDIC machine or vice versa.

# LINEAR

uses a linear search with IF statements that have categories in the order of the class levels. This method is slow if there are many categories.

# **SELECT**

uses a SELECT statement.

The default is LOOKUP=AUTO.

# RESIDUAL | NORESIDUAL

specifies whether to generate code to compute residual values. If you request code for residuals and then score a data set that does not contain target values, the residuals will have missing values. The default is NORESIDUAL, which specifies that the code for residuals not be generated.

# **EFFECT Statement**

This section applies to the following procedures:

GLIMMIX, GLMSELECT, HPMIXED, LOGISTIC, ORTHOREG, PHREG, PLS, QUANTLIFE, QUANTREG, QUANTSELECT, ROBUSTREG, SURVEYLOGISTIC, and SURVEYREG.

The EFFECT statement enables you to construct special collections of columns for design matrices. These collections are referred to as *constructed effects* to distinguish them from the usual model effects that are formed from continuous or classification variables, as discussed in the section "GLM Parameterization of Classification Variables and Effects" on page 387. For example, the terms A, B, x, A\*x, A\*B, and sub in the following statements define fixed, random, and subject effects of the usual type in a mixed model:

```
proc glimmix;
   class A B sub;
   model y = A B x A*x;
   random A*B / subject=sub;
run;
```

A constructed effect, on the other hand, is assigned through the EFFECT statement. For example, in the following program, the EFFECT statement defines a constructed effect named spl:

```
proc glimmix;
   class A B SUB;
   effect spl = spline(x);
   model y = A B A*spl;
   random A*B / subject=sub;
run;
```

The columns of spl are formed from the data set variable x as a cubic B-spline basis with three equally spaced interior knots.

Each constructed effect corresponds to a collection of columns that are referred to by using the name you supply. You can specify multiple EFFECT statements, and all EFFECT statements must precede the MODEL statement.

The general syntax for the EFFECT statement with *effect-specification* is

```
EFFECT effect-name = effect-type (var-list < / effect-options >);
```

The name of the effect is specified after the EFFECT keyword. This name can appear in only one EFFECT statement and cannot be the name of a variable in the input data set. The effect-type is specified after an equal sign, followed by a list of variables within parentheses which are used in constructing the effect. Effect-options that are specific to an effect-type can be specified after a slash (/) following the variable list. The following *effect-types* are available and are discussed in the following sections:

COLLECTION	is a collection effect that defines one or more	e variables as a single effect with
------------	---	-------------------------------------

multiple degrees of freedom. The variables in a collection are considered as

a unit for estimation and inference.

LAG is a classification effect in which the level that is used for a given period

corresponds to the level in the preceding period.

**MULTIMEMBER | MM** is a multimember classification effect whose levels are determined by one or

more variables that appear in a CLASS statement.

**POLYNOMIAL | POLY** is a multivariate polynomial effect in the specified numeric variables.

**SPLINE** is a regression spline effect whose columns are univariate spline expansions

of one or more variables. A spline expansion replaces the original variable

with an expanded or larger set of new variables.

Table 19.11 summarizes the *options* available in the EFFECT statement.

Table 19.11 EFFECT Statement Options

Option	Description		
Collection Effects Optio DETAILS	ns Displays the constituents of the collection effect		
Lag Effects Options DESIGNROLE=	Names a variable that controls to which lag design an observation is assigned		
DETAILS	Displays the lag design of the lag effect		
NLAG=	Specifies the number of periods in the lag		
PERIOD=	Names the variable that defines the period		
WITHIN=	Names the variable or variables that define the group within which each period is defined		
<b>Multimember Effects Options</b>			
NOEFFECT WEIGHT=	Specifies that observations with all missing levels for the multi- member variables should have zero values in the corresponding design matrix columns Specifies the weight variable for the contributions of each of the classification effects		
Polynomial Effects Options			
DEGREE= MDEGREE=	Specifies the degree of the polynomial Specifies the maximum degree of any variable in a term of the polynomial		
STANDARDIZE=	Specifies centering and scaling suboptions for the variables that define the polynomial		
Spline Effects Options BASIS=	Specifies the type of basis (B-spline basis or truncated power function basis) for the spline effect		
DEGREE= KNOTMETHOD=	Specifies the degree of the spline effect Specifies how to construct the knots for the spline effect		

# **Collection Effects**

# **EFFECT** name=**COLLECTION** (var-list </ **DETAILS**>);

You use a collection effect to define a set of variables that are treated as a single effect with multiple degrees of freedom. The variables in *var-list* can be continuous or classification variables. The columns in the design matrix that are contributed by a collection effect are the design columns of its constituent variables in the order in which they appear in the definition of the collection effect. If you specify the DETAILS option, then a table that shows the constituents of the collection effect is displayed.

# Lag Effects

# EFFECT name=LAG (variable / lag-options);

A lag effect is a classification effect for the CLASS variable that is given after the keyword LAG. A lag effect is used to represent the effect of a previous value of the lagged variable when there is some inherent ordering of the observations of this variable. A typical example where lag effects are useful is a study in which different subjects are given sequences of treatments and you want to investigate whether the treatment in the previous period is important in understanding the outcome in the current period. You can do this by including a lagged treatment effect in your model.

The precise definition of a LAG effect depends on a subdivision of the data into disjoint subsets, often referred to as "subjects," and an ordering into units called "periods" of the observations within a subject. For an observation that belongs to a given subject and at a given period, the design matrix columns of the lagged variable are the usual design matrix columns of that variable except for the observation at the preceding period for that subject. Observations at the initial period do not have a preceding value, and so the design matrix columns of the lag effect for these observations are set to zero. You can also define lag effects where the number of periods that are lagged is greater than one. If the number of periods that are lagged is n, then the design matrix columns of observations in periods less than or equal to n are set to zero. The design matrix columns that correspond to a subject at period p, where p > n, are the usual design matrix columns of the lagged variable for that subject at period p - n.

A convenient way to represent the organization of observations into subjects and periods is to form the lag design matrix. The rows and columns of this matrix correspond to the subjects and periods respectively. The lag design matrix entry is the treatment for the corresponding subject and period. In a valid lag design there is at most one observation for a given period and subject. For example, the following set of treatments by subject and period form a valid lag design:

Subject	Period	<b>Treatment</b>
Sheila	1	В
Joey	1	A
Athena	1	A
Gelindo	1	A
Sheila	2	C
Joey	2	A
Athena	2	
Gelindo	2	В
Sheila	3	В
Joey	3	C
Athena	3	A
Gelindo	3	В

The associated lag design matrix is

	Period		
Subject	1	2	3
Athena	Α		A
Gelindo	Α	В	В
Joey	Α	A	C
Sheila	В	C	В

Note that the subject Athena did not receive a treatment at period 2, and so the corresponding entry in the lag design matrix is missing. You can define a lag effect for this lag design with the following statements:

```
CLASS treatment;
EFFECT Lag = LAG( treatment / WITHIN=subject PERIOD=period);
```

When GLM coding is used for the CLASS variable treatment, the design matrix columns Lag\_A, Lag\_B, and Lag\_C for the constructed effect Lag are as follows:

Subject	Period	<b>Treatment</b>	Lag_A	Lag_B	Lag_C
Athena	1	A	0	0	0
Athena	2		1	0	0
Athena	3	A			
Gelindo	1	A	0	0	0
Gelindo	2	В	1	0	0
Gelindo	3	В	0	1	0
Joey	1	A	0	0	0
Joey	2	A	1	0	0
Joey	3	C	1	0	0
Sheila	1	В	0	0	0
Sheila	2	C	0	1	0
Sheila	3	В	0	0	1

The design matrix columns for each subject at period 1 are all zero because there are no lagged observations for period 1. You can also see that the design matrix columns at period 3 for subject Athena are missing because Athena did not receive a treatment at period 2. Nevertheless, the design matrix columns for Athena at period 2 are nonmissing and correspond to the treatment "A" that she received in period 1.

The following *lag-options* are required:

# PERIOD=variable

specifies the period variable of the LAG design. The number of periods is the number of unique formatted values of the PERIOD= variable, and the ordering of the period is formed by sorting these formatted values in ascending order. You must specify a PERIOD= variable.

# WITHIN=(variables)

# WITHIN=variable

specifies a variable (or a list of variables within parentheses) that defines the subject grouping of the lag design. If there is only one WITHIN= variable, then the parentheses are not required. Each subject is defined by the unique set of formatted values of the variables in the WITHIN= list. The subjects are sorted in ascending lexicographic order. You must specify a WITHIN= variable.

You can also specify the following *lag-options*:

# **DESIGNROLE**=variable

specifies a numeric variable that is used to subset observations into a fitting group in which the value of the DESIGNROLE= variable is nonzero and a second group in which the value of the specified variable is zero. The observations in the fitting group are used to form the LAG design matrix that is used in fitting the model. The LAG design that corresponds to the non-fitting group is used when scoring observations in the input data set that do not belong to the fitting group. This option is useful when you want to obtain predicted values in an output data set for observations that are not used in fitting the model. If you do not specify a DESIGNROLE= variable, then all observations are assigned to the fitting group.

#### **DETAILS**

requests a table that shows the lag design matrix of the lag effect.

# NLAG = n

specifies the number of lags. By default NLAG=1.

# **Multimember Effects**

```
EFFECT name=MULTIMEMBER (var-list < / mm-options >);
```

**EFFECT** name=**MM** (var-list < / mm-options >);

A multimember effect is formed from one or more classification variables in such a way that each observation can be associated with one or more levels of the union of the levels of the classification variables. In other words, a multimember effect is a classification-type effect with possibly more than one nonzero column entry for each observation. Multimember effects are useful, for example, in modeling the following:

- nurses' effects on patient recovery in hospitals
- teachers' effects on student scores
- lineage effects in genetic studies. See Example 44.16 in Chapter 44, "The GLIMMIX Procedure," for an application with random multimember effects in a genetic diallel experiment.

The levels of a multimember effect consist of the union of formatted values of the variables that define this effect. Each such level contributes one column to the design matrix. For each observation, the value that corresponds to each level of the multimember effect in the design matrix is the number of times that this level occurs for the observation.

For example, the following data provide teacher information and end-of-year test scores for students after two semesters:

Student	Score	Teacher1	Teacher2
Mary	87	Tobias	Cohen
Tom	89	Rodriguez	Tobias
Fred	82	Cohen	Cohen
Jane	88	Tobias	
Jack	99		

For example, Mary had different teachers in the two semesters, Fred had the same teacher in both semesters, and Jane received instruction only in the first semester.

You can model the effect of the teachers on student performance by using a multimember effect specified as follows:

```
CLASS teacher1 teacher2;
EFFECT teacher = MM(teacher1 teacher2);
```

The levels of the teacher effect are Cohen, Rodriguez, and Tobias, and the associated design matrix columns are as follows:

Student	Cohen	Rodriguez	Tobias
Mary	1	0	1
Tom	0	1	1
Fred	2	0	0
Jane	0	0	1
Jack		•	

You can specify the following *mm-options* after a slash (/):

#### **DETAILS**

requests a table that shows the levels of the multimember effect.

# **NOEFFECT**

specifies that, for observations with all missing levels of the multimember variables, the values in the corresponding design matrix columns be set to zero. If, in the preceding example, the teacher effect is defined by

```
EFFECT teacher = MM(teacher1 teacher2 / noeffect);
```

then the associated design matrix columns' values for Jack are all zero. This enables you to include Jack in the analysis even though there is no effect of teachers on his performance.

A situation where it is important to designate observations as having no effect due to a classification variable is the analysis of crossover designs, where lagged treatment levels are used to model the carryover effects of treatments between periods. Since there is no carryover effect for the first period, the treatment lag effect in a crossover design can be modeled with a multimember effect that consists of a single classification variable and the NOEFFECT option, as in the following statements:

```
CLASS Treatment lagTreatment;
EFFECT Carryover = MM(lagTreatment / noeffect);
```

The lagTreatment variable contains a missing value for the first period. Otherwise, it contains the value of the treatment variable for the preceding period.

# **STDIZE**

specifies that for each observation, the entries in the design matrix that corresponds to the multimember effect be scaled to have a sum of one.

# WEIGHT=wght-list

specifies numeric variables used to weigh the contributions of each of the classification effects that define the constructed multimember effect. The number of variables in *wght-list* must match the number of classification variables that define the effect.

# **Polynomial Effects**

```
EFFECT name=POLYNOMIAL (var-list </polynomial-options>); 
EFFECT name=POLY (var-list </polynomial-options>);
```

The variables in *var-list* must be numeric. A design matrix column is generated for each term of the specified polynomial. By default, each of these terms is treated as a separate effect for the purpose of model building. For example, the statements

```
proc glmselect;
    effect MyPoly = polynomial(x1-x3/degree=2);
    model y = MyPoly;
run;
yield the identical analysis to the statements

proc glmselect;
    model y = x1 x2 x3 x1*x1 x1*x2 x1*x3 x2*x2 x2*x3 x3*x3;
run:
```

You can specify the following *polynomial-options* after a slash (/):

# **DEGREE**=n

specifies the degree of the polynomial. The degree must be a positive integer. The degree is typically a small integer, such as 1, 2, or 3. The default is DEGREE=1.

# **DETAILS**

requests a table that shows the details of the specified polynomial, including the number of terms generated. If you also specify the STANDARDIZE option, then a table that shows the standardization details is also produced.

# LABELSTYLE=(style-opts)

# LABELSTYLE=style-opt

specifies how the terms in the polynomial are labeled. By default, powers are shown with  $\hat{}$  as the exponentiation operator and \* as the multiplication operator. For example, a polynomial term such as  $x_1^3x_2x_3^2$  is labeled x1^3\*x2\*x3^2. You can change the style of the label by using the following *style-opts* within parentheses. If you specify a single *style-opt*, then you can omit the enclosing parentheses.

# **EXPAND**

specifies that each variable with an exponent greater than 1 be written as products of that variable. For example, the term  $x_1^3x_2x_3^2$  receives the label  $x_1^*x_1^*x_1^*x_2^*x_3^*x_3$ .

# **EXPONENT** <= quoted string >

specifies that each variable with an exponent greater than 1 be written using exponential notation. By default, the symbol ^ is used as the exponentiation operator. If you supply the optional quoted string after an equal sign, then that string is used as the exponentiation operator. For example, if you specify

```
LABELSTYLE= (EXPONENT="**")
```

then the term  $x_1^3x_2x_3^2$  receives the label x1\*\*3\*x2\*x3\*\*2.

# **INCLUDENAME**

specifies that the name of the effect followed by an underscore be used as a prefix for term labels. For example, the following statement generates terms with labels MyPoly x1 and MyPoly x1^2:

```
EFFECT MyPoly=POLYNOMIAL(x1/degree=2 labelstyle=INCLUDENAME)
```

The INCLUDENAME option is ignored if you also specify the NOSEPARATE option in the EFFECT=POLYNOMIAL statement.

# PRODUCTSYMBOL=NONE | quoted string

specifies that the supplied string be used as the product symbol. For example, the following statement generates terms with labels x1, x2, and x1 x2:

If you specify PRODUCTSYMBOL=NONE, then the labels are formed by juxtaposing the constituent variable names.

# MDEGREE=n

specifies the maximum degree of any variable in a term of the polynomial. This degree must be a positive integer. The default is the degree of the specified polynomial. For example, the following statement generates the terms  $x_1$ ,  $x_2$ ,  $x_1^2$ ,  $x_1x_2$ ,  $x_2^2$ ,  $x_1^2x_2$ ,  $x_1x_2^2$  and  $x_1^2x_2^2$ :

```
EFFECT MyPoly=POLYNOMIAL(x1 x2/degree=4 MDEGREE=2);
```

# **NOSEPARATE**

specifies that the polynomial be treated as a single effect with multiple degrees of freedom. The effect name that you specify is used as the constructed effect name, and the labels of the terms are used as labels of the corresponding parameters.

specifies that the variables that define the polynomial be standardized. By default, the standardized variables receive prefix "s\_" in the variable names.

You can use the following centerscale-opts to specify how the center and scale are estimated:

# **METHOD=MOMENTS**

specifies that the center be estimated by the variable mean and the scale be estimated by the standard deviation. If a weight variable is specified using a WEIGHT statement, the observations with invalid weights are ignored when forming the mean and standard deviation, but the weights are otherwise not used. Only observations that are used in performing the analysis are used for the standardization.

#### METHOD=RANGE

specifies that the center be estimated by the midpoint of the variable range and the scale be estimated as half the variable range. Any observation that has a missing value for any regressor used in the model is ignored when computing the range of variables in a polynomial effect. Observations with valid regressor values but missing or invalid values of frequency variables, weight variables, or dependent variables are used in computing variable ranges. The default (if you do not specify the METHOD= suboption) is METHOD=RANGE.

# **METHOD=WMOMENTS**

is the same as METHOD=MOMENTS except that weighted means and weighted standard deviations are used.

Let

n = number of observations used in the analysis w = weight variable f = frequency variable x = variable to be standardized  $x_{(n)} = \text{Max}_{i=1}^{n}(x_i)$   $x_{(1)} = \text{Min}_{i=1}^{n}(x_i)$  F = sum of frequencies  $= \sum_{i=1}^{n} f_i$ WF = sum of weighted frequencies  $= \sum_{i=1}^{n} w_i f_i$ 

Table 19.12 shows how the center and scale are computed for each of the supported methods.

 Method
 Center
 Scale

 Range
  $(x_{(n)} + x_{(1)})/2$   $(x_{(n)} - x_{(1)})/2$  

 Moments
  $\bar{x} = \sum_{i=1}^{n} f_i x_i / F$   $\sqrt{\sum_{i=1}^{n} f_i (x_i - \bar{x})^2 / (F - 1)}$  

 WMoments
  $\bar{x}_w = \sum_{i=1}^{n} w_i f_i x_i / WF$   $\sqrt{\sum_{i=1}^{n} w_i f_i (x_i - \bar{x}_w)^2 / (F - 1)}$ 

Table 19.12 Center and Scale Estimates by Method

# PREFIX=NONE | quoted-string

specifies the prefix that is appended to standardized variables when forming the term labels. If you omit this option, the default prefix is "s\_". If you specify PREFIX=NONE, then standardized variables are not prefixed.

You can control whether the standardization is to center, scale, or both center and scale by specifying a *standardize-opt*:

#### **CENTER**

specifies that variables be centered but not scaled. For a variable x,

$$s_x = x - center$$

# **CENTERSCALE**

specifies that variables be centered and scaled. This is the default if you do not specify a standardization-opt. For a variable x,

$$s_x = \frac{x - \text{center}}{\text{scale}}$$

# **NONE**

specifies that no standardization be performed.

# **SCALE**

specifies that variables be scaled but not centered. For a variable x,

$$s_x = \frac{x}{\text{scale}}$$

# **Spline Effects**

This section discusses the construction of spline effects through the EFFECT statement. You can also include spline effects in statistical models by other means. The TRANSREG procedure has dedicated facilities for including regression splines in your model and controlling the construction of the splines. For example, you can use the TRANSREG procedure to fit a spline function but restrict the function to be always increasing or decreasing (monotone). See the section "Using Splines and Knots" on page 8614 in Chapter 104, "The TRANSREG Procedure," for more information about using splines with the TRANSREG procedure. The

GAM and TPSPLINE procedures also can model the effects of regressor variables in terms of smooth functions that are generated from spline bases. For more information see Chapter 41, "The GAM Procedure," and Chapter 103, "The TPSPLINE Procedure."

A spline effect expands variables into spline bases whose form depends on the options that you specify. You can find details about regression splines and spline bases in the section "Splines and Spline Bases" on page 411. You request a spline effect with the syntax

# **EFFECT** name=**SPLINE** (var-list < / spline-options >);

The variables in *var-list* must be numeric. Design matrix columns are generated separately for each of these variables, and the set of columns is collectively referred to with the specified name. By default, the spline basis that is generated for each variable is a cubic B-spline basis with three equally spaced knots positioned between the minimum and maximum values of that variable. This yields by default seven design matrix columns for each of the variables in the SPLINE effect.

You can specify the following *spline-options* after a slash (/):

# **BASIS=BSPLINE**

specifies a B-spline basis for the spline expansion. For splines of degree d defined with n knots, this basis consists of n + d + 1 columns. In order to completely specify the B-spline basis, d left-side boundary knots and  $\max\{d,1\}$  right-side boundary knots are also required. See the suboptions KNOTMETHOD=, DATABOUNDARY, KNOTMIN=, and KNOTMAX= for details about how to specify the positions of both the internal and boundary knots. This is the default if you do not specify the BASIS= suboption.

# BASIS=TPF(options)

specifies a truncated power function basis for the spline expansion. For splines of degree d defined with n knots for a variable x, this basis consists of an intercept, polynomials x,  $x^2$ , ...,  $x^d$  and one truncated power function for each of the n knots. Unlike the B-spline basis, no boundary knots are required. See the suboption KNOTMETHOD= for details about how you can specify the position of the internal knots.

You can modify the number of columns when you request BASIS=TPF with the following options:

# **NOINT**

excludes the intercept column.

# **NOPOWERS**

excludes the intercept and polynomial columns.

# **DATABOUNDARY**

specifies that the extremes of the data be used as boundary knots when building a B-spline basis.

# **DEGREE**=n

specifies the degree of the spline transformation. The degree must be a nonnegative integer. The degree is typically a small integer, such as 0, 1, 2, or 3. The default is DEGREE=3.

# **DETAILS**

requests tables that show the knot locations and the knots associated with each spline basis function.

# KNOTMAX=value

specifies that, for each variable in the EFFECT statement, the right-side boundary knots be equally spaced starting at the maximum of the variable and ending at the specified value. This option is ignored for variables whose maximum value is greater than the specified value or if the DATABOUNDARY option is also specified.

# **KNOTMETHOD**=knot-method< (knot-options)>

specifies how to construct the knots for spline effects. You can choose from the following *knot-methods* and affect the knot construction further with the method-specific *knot-options*:

# EQUAL<(n)>

specifies that n equally spaced knots be positioned between the extremes of the data. The default is n = 3. For a B-spline basis, any needed boundary knots continue to be equally spaced unless the DATABOUNDARY option has also been specified. KNOTMETHOD=EQUAL is the default if no *knot-method* is specified.

# LIST(number-list)

specifies the list of internal knots to be used in forming the spline basis columns. For a B-spline basis, the data extremes are used as boundary knots.

# **LISTWITHBOUNDARY**(number-list)

specifies the list of all knots that are used in forming the spline basis columns. When you use a truncated power function basis, this list is interpreted as the list of internal knots. When you use a B-spline basis of degree d, then the first d entries are used as left-side boundary knots and the last MAX(d, 1) entries in the list are used as right-side boundary knots.

# MULTISCALE < (multiscale-options) >

specifies that multiple B-spline bases be generated, corresponding to sets with an increasing number of internal knots. As you increase the number of internal knots, the spline basis you generate is able to approximate features of the data at finer scales. So, by generating bases at multiple scales, you facilitate the modeling of both coarse- and fine-grained features of the data. For scale i, the spline basis corresponds to  $2^i$  equally spaced internal knots. By default, the bases for scales 0-7 are generated. For each scale, a separate spline effect is generated. The name of the constructed spline effect at scale i is formed by appending Si to the effect name that you specify in the EFFECT statement. If you specify multiple variables in the EFFECT statement, then spline bases are generated separately for each variable at each scale and the name of the corresponding effect is obtained by appending the variable name followed by Si to the name in the EFFECT statement. For example, the following statement generates effects named splSi0, splSi1, splSi2, splSi3, splSi3, splSi4, splSi5, s

# EFFECT spl = spline(x1 x2 / knotmethod=multiscale);

The MULTISCALE option is ignored if you specify the BASIS=TPF *spline-option*. The MULTI-SCALE option is not available for spline effects that are specified in the RANDOM statement of the GLIMMIX procedure.

You can control which scales are included with the following *multiscale-options*:

#### STARTSCALE=n

specifies the start scale, where n is a positive integer. The default is STARTSCALE=0.

# **ENDSCALE**=n

specifies the end scale, where n is a positive integer. The default is ENDSCALE=7.

# PERCENTILES(n)

requests that internal knots be placed at *n* equally spaced percentiles of the variable or variables named in the EFFECT statement. For example, the following statement positions internal knots at the deciles of the variable x. For a B-spline basis, the extremes of the data are used as boundary knots:

```
EFFECT spl = spline(x / knotmethod=percentiles(9));
```

# **RANGEFRACTIONS**(fraction-list)

requests that internal knots be placed at each fraction of the ranges of the variables in the EFFECT statement. For example, if variable x1 ranges between 1 and 3, and variable x2 ranges between 0 and 20, then the following EFFECT statement uses internal knots 1.2, 2, and 2.5 for variable x1 and internal knots 2, 10, and 15 for variable x2:

```
EFFECT spl = spline(x1 x2 / knotmethod=rangefractions(.1 .5 .75));
```

For a B-spline basis, the data extremes are used as boundary knots.

# KNOTMIN=value

specifies that for each variable in the EFFECT statement, the left-side boundary knots be equally spaced starting at the specified value and ending at the minimum of the variable. This option is ignored for variables whose minimum value is less than the specified value or if the DATABOUNDARY option is also specified.

# **NATURALCUBIC**

specifies a natural cubic spline basis for the spline expansion. Natural cubic splines, also known as restricted cubic splines, are cubic splines that are constrained to be linear beyond the extreme knots. The natural cubic spline basis that is produced by the EFFECT statement is obtained by starting from the unrestricted truncated power function cubic spline basis that is defined with n distinct knots and imposes the linearity constraints beyond the extreme knots. This basis consists of an intercept, the polynomial x, and n-2 functions that are all linear beyond the largest knot. The ith function,  $i=1,2,\ldots,n-2$ , is zero to the left of the ith knot, which is called the "break knot." See the section "Splines and Spline Bases" on page 411 for details of this basis. You can use the NOINT and NOPOWERS suboptions of the BASIS=TPF option to suppress the intercept and polynomial x when forming the columns of the natural cubic spline basis. When you specify the NATURALCUBIC option, the options BASIS=BSPLINE, DATABOUNDARY, DEGREE=, and KNOTMETHOD=MULTISCALE are not applicable.

# **SEPARATE**

specifies that when multiple variables are specified in the EFFECT statement, the spline basis for each variable be treated as a separate effect. The names of these separated effects are formed by appending an underscore followed by the name of the variable to the name that you specify in the EFFECT statement. For example, the effect names generated with the following statement are spl\_x1 and spl\_x2:

In procedures that support variable selection, such as the GLMSELECT procedure, these two effects can enter or leave the model independently during the selection process. Separated effects are not supported in the RANDOM statement of the GLIMMIX procedure.

# **SPLIT**

specifies that each individual column in the design matrix that corresponds to the spline effect be treated as a separate effect that can enter or leave the model independently. Names for these split effects are generated by appending the variable name and an index for each column to the name that you specify in the EFFECT statement. For example, the effects generated for the spline effect in the following statement are spl x1:1, spl x1:2,..., spl x1:2, and spl x2:1, spl x2:2,..., spl x2:7:

```
EFFECT spl = spline(x1 x2 / split);
```

The SPLIT option is not supported in the GLIMMIX procedure.

# **Splines and Spline Bases**

This section provides details about the construction of spline bases with the EFFECT statement. A spline function is a piecewise polynomial function in which the individual polynomials have the same degree and connect smoothly at join points whose abscissa values, referred to as knots, are prespecified. You can use spline functions to fit curves to a wide variety of data.

A spline of degree 0 is a step function with steps located at the knots. A spline of degree 1 is a piecewise linear function where the lines connect at the knots. A spline of degree 2 is a piecewise quadratic curve whose values and slopes coincide at the knots. A spline of degree 3 is a piecewise cubic curve whose values, slopes, and curvature coincide at the knots. Visually, a cubic spline is a smooth curve, and it is the most commonly used spline when a smooth fit is desired. Note that when no knots are used, splines of degree *d* are simply polynomials of degree *d*.

More formally, suppose you specify knots  $k_1 < k_2 < k_3 < \cdots < k_n$ . Then a spline of degree  $d \ge 0$  is a function S(x) with d-1 continuous derivatives such that

$$S(x) = \begin{cases} P_0(x) & x < k_1 \\ P_i(x) & k_i \le x < k_{i+1}; i = 1, 2, \dots, n-1 \\ P_n(x) & x \ge k_n \end{cases}$$

where each  $P_i(x)$  is a polynomial of degree d. The requirement that S(x) has d-1continuous derivatives is satisfied by requiring that the function values and all derivatives up to order d-1 of the adjacent polynomials at each knot match.

A counting argument yields the number of parameters that define a spline with n knots. There are n + 1 polynomials of degree d, giving (n + 1)(d + 1) coefficients. However, there are d restrictions at each of the n knots, so the number of free parameters is (n + 1)(d + 1) - nd = n + d + 1. In mathematical terminology this says that the dimension of the vector space of splines of degree d on n distinct knots is n + d + 1. If you have n + d + 1 basis vectors, then you can fit a curve to your data by regressing your dependent variable by

using this basis for the corresponding design matrix columns. In this context, such a spline is known as a regression spline. The EFFECT statement provides a simple mechanism for obtaining such a basis.

If you remove the restriction that the knots of a spline must be distinct and allow repeated knots, then you can obtain functions with less smoothness and even discontinuities at the repeated knot location. For a spline of degree d and a repeated knot with multiplicity  $m \le d$ , the piecewise polynomials that join such a knot are required to have only d-m matching derivatives. Note that this increases the number of free parameters by m-1 but also decreases the number of distinct knots by m-1. Hence the dimension of the vector space of splines of degree d with n knots is still n+d+1, provided that any repeated knot has a multiplicity less than or equal to d.

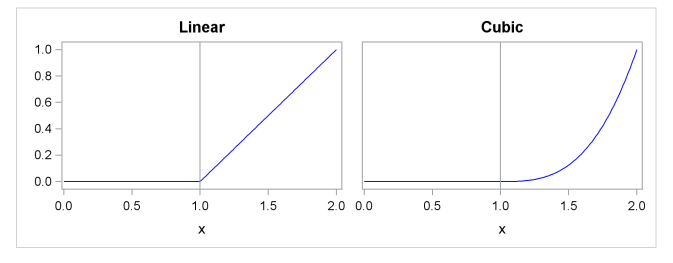
The EFFECT statement provides support for the commonly used *truncated power function* basis and *B-spline* basis. With exact arithmetic and by using the complete basis, you obtain the same fit with either of these bases. The following sections provide details about constructing spline bases for the space of splines of degree d with n knots that satisfies  $k_1 \le k_2 \le k_3 < \cdots \le k_n$ .

# **Truncated Power Function Basis**

A truncated power function for a knot  $k_i$  is a function defined by

$$t_i(x) = \begin{cases} 0 & x < k_i \\ (x - k_i)^d & x \ge k_i \end{cases}$$

Figure 19.1 shows such functions for d = 1 and d = 3 with a knot at x = 1.



**Figure 19.1** Truncated Power Functions with Knot at x = 1

The name is derived from the fact that these functions are shifted power functions that get truncated to zero to the left of the knot. These functions are piecewise polynomial functions with two pieces whose function values and derivatives of all orders up to d-1 are zero at the defining knot. Hence these functions are splines of degree d. It is easy to see that these n functions are linearly independent. However, they do not form a basis, because such a basis requires n+d+1 functions. The usual way to add d+1 additional basis

functions is to use the polynomials  $1, x, x^2, \dots, x^d$ . These d+1 functions together with the n truncated power functions  $t_i(x)$ ,  $i=1,2,\ldots,n$  form the truncated power basis.

Note that each time a knot is repeated, the associated exponent used in the corresponding basis function is reduced by 1. For example, for splines of degree d with three repeated knots  $k_i = k_{i+1} = k_{i+2}$  the corresponding basis functions are  $t_i(x) = (x - k_i)_+^d$ ,  $t_{i+1}(x) = (x - k_i)_+^{d-1}$ , and  $t_{i+2}(x) = (x - k_i)_+^{d-2}$ . Provided that the multiplicity of each repeated knot is less than or equal to the degree, this construction continues to yield a basis for the associated space of splines.

The main advantage of the truncated power function basis is the simplicity of its construction and the ease of interpreting the parameters in a model that corresponds to these basis functions. However, there are two weaknesses when you use this basis for regression. These functions grow rapidly without bound as x increases, resulting in numerical precision problems when the x data span a wide range. Furthermore, many or even all of these basis functions can be nonzero when evaluated at some x value, resulting in a design matrix with few zeros that precludes the use of sparse matrix technology to speed up computation. This weakness can be addressed by using a B-spline basis.

# **B-Spline Basis**

A B-spline basis can be built by starting with a set of Haar basis functions, which are functions that are 1 between adjacent knots and 0 elsewhere, and then applying a simple linear recursion relationship d times, yielding the n+d+1 needed basis functions. For the purpose of building the B-spline basis, the n prespecified knots are referred to as internal knots. This construction requires d additional knots, known as boundary knots, to be positioned to the left of the internal knots, and MAX(d,1) boundary knots to be positioned to the right of the internal knots. The actual values of these boundary knots can be arbitrary. The EFFECT statement provides several methods for placing the needed boundary knots, including the common method of using repeated values of the data extremes as the boundary knots. The boundary knot placement affects the precise form of the basis functions that are generated, but it does not affect the following two desirable properties:

- 1. The B-spline basis functions are nonzero over an interval that spans at most d + 2 knots. This yields design matrix columns each of whose rows contain at most d + 2 adjacent nonzero entries.
- 2. The computation of the basis functions at any *x* value is numerically stable and does not require evaluating powers of this value.

The following figures show the B-spline bases defined on [0, 1] with four equally spaced internal knots at 0.2, 0.4, 0.6, and 0.8.

Figure 19.2 shows a linear B-spline basis. Note that this basis consists of six functions each of which is nonzero over an interval that spans at most three knots.

Figure 19.2 Linear B-Spline Basis with Four Equally Spaced Interior Knots

Figure 19.3 shows a cubic B-spline basis where the needed boundary knots are positioned at x = 0 and x = 1. Note that this basis consists of eight functions, each of which is nonzero over an interval spanning at most five knots.

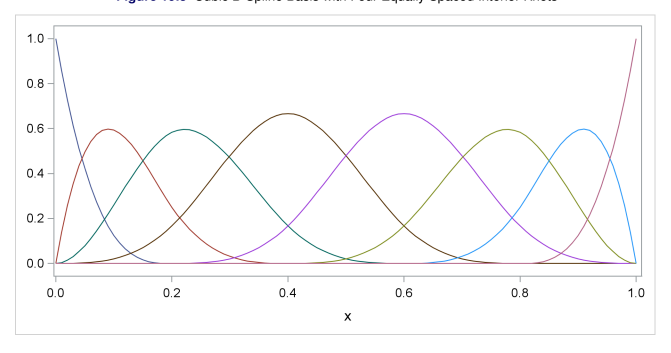


Figure 19.3 Cubic B-Spline Basis with Four Equally Spaced Interior Knots

Figure 19.4 shows a different cubic B-spline basis where the needed left-side boundary knots are positioned at -0.6, -0.4, -0.2, and 0. The right-side boundary knots are positioned at 1, 1.2, 1.4, and 1.6. Note that, as in the basis shown in Figure 19.3, this basis consists of eight functions, each of which is nonzero over an interval spanning at most five knots. The different positioning of the boundary knots has merely changed the shape of the individual basis functions.

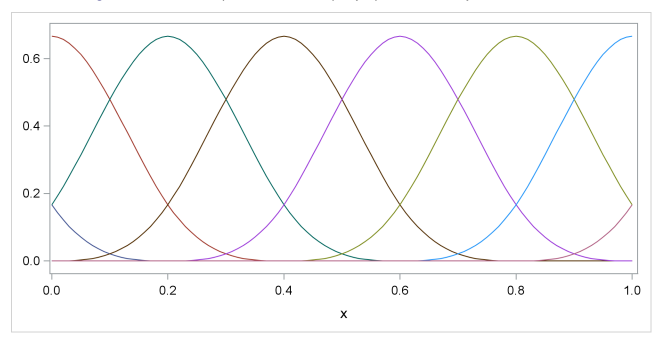


Figure 19.4 Cubic B-Spline Basis with Equally Spaced Boundary and Interior Knots

You can find details about this construction in Hastie, Tibshirani, and Friedman (2001).

# **Natural Cubic Spline Basis**

Natural cubic splines are cubic splines with the additional restriction that the splines are required to be linear beyond the extreme knots. Some authors prefer the terminology "restricted cubic splines" to "natural cubic splines." The space of unrestricted cubic splines on n knots has the dimension n + 4. Imposing the restrictions that the cubic polynomials beyond the first and last knot reduce to linear polynomials reduces the number of degrees of freedom by 4, and so a basis for the natural cubic splines consists of n functions. Starting from the truncated power function basis for the unrestricted cubic splines, you can obtain a reduced basis by imposing linearity constraints. You can find details about this construction in Hastie, Tibshirani, and Friedman (2001). Figure 19.5 shows this natural cubic spline basis defined on [0, 1] with four equally spaced internal knots at 0.2, 0.4, 0.6, and 0.8. Note that this basis consists of four basis functions that are all linear beyond the extreme knots at 0.2 and 0.8.

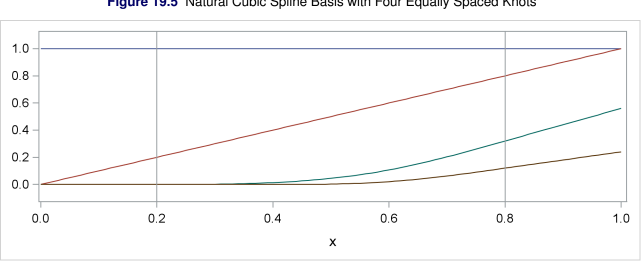


Figure 19.5 Natural Cubic Spline Basis with Four Equally Spaced Knots

# **EFFECTPLOT Statement**

This statement applies to the following SAS/STAT procedures: GENMOD, LOGISTIC, ORTHOREG, and PLM. It also applies to the RELIABILITY procedure in SAS/QC software.

The EFFECTPLOT statement produces a display (effect plot) of a complex fitted model and provides options for changing and enhancing the display. One simple effect plot is the display for a linear regression of the response Y on a single predictor X: the regression line is drawn with the predicted response on the Y axis and the covariate on the X axis. The regression line can be enhanced by displaying the observations and adding confidence and prediction limits. When your model is more complicated—with more continuous and categorical covariates, nestings and interactions, and link functions—the effect plots display the behavior of some covariates over their ranges while holding other covariates at some fixed values; this can enable easier interpretation and explanation of the resulting model.

By default, a single plot is produced based on the type of response variable and the number of continuous and classification covariates in the model. You can also specify options to do the following:

- select the variables to display in the plots
- produce multiple plots based on the following: the levels of classification covariates; the minimum, maximum, mean, or middle (midrange) value of continuous covariates; and specified values of the covariates
- specify different fixed values for continuous and classification covariates that are not displayed in the plot
- panel and unpanel plots
- select variables to slice or group by
- display (or remove from display) observations and confidence limits

# Syntax: EFFECTPLOT Statement

**EFFECTPLOT** < plot-type < (plot-definition-options) >> </ options>;

The available plot-types and their plot-definition-options are described in Table 19.13. Table 19.15 lists the options that can be specified after a slash (/) for any plot-type, and Table 19.16 lists additional options that enhance specific plot-types. Full descriptions of the plot-definition-options and the other options are provided in the section "Dictionary of Options" on page 418.

 Table 19.13
 Plot-Types and Plot-Definition-Options

Plot-Type and Description	Plot-Definition-Options
BOX Displays a box plot of continuous response data at each level of a CLASS effect, with predicted values superimposed and connected by a line. This is an alternative to the INTERACTION plot-type.	PLOTBY= variable or CLASS effect X= CLASS variable or effect
<b>CONTOUR</b> Displays a contour plot of predicted values against two continuous covariates	PLOTBY= variable or CLASS effect X= continuous variable Y= continuous variable
FIT Displays a curve of predicted values versus a continuous variable	PLOTBY= variable or CLASS effect X= continuous variable
INTERACTION Displays a plot of predicted values (possibly with error bars) versus the levels of a CLASS effect. The predicted values are connected with lines and can be grouped by the levels of another CLASS effect.	PLOTBY= variable or CLASS effect SLICEBY= variable or CLASS effect X= CLASS variable or effect
MOSAIC Displays a mosaic plot of predicted values by using up to three CLASS effects	PLOTBY= variable or CLASS effect X= CLASS effects
SLICEFIT Displays a curve of predicted values versus a continuous variable, grouped by the levels of a CLASS effect	PLOTBY= variable or CLASS effect SLICEBY= variable or CLASS effect X= continuous variable

By default, a single plot is produced based on the type of response variable and the number of continuous and classification covariates in the model, as shown in Table 19.14. If you have a polytomous response model, then the response variable is treated as the grouping classification variable in this table. If your model does not fit into Table 19.14, then a default plot is not produced; however, specifying the *plot-type* argument displays a plot with the extra continuous covariates fixed at their mean values and the extra classification covariates fixed at their reference levels.

**Table 19.14** Default *Plot-Types* 

Number of Covariates		Type of Response Variable	
Classification	Continuous	<b>Continuous or Binary</b>	Polytomous
1	0	INTERACTION	INTERACTION with groups
2	0	INTERACTION with groups	None
0	1	FIT	SLICEFIT
0	2	CONTOUR	None
1	1	SLICEFIT	None

**Table 19.15** Available *Options* for All *Plot-Types* 

AT< args>			NROWS=ab	PREDLABEL=
ATLEN=	INDIVIDUAL <sup>a</sup>		OBS<(options)>	UNPACK
ATORDER=	$LINK^b$	$NOOBS^{ab}$	PLOTBYLEN=	

<sup>&</sup>lt;sup>a</sup> Not available for the BOX *plot-type*.

**NOTE:** If your model contains an offset variable and the MOFF option is not specified or not valid, then the predicted values are computed only at the observations. In this case, the FIT and SLICEFIT *plot-types* display scatter plots of the predicted values, the CONTOUR *plot-type* displays the residuals against two continuous covariates but with no fitted surface, the INTERACTION *plot-type* does not connect the predicted values with lines, and the BOX *plot-type* is unchanged.

Table 19.16 Additional Options for Each Plot-Type

Plot-Type	Options
BOX	CLUSTER, CONNECT, NOCLUSTER, NOCONNECT, YRANGE=
CONTOUR	EXTEND=, GRIDSIZE=
FIT	ALPHA=, EXTEND=, GRIDSIZE=, NOCLI, NOCLM, NOLIMITS, SMOOTH, YRANGE=
INTERACTION	ALPHA=, CLI, CLM, CLUSTER, CONNECT, LIMITS, NOCLUSTER, NOCONNECT, POLYBAR, YRANGE=
MOSAIC	ADDCELL, BIN, EQUAL, NOBORDER, TYPE=
SLICEFIT	ALPHA=, CLI, CLM, EXTEND=, GRIDSIZE=, LIMITS, YRANGE=

# **Dictionary of Options**

You can specify the following EFFECTPLOT options.

#### ADDCELL<=value>

adds *value* to the weight of every cell in the MOSAIC *plot-type*. You can specify *value* as any nonnegative number. If you do not specify a *value*, then *value*=0.5. This enables you to add some dimension to zero frequency cells.

## ALPHA=value

specifies the significance level,  $0 \le value \le 1$ , for producing 100(1 - value/2)% prediction and confidence limits. By default, value=0.05.

<sup>&</sup>lt;sup>b</sup> Not available for the MOSAIC *plot-type*.

AT < contopt > < classopt > < variable1< (CODED) >= varopt < variable2< (CODED) >= varopt... >> where contopt= MEAN | MIN | MAX | MIDRANGE classopt= ALL | REF

varopt= contopts | number-list | classopts | 'class-level'... 'class-level' specifies values at which to fix continuous and CLASS variables when they are not used in X=, Y=, SLICEBY=, or PLOTBY= effects. The contopt keyword fixes continuous variables at their mean, minimum, maximum, or midrange=  $\frac{1}{2}$ (minimum + maximum); the default is to use the mean. The classopt keyword either fixes a CLASS variable at its reference (last) level or indicates that all levels of the CLASS variable should be processed; the default is to use the reference level. The varopt values enable you to specify contopt and classopt keywords or to specify lists of numbers or class levels. You can specify a CLASS variable only once in the AT specification, but you can specify a continuous variable multiple times; for example, the following syntax is valid when X is a continuous variable:

effectplot / at(x=min max x=0 to 2 by 1 x=2 5 7);

Duplicate AT values are suppressed, so the last X=2 value is ignored.

You can also specify *coded plug-in values* for CLASS variable levels when computing the predicted values  $\mathbf{x}'\boldsymbol{\beta}$ . For example, suppose a CLASS variable A with two levels= $\{0,1\}$  is in the model. Then instead of using the coding for A in the x vector by specifying  $\mathbf{AT}$  ( $\mathbf{A}=\mathbf{all}$ ),  $\mathbf{AT}$  ( $\mathbf{A}=\mathbf{ref}$ ) or  $\mathbf{AT}$  ( $\mathbf{A}=\mathbf{ref}$ )

The coded plug-in values are distributed through the rest of the model effects in the following fashion. If a variable is nested within a plug-in variable, then its coding is multiplied by the plug-in value for the level it is nested in. If a variable interacts with a plug-in variable, its coding is multiplied by the appropriate plug-in value for the level it is interacting with. Lag, multimember, polynomial, and spline constructed effects are affected only by interactions and nestings. If the plug-in variable is part of a collection effect, then its values are replaced by the plug-in values; collection effects are also affected by interactions and nestings.

The AT levels are used for computing the predicted values. If the OBS option is also specified, then all observations are still displayed in all the plots. For example, if you specify the options AT (A='1') OBS, then the fitted values are computed by using A=1, but all the observations are displayed with their predicted values computed at their observed level of A. If you want to display only a subset of the observations based on the levels of a CLASS variable, then you must specify either the PLOTBY= option or the OBS(BYAT) option.

#### ATLEN=n

specifies the maximum length ( $1 \le n \le 256$ ) of the levels of the AT variables that are displayed in footnotes and headers. By default, up to 256 characters of the CLASS levels are displayed, and the continuous AT levels are displayed with a BEST format that has a width greater than or equal to 5,

which distinguishes each level. **CAUTION:** If the levels of your AT variables are not unique when the first n characters are displayed, then the levels are combined in the plots but not in the underlying computations. Also, at most n characters for continuous AT variables are displayed.

### ATORDER=ASCENDING | DESCENDING

uses the AT values for continuous variables in ascending or descending order as specified. By default, values are used in the order of their first appearance in the AT option.

#### BIN

displays the statistic for the MOSAIC *plot-type* with a discrete coloring scheme.

# CLI

displays normal (Wald) prediction limits. This option is available only for normal distributions with identity links. If your model is from a Bayesian analysis, then sampling-based intervals are computed; for more information, see the section "Analysis Based on Posterior Estimates" on page 6180 in Chapter 75, "The PLM Procedure."

#### **CLM**

displays confidence limits. These are computed as the normal (Wald) confidence limits for the linear predictor, and if the ILINK option is specified, the limits are also back-transformed by the inverse link function. If your model is from a Bayesian analysis, then sampling-based intervals are computed; for more information, see the section "Analysis Based on Posterior Estimates" on page 6180 in Chapter 75, "The PLM Procedure."

# CLUSTER < = percent >

modifies the BOX and INTERACTION plot-types by displaying the levels of the SLICEBY= effect side by side. You can specify *percent* as a percentage of half the distance between X levels. The percent value must be between 0.1 and 1; the default percent depends on the number of X levels, the number of SLICEBY levels, and the number of PLOTBY levels for INTERACTION plot-types. You can remove default clustering by specifying the NOCLUSTER option.

# **CONNECT**

modifies the BOX and INTERACTION plot-types by connecting the predicted values with a line. You can remove default connecting lines by specifying the NOCONNECT option.

# **EQUAL**

causes every cell in the MOSAIC plot-type to have the same dimensions.

## **EXTEND=DATA** | value

extends continuous covariate axes by  $value \times \frac{1}{2} range$  in both directions, where range is the range of the X axis. Specifying the DATA keyword displays curves to the range of the data within the appropriate SLICEBY=, PLOTBY=, and AT level. For the CONTOUR plot-type, value=0.05 by default; other plot-types set the default value to 0. When constructed effects are present, only the EXTEND=DATA option is available.

#### **GRIDSIZE=**n

specifies the resolution of curves by computing the predicted values at n equally spaced values on the X axis and specifies the resolution of surfaces by computing the predicted values on an  $n \times n$  grid of points. Default values are n = 200 for curves and bands, n = 50 for surfaces, and n = 2 for lines. If results of a Bayesian or bootstrap analysis are being displayed, then the defaults are n = 500000/B, where B is the number of samples, the upper limit is equal to the usual defaults, and the lower limit is equal to 20.

#### **ILINK**

displays the fit on the scale of the inverse link function. In particular, the results are displayed on the probability scale for logistic regression. By default, a procedure displays the fit on either the link or inverse link scale.

#### **INDIVIDUAL**

displays individual probabilities for polytomous response models with cumulative links on the scale of the inverse link function. This option is not available when the LINK option is specified, and confidence limits are not available when you specify this option.

#### **LIMITS**

invokes the CLI and CLM options.

#### LINK

displays the fit on the scale of the link function—that is, the linear predictor. Probabilities or observed proportions near 0 and 1 are transformed to  $\pm 20$ . By default, a procedure displays the fit on either the link or inverse link scale.

#### **MOFF**

moves the offset for a Poisson regression model to the response side of the equation. If the ILINK option is also specified, then the rate is displayed on the Y axis; if the LINK option is also specified, then the log of the rate is displayed on the Y axis. Without the MOFF option, the predicted values are computed and displayed only for the observations.

#### NCOLS=n

specifies the maximum number of columns in a paneled plot. This option is not available with the BOX *plot-type*.

The default choice of NROWS= and NCOLS= is based on the number of PLOTBY= and AT levels. If only one plot is displayed in a panel, then NROWS=1 and NCOLS=1 and the plots are produced as if you specified only the UNPACK option. If only two plots are displayed in a panel, then NROWS=1 and NCOLS=2. For all other cases, a  $2\times2$ ,  $2\times3$ , or  $3\times3$  panel is chosen based on how much of the last panel is used, with ties going to the larger panels. For example, if 14 plots are being created, then this requires either four  $2\times2$  panels with 50% of the last panel filled, three  $2\times3$  panels with 33% of the last panel filled, or two  $3\times3$  panels with 55% of the last panel filled; in this case, the  $3\times3$  panels are chosen.

If you specify both the NROWS= and NCOLS= options, then those are the values used. However, if you specify only one of the options but have fewer plots, then the panel size is reduced; for example, if you specify NROWS=6 but have only four plots, then a plot that has four rows and one column is produced.

#### **NOBORDER**

removes the border from the cells in the MOSAIC *plot-type*. Otherwise, the color of the cells that were not observed in the data set is hidden by the border.

### **NOCLI**

suppresses the prediction limits.

#### **NOCLM**

suppresses the confidence limits.

### **NOCLUSTER**

modifies the BOX and INTERACTION *plot-types* by preventing the side-by-side display of the levels of the SLICEBY= effect.

#### **NOCONNECT**

modifies the BOX and INTERACTION *plot-types* by suppressing the line that connects the predicted values.

## **NOLIMITS**

invokes the NOCLI and NOCLM options.

#### **NOOBS**

suppresses the display of observations and overrides the specification of the OBS= option.

#### NROWS=n

specifies the maximum number of rows in a paneled plot. This option is not available with the BOX *plot-type*. For more information, see the NCOLS= option.

## OBS<(obs-options)>

displays observations in the effect plots. An input data set is required; hence the OBS option is not available with PROC PLM. The OBS option is overridden by the NOOBS option. When the ILINK option is specified with binary response variables, then either the observed proportions or a coded value of the response is displayed. For polytomous response variables, the observed values are overlaid onto the fitted curves unless the LOCATION= option is specified. Whether or not observations are displayed by default depends on the procedure. If the PLOTBY= option is specified, then the observations that are displayed in each plot are from the corresponding PLOTBY= level for classification effects; for continuous effects, all observations are displayed in every plot.

You can specify the following *obs-options*:

# **BYAT**

subsets the observations by AT level and by the PLOTBY= level. If you specify the PLOTBY= option without specifying this option, the observations are displayed in the plots that correspond to their PLOTBY= level without regard to any classification variables specified in the AT option. However, for FIT *plot-types* a distance can be computed and displayed (for more information, see the DISTANCE option). This option is ignored when there are no AT variables.

# CDISPLAY=NONE | OUTLINE | GRADIENT | OUTLINEGRADIENT

controls the display of observations in contour plots. The keyword OUTLINE displays the observations as circles, GRADIENT displays gradient-colored dots, OUTLINEGRADIENT displays gradient-filled-circles, and NONE suppresses the display of the observations. The default is CDISPLAY=OUTLINEGRADIENT.

# CGRADIENT=RESIDUAL | DEPENDENT

specifies what the gradient shading of the observed values in the CONTOUR *plot-type* represents. The RESIDUAL keyword shades the observations by the raw residual value and displays the fitted surface as a line contour plot. The DEPENDENT keyword shades the observations by the response variable value and displays the fitted surface as a contour shaded on the same scale. The default is CGRADIENT=DEPENDENT.

## **DEPTH**=depth

specifies the number of overlapping observations that can be distinguished by adjusting their transparency; you can specify  $1 \le depth \le 100$ . By default, DEPTH=1. The DEPTH= option is available with the FIT, SLICEFIT, and INTERACTION *plot-types*.

# **DISTANCE**

displays observations in FIT *plot-types* with a color gradient that indicates how far the observation is from the AT and PLOTBY= level. This option is ignored unless an AT or PLOTBY= option is specified.

The distance is computed as the square root of the following number: for each continuous AT and PLOTBY= variable, add the square of the difference from the observed value divided by the range of the variable; for each CLASS AT and PLOTBY= variable, add 1 if the CLASS levels are different. Thus the largest possible distance is the square root of the number of AT and PLOTBY= variables. Observations at zero distance are displayed by using the darkest color, and the color fades as the distance increases.

The unpacked panels compute the maximum distance within each panel and hence do not use the same gradient across all panels. Also, the PANELS *panel-type* computes the maximum distance within each PLOTBY= level, so a different gradient is used for each PLOTBY= level. All other *panel-types* compute the maximum distance across all observations and therefore use the same gradient in every plot.

#### **FITATCLASS**

computes fitted values only for class levels that are observed in the data set. This option is ignored when the GLM parameterization is used.

### **FRINGE**

displays observations in a fringe (rug) plot at the bottom of the plot. This option is available only with the FIT and SLICEFIT *plot-types*.

### JITTER<(jitter-options)>

shifts (*jitters*) the observations. By default, the jittering in the X direction is achieved by adding a random number that is generated according to a normal distribution with mean=0 and standard deviation= x-jitter/2 and truncating at  $\pm x$ -jitter, where x-jitter=0.01 times the range of the X axis; the jittering in the Y direction is performed independently but in the same fashion. The JITTER option is not available with the BOX plot-type. You can specify the following jitter-options:

- **FACTOR**=*factor* sets the jitter to *factor* times the range of the axis, and jitters in both the X and Y directions. You can specify  $0 \le factor \le 1$ .
- **SEED**=*seed* specifies an integer to use as the initial seed for the random number generator. If you do not specify a seed, or if you specify a value less than or equal to zero, then the time of day from the computer clock is used to generate an initial seed.
- **X**=x-jitter sets the jitter to x-jitter for the X direction; the jitter in the Y direction is assumed to be 0 unless the Y= option is also specified. You can specify x-jitter  $\geq 0$ . The X= option is not available for the INTERACTION plot-type. This option is ignored if the FACTOR= option is also specified.
- **Y**=*y*-jitter sets the jitter to *y*-jitter for the Y direction; the jitter in the X direction is assumed to be 0 unless the X= option is also specified. You can specify *y*-jitter  $\geq$  0. This option is ignored if the FACTOR= option is also specified.

#### LABEL< =OBS>

labels markers with their observation number.

#### **LOCATION**=location

specifies where the observed values for polytomous response models are displayed when the SLICEBY= variable is the response. This option is available only with the SLICEFIT and INTERACTION *plot-types*. The observations are always displayed at their appropriate X-axis value, but their Y-axis location can depend on the specification of the YRANGE= option or on the minimum and maximum computed predicted values in addition to the specified *location*. You can specify the following *locations*:

**BOTTOM**<=factor> displays the first response level at the minimum predicted value, and displays succeeding response levels above the first level at  $factor \times range$  intervals, where range is the range of the predicted values. You can specify  $0 \le factor \le 1$ , but the largest usable value, which corresponds to LOCATION=SPREAD, is  $factor = \frac{1}{k}$ , where k+1 is the number of response levels that are displayed. By default, factor = 0.03.

**CURVE** displays the observations for polytomous response models at their predicted values. For displays on the LINK scale, the reference level is displayed at the maximum value. This method is the default.

**FIRST** displays the observations for a response level at the first displayed predicted value for that response level.

**MAX** displays the observations for a response level at the maximum displayed predicted value for that response level.

**MIDDLE** displays the observations for a response level at the middle of the displayed predicted values for that response level.

**MIN** displays the observations for a response level at the minimum displayed predicted value for that response level.

**SPREAD** displays the observations with the response levels evenly spread across the Y axis.

**TOP**<=factor> displays the last response level at the maximum predicted value, and displays preceding response levels below the last level at  $factor \times range$  intervals, where range is the range of the predicted values. You can specify  $0 \le factor \le 1$ , but the largest usable value, which corresponds to LOCATION=SPREAD, is  $factor = \frac{1}{k}$ , where k+1 is the number of response levels that are displayed. By default, factor = 0.03.

# PLOTBY<(panel-type)>=effect<=numeric-list>

specifies a variable or CLASS effect at whose levels the predicted values are computed and the plots are displayed. You can specify the response variable as the *effect* for polytomous response models. The *panel-type* argument specifies the method in which the plots are grouped for the display. You can specify the following *panel-types*:

# **COLUMNS**

specifies that the columns within each panel correspond to different levels of the PLOTBY= effect and hence the rows correspond to different AT levels.

#### **PACK**

specifies that plots be displayed in the panels as they are produced, with no control over the placement of the PLOTBY= and AT levels.

# **PANELS | LEVELS**

specifies that each level of the PLOTBY= effect begin a new panel of plots and the AT levels define the plots within the panels.

#### **ROWS**

specifies that the rows within each panel correspond to different levels of the PLOTBY= effect and hence the columns correspond to different AT levels.

This option is ignored with the BOX *plot-type*; box plots are always displayed in an unpacked fashion, grouped by the PLOTBY= and AT levels. If you specify a continuous variable as the *effect*, then either you can specify a *numeric-list* of values at which to display that variable or, by default, five equally spaced values from the minimum variable value to its maximum are displayed.

The default *panel-type* is based on the number of PLOTBY= and AT levels, as shown in the following table.

Number of PLOTBY= Levels	Number of AT Levels	Resulting panel-type
1	1	(UNPACK)
>1	1	PACK
1	>1	PACK
2	>1	ROWS
3	>1	COLUMNS
>3	>1	PANELS

The default dimensions of the panels are also based on the number of PLOTBY= and AT levels; for more information, see the NCOLS= option.

Specification of the *panel-type* is honored except in the following cases. If you specify a *panel-type* but produce only one plot, specify the NROWS=1 and NCOLS=1 options, or specify the UNPACK option, then the plots are produced as if you specified only the UNPACK option. If you specify the PANELS *panel-type* with only one AT level, then the plots are produced with the UNPACK option. However, if you specify the PANELS *panel-type* but the PLOTBY= effect has only one level, then the *panel-type* is changed to PACK.

### **PLOTBYLEN**=n

specifies the maximum length ( $1 \le n \le 256$ ) of the levels of the PLOTBY= variables, which are displayed in footnotes and headers. By default, up to 256 characters of the CLASS levels are displayed. **CAUTION:** If the levels of your PLOTBY= variables are not unique when the first *n* characters are displayed, then the levels are combined in the plots but not in the underlying computations.

## **POLYBAR**

displays polytomous response data as a stacked histogram whose bar heights are defined by the individual predicted value. Your response variable must be the *effect* that is specified in the SLICEBY= option. If you specify the INDIVIDUAL option, then the histogram bars are displayed side by side. If you specify the CLM option, then error bars are displayed on the side-by-side histogram bars.

#### PREDLABEL='label'

specifies a label to be displayed on the Y axis. The default Y-axis label is determined by your model. For the CONTOUR plot-type, this option changes the title to "label for Y."

#### SHOWCLEGEND

displays the gradient legend for the CONTOUR plot-type. This option has no effect when the OBS(CGRADIENT=RESIDUAL) option is also specified.

#### SLICEBY=NONE | effect< =numeric-list>

displays the fitted values at the different levels of the specified variable or CLASS effect. You can specify the response variable as the *effect* for polytomous response models. Use this option to modify the SLICEFIT, INTERACTION, and BOX plot-types. If you specify a continuous variable as the effect, then either you can specify a numeric-list of values at which to display that variable or, by default, five equally spaced values from the minimum variable value to its maximum are displayed. The NONE keyword is available for preventing the INTERACTION plot-type from slicing by a second classification covariate. The SLICEBY=NONE option is not available for the SLICEFIT plot-type, because that is the same as the FIT plot-type. The BOX plot-type accepts only classification effects.

#### **SMOOTH**

overlays a loess smooth on the FIT plot-type for models that have only one continuous predictor. This option is not available for binary or polytomous response models.

## TYPE=PREDICTED | PARQUET | GOF

specifies the type of display for the MOSAIC plot-types. For effects that are specified as in the X= option, the TYPE=PREDICTED and TYPE=GOF mosaic plots create cells by dividing the X axis proportional to the total weight in each level of the x-effect, then dividing the Y axis according to the weight in each level of the *y-effect* within the *x-effect* levels, and dividing the X2 axis according to the weight in each level of the x2-effect within the x-effect and y-effect levels. The TYPE=PARQUET plot uses the predicted probabilities instead of the weights to determine the dimensions of the cells.

The default TYPE=PREDICTED mosaic plot colors the cells according to their predicted values (probabilities for binary and polytomous response models) computed at the AT and PLOTBY levels. The TYPE=GOF plot displays the Pearson goodness-of-fit statistic as in Friendly (2000), with the expected value computed at the AT and PLOTBY levels. For a cell  $i_v$  defined by the axis levels, the PLOTBY and AT levels, and the response level y, let  $W_{i_y}$  be the sum of all the weights of the observations in that cell, let  $W_i = \sum_y W_{i_y}$  be the sum of the weights across all response levels, and let  $\hat{Y}_y$  be the predicted response for that cell, where y is the event level for binary response models, and  $\hat{Y}_y = \Pr(Y = y | i_y)$  for binary and multinomial models. Then the Pearson goodness-of-fit statistic is computed as

$$\frac{W_{i_y} - W_i \, \hat{Y}}{\sqrt{W_i \, \hat{Y}}}$$

The TYPE=GOF plot is not available when you have continuous covariates in the model. The TYPE=PARQUET plot shades the cells with their observed weights and is available only with binary or polytomous response data.

#### **UNPACK**

suppresses paneling. By default, multiple plots can appear in some output *panels*. Specify UNPACK to display each plot separately.

## $X=effect \mid (x-effect < y-effect < x2-effect >>)$

specifies values to display on the X axis. For the BOX and INTERACTION plot-types, effect can be a CLASS effect in the MODEL statement. For the FIT, SLICEFIT, and CONTOUR plot-types, effect can be any continuous variable in the model. For the MOSAIC plot-types, you can specify CLASS effects (or the response variable if you have a multinomial model) as the effect or x-effect to display on the X axis, as the y-effect to display on the Y axis, and as the x2-effect to display on the X2 (upper) axis.

#### Y=args

specifies values to display on the Y axis for the CONTOUR *plot-type*. The Y= argument can be any continuous variable in the model.

# **YRANGE=CLIP** | (< min >< ,max >)

displays the predicted values on the Y axis in the range [min,max]. The YRANGE=CLIP option has the same effect as specifying the minimum predicted value as min and the maximum predicted value as max. The axis might extend beyond your specified values. By default, when the Y axis displays predicted probabilities, the entire Y axis, [0,1], is displayed. This option is useful if your predicted probabilities are all contained in some subset of this range. This option is not available with the CONTOUR plot-type.

# **ODS Graphics: EFFECTPLOT Statement**

To produce the EFFECTPLOT displays, ODS Graphics must be enabled. For more information about ODS Graphics, see Chapter 21, "Statistical Graphics Using ODS." The available graph names are provided in Table 19.17.

Table 19.17 Graphs Produced by the EFFECTPLOT Statement

<b>ODS Graph Name</b>	Plot Description
BoxFitPlot	A box plot of the responses at each level of one classification effect, overlaid
	with a plot of the predicted values
ContourFitPlot	A contour plot of the fitted surface against two continuous covariates
ContourFitPanel	A panel of ContourFitPlots
FitPlot	A curve of the predicted values plotted against one continuous covariate
FitPanel	A panel of FitPlots
InteractionPlot	A plot of the predicted values (connected by a line) against one classification
	effect, possibly for each level of a second classification effect
InteractionPanel	A panel of InteractionPlots
MosaicFitPlot	A mosaic plot of the predicted values categorized by one to three classifica-
	tion effects
SliceFitPlot	A curve of the predicted values against one continuous covariate for each
	level of a second classification covariate
SliceFitPanel	A panel of SliceFitPlots

# **Examples: EFFECTPLOT Statement**

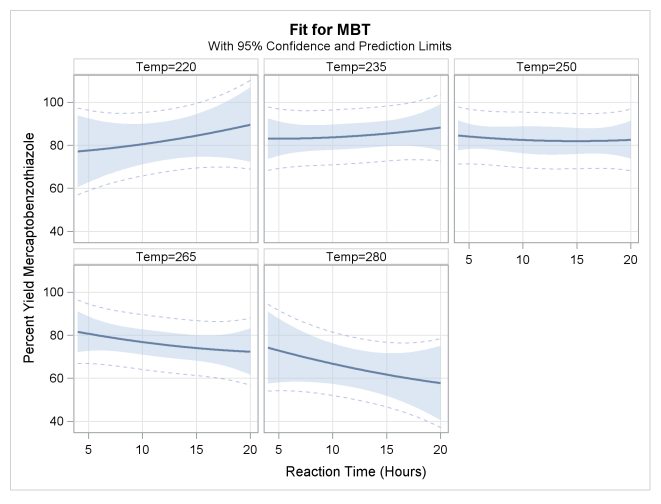
# **Example 19.1: A Saddle Surface**

Myers (1976) analyzes an experiment reported by Frankel (1961) that is aimed at maximizing the yield of mercaptobenzothiazole (MBT) by varying processing time and temperature. Myers uses a two-factor model in which the estimated surface does not have a unique optimum. The objective is to find the settings of time and temperature in the processing of the chemical that maximize the yield. The following statements create the data set d:

In the following statements, the ORTHOREG procedure fits a response surface regression model to the data and uses the EFFECTPLOT statement to create a slice of the response surface. The FIT *plot-type* requests plots of the predicted yield against the Time variable, and the PLOTBY= option specifies that the Temp variable is fixed at five equally spaced values so that five fitted regression curves are displayed in Output 19.1.1.

```
ods graphics on;
proc orthoreg data=d;
   model MBT=Time|Time|Temp|Temp@2;
   effectplot fit(x=time plotby=temp);
run;
ods graphics off;
```

The displays in Output 19.1.1 show that the slope of the surface changes as the temperature increases.

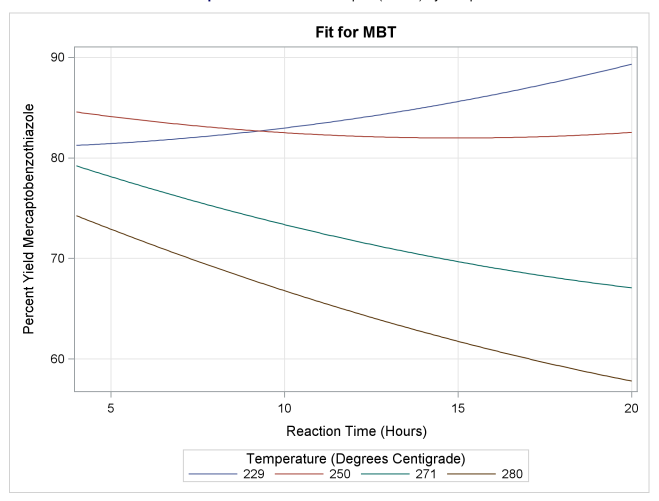


Output 19.1.1 Panel of Fit Plots

It might be more informative to see these results in one graphic, so the following statements specify the SLICEFIT *plot-type* to overlay plots of the predicted yield versus time, fixed at several values of temperature. In this case, the SLICEBY= option is specified to explicitly use the same four temperatures as used in the experiment.

```
ods graphics on;
proc orthoreg data=d;
  model MBT=Time|Time|Temp|Temp@2;
  effectplot slicefit(x=time sliceby=temp=229 250 271 280);
run;
ods graphics off;
```

Output 19.1.2 shows that to optimize the yield you should choose either low temperatures and long times or high temperatures and short times.

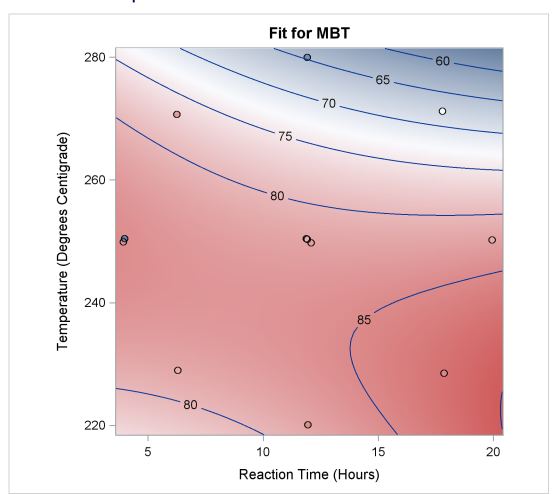


Output 19.1.2 Fit Plot Grouped (Sliced) by Temp

Another plot might explain the reason for these conflicting results more clearly. The following statements produce the default EFFECTPLOT statement display, enhanced by the OBS(JITTER) option to jitter the observations so that you can see the replicated points:

```
ods graphics on;
proc orthoreg data=d;
  model MBT=Time|Time|Temp|Temp@2;
  effectplot / obs(jitter(seed=39393));
run;
ods graphics off;
```

Output 19.1.3 shows that the reason for the changing slopes is that the surface is at a saddle point. This surface does not have an optimum point.



Output 19.1.3 Contour Fit Plot with Jittered Observations

# **Example 19.2: Unbalanced Two-Way ANOVA**

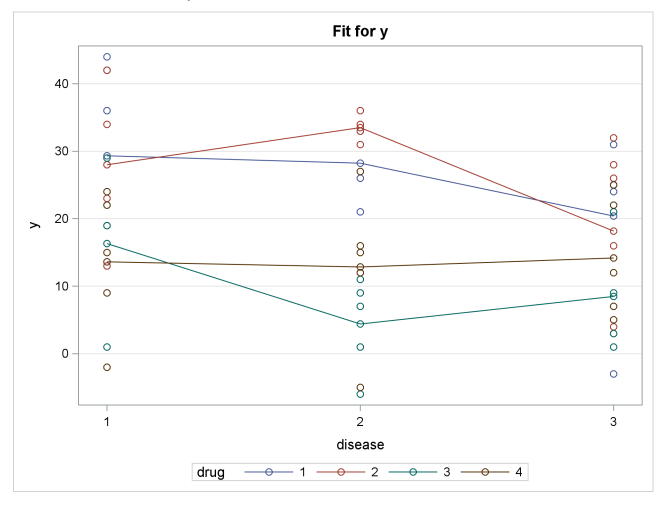
This example uses data from Kutner (1974, p. 98) to illustrate a two-way analysis of variance. The original data source is Afifi and Azen (1972, p. 166). The following statements create the data set a:

```
data a:
  input drug disease @;
  do i=1 to 6;
     input y @;
      output;
  end;
  datalines;
1 1 42 44 36 13 19 22
1 2 33 . 26 . 33 21
1 3 31 -3 . 25 25 24
2 1 28 . 23 34 42 13
2 2 . 34 33 31 . 36
2 3 3 26 28 32 4 16
3 1
    . .
         1 29
                . 19
3 2 . 11 9 7 1 -6
3 3 21 1 .
             9 3 .
4 1 24 . 9 22 -2 15
4 2 27 12 12 -5 16 15
4 3 22 7 25 5 12 .
```

In the following statements, PROC GENMOD fits two classification variables and their interaction to Y. The first EFFECTPLOT statement displays the default graphic, which plots the predicted values against Disease for each of the three Drug levels. The OBS option also displays the observations in the plot. The second EFFECTPLOT statement modifies the default to plot the predicted values against Drug for each of the three Disease levels. The CLM option is specified to produce 95% confidence bars for the means.

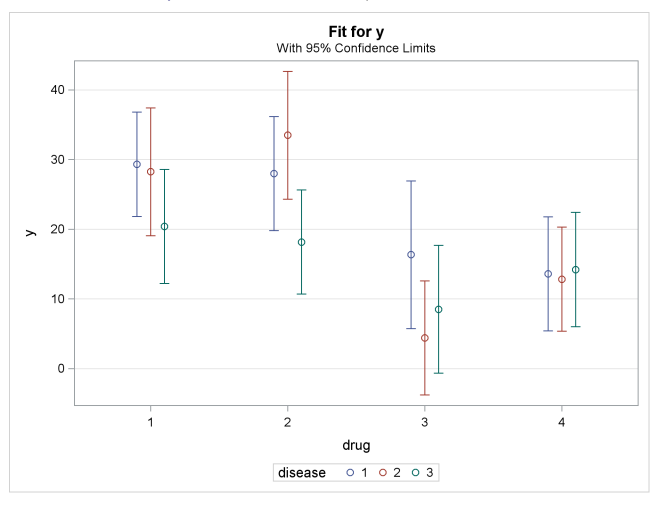
```
ods graphics on;
proc genmod data=a;
   class drug disease;
   model y=disease drug disease*drug / d=n;
   effectplot / obs;
   effectplot interaction(sliceby=disease) / clm;
run;
ods graphics off;
```

In Output 19.2.1, the default interaction plot is produced, and the observations are also displayed. From this plot, you can compare the performance of the drugs for a given disease. The predicted values are connected with a line to provide something for your eye to follow; obviously a line has no intrinsic meaning in this graphic. Drugs 3 and 4 are consistently outperformed by the first two drugs.



Output 19.2.1 Interaction Plot: Default with Observations

By default, the first classification variable is displayed on the X axis and the second classification variable is used for grouping. Specifying the SLICEBY=DISEASE option in the second EFFECTPLOT statement does the reverse: it displays the classification variable that has the most levels on the X axis, and slices by fewer levels, resulting in a more readable display. Output 19.2.2 shows how well a given drug performs for each disease.



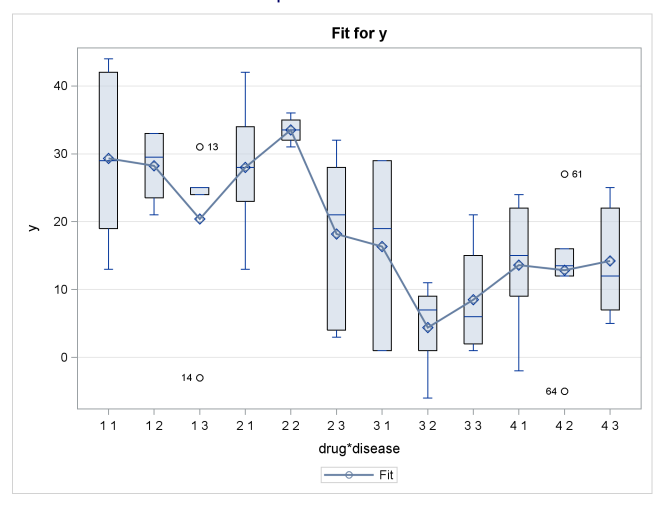
Output 19.2.2 Interaction Plot with Specified SLICEBY= Effect

In the following statements, the BOX *plot-type* is requested to display box plots of the predictions by each combination of drug and disease. The second EFFECTPLOT statement displays the same information by using an INTERACTION *plot-type* and specifies the OBS option to display the individual observations. The third EFFECTPLOT statement displays the predictions in a MOSAIC plot. The fourth EFFECTPLOT statement creates an interaction plot of predictions versus drug for each of the Disease levels and displays them in a panel.

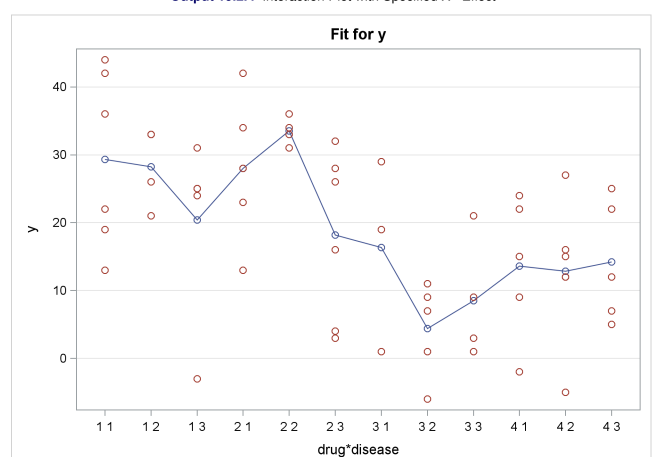
```
ods graphics on;
proc genmod data=a;
   class drug disease;
   model y=drug disease drug*disease / d=n;
   effectplot box;
   effectplot interaction(x=drug*disease) / obs;
   effectplot mosaic;
   effectplot interaction(plotby=disease);
run;
ods graphics off;
```

In the box plot in Output 19.2.3, the predicted values are displayed as circles; they coincide with the mean of the data at each level, all of which are displayed as diamonds. The predicted values are again connected by lines. It is difficult to draw any conclusions from this graphic.

Output 19.2.3 Box Fit Plot



Output 19.2.4 shows the interaction plot at every combination of Drug and Disease. This plot is identical to the box plot in Output 19.2.3, except the boxes are replaced by the actual observations. Again, it is difficult to see any pattern in the plot.

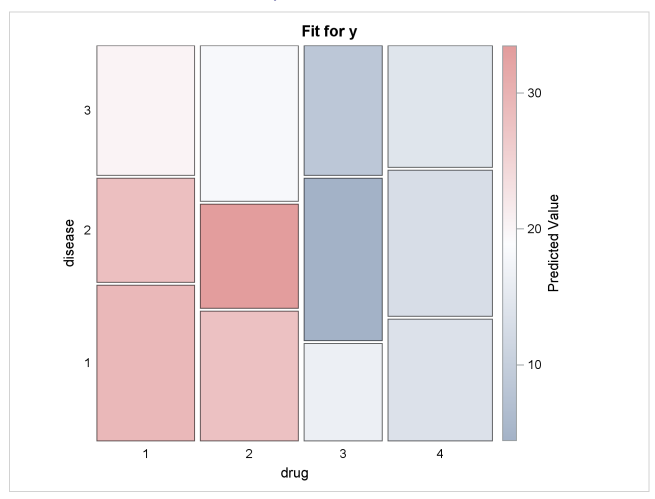


Observed — Fit

Output 19.2.4 Interaction Plot with Specified X= Effect

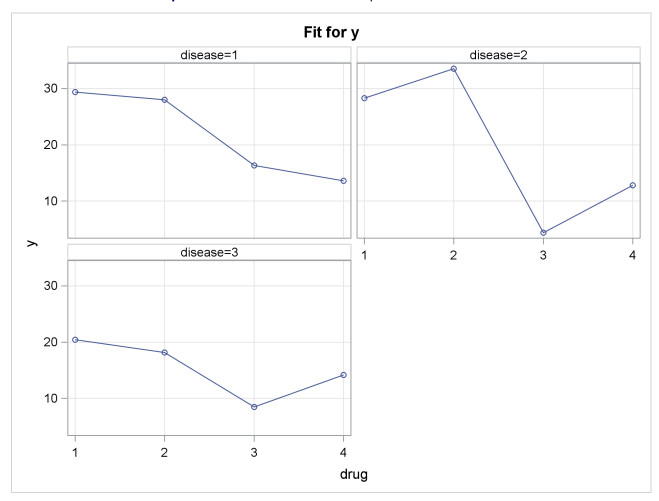
Output 19.2.5 displays the mosaic plot. You can see that drugs 1 and 2 consistently outperform the other two drugs.

Output 19.2.5 Mosaic Plot



Output 19.2.6 groups the observations by Disease, and for each disease displays the effectiveness of the four drugs in a panel of plots.

Output 19.2.6 Interaction Plot with Specified PLOTBY= Effect



# **Example 19.3: Logistic Regression**

Consider a study of the analgesic effects of treatments on elderly patients with neuralgia. Two test treatments and a placebo are compared. The response variable is whether the patient reported pain or not. Researchers recorded the age and gender of 60 patients and the duration of complaint before the treatment began. The following DATA step creates the data set Neuralgia:

```
data Neuralgia;
   input Treatment $ Sex $ Age Duration Pain $ @@;
   datalines;
      68
                         74
                                                30
                                                    No
           1
             No
                   B M
                             16
                                 No
                                      P
      66
          26 Yes B F
                         67
                                 No
                                            77
P
  M
                             28
                                     B F
                                                16
                                                    No
A
  F
      71
          12
              No
                   B F
                         72
                             50
                                 No
                                     В
                                        F
                                            76
                                                 9
                                                    Yes
      71
              Yes A F
                         63
                                                18
  M
          17
                             27
                                 No
                                     A F
                                            69
Α
                                                    Yes
                         62
В
  F
      66
          12
              No
                   A M
                             42
                                 No
                                     Ρ
                                       F
                                            64
                                                 1
                                                    Yes
  F
      64
          17
                   P M
                         74
                              4
                                            72
                                                25
A
              No
                                 No
                                      Α
                                       F
                                                    No
      70
                         66
P
  М
           1
              Yes
                   В
                      М
                             19
                                 No
                                      В
                                         М
                                            59
                                                29
                                                    No
A
  F
      64
          30
              No
                   A M
                         70
                             28
                                 No
                                     Α
                                        M
                                            69
                                                 1
                                                    No
  F
      78
           1
              No
                   P M
                         83
                              1
                                 Yes B
                                       F
                                            69
                                               42
                                                    No
В
                         77
      75
                                            79
В
  М
          30
              Yes P
                      M
                             29
                                 Yes P
                                         F
                                                20
                                                    Yes
  М
      70
          12
                     F
                         69
                             12
                                        F
                                            65
                                                14
Α
              No
                   Α
                                 No
                                     R
                                                    No
В
  M
      70
           1
              No
                   B M
                         67
                             23
                                 No
                                     Α
                                        М
                                            76
                                                25
                                                    Yes
P
  М
      78
          12
                   в м
                         77
                              1
                                 Yes B
                                        F
                                            69
                                                24
              Yes
                                                    No
P
  М
      66
           4
              Yes
                   Ρ
                      F
                         65
                             29
                                 No
                                      Ρ
                                         М
                                            60
                                                26
                                                    Yes
      78
                   в м
                         75
                             21
                                 Yes A F
                                            67
                                                11
  М
          15
              Yes
                                                    No
A
P
  F
      72
          27
              No
                   P F
                         70
                             13
                                 Yes A M
                                            75
                                               6
                                                    Yes
                   P F
В
  F
      65
           7
              No
                         68
                             27
                                 Yes P M
                                           68 11
                                                    Yes
      67
          17
                   В
                         70
                             22
                                            65
                                                15
P
              Yes
                      M
                                 No
                                      Α
                                         М
                                                    No
  F
                         67
P
      67
           1
              Yes A M
                             10
                                 No
                                    P
                                         F
                                            72
                                                11
                                                    Yes
      74
           1
                   В
                      М
                         80
                             21
A
                                 Yes A
                                                    No
```

The Neuralgia data set contains five variables. The Pain variable is the response. A specification of Pain=Yes indicates that the patient felt pain, and Pain=No indicates that the patient did not feel pain. The variable Treatment is a categorical variable with three levels: A and B represent the two test treatments, and P represents the placebo treatment. The gender of the patients is given by the categorical variable Sex. The variable Age is the age of the patients, in years, when treatment began. The duration of complaint, in months, before the treatment began is given by the variable Duration.

In the following statements, a complex model that includes classification and continuous covariates and an interaction term is fit to the Neuralgia data. When you try to create a default effect plot from this model, computations stop because the best type of plot cannot easily be determined.

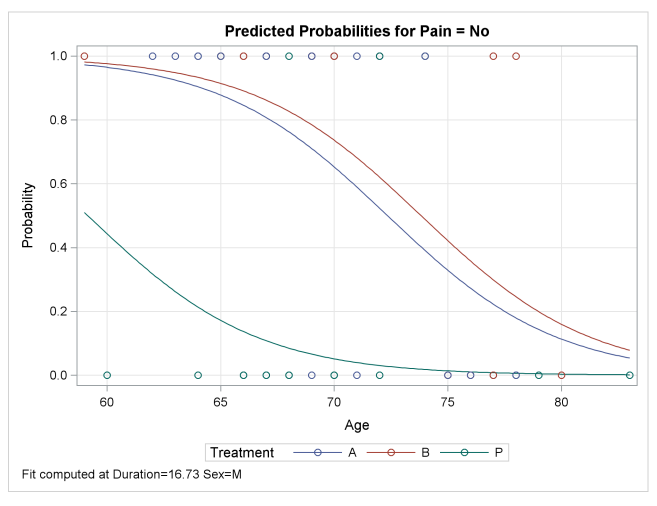
```
ods graphics on;
proc logistic data=Neuralgia;
   class Treatment Sex / param=ref;
   model Pain= Treatment|Sex Age Duration;
   effectplot;
run;
ods graphics off;
```

To produce an effect plot for this model, you need to first choose the type of plot to be created. In this case, since there are both classification and continuous covariates on the model, a SLICEFIT plot-type displays the

first continuous covariate (Age) on the X axis and displays fit curves that correspond to each level of the first classification covariate (Treatment). The following statements produce Output 19.3.1.

```
ods graphics on;
proc logistic data=Neuralgia;
   class Treatment Sex / param=ref;
   model Pain= Treatment|Sex Age Duration;
   effectplot slicefit;
run;
ods graphics off;
```

By default, effect plots from PROC LOGISTIC are displayed on the probability scale. The predicted values are computed at the mean of the Duration variable, 16.73, and at the reference level of the Sex variable, M. Observations are also displayed on the sliced-fit plot in Output 19.3.1. While the display of binary responses can give you a feel for the spread of the data, it does not enable you to evaluate the fit of the model.

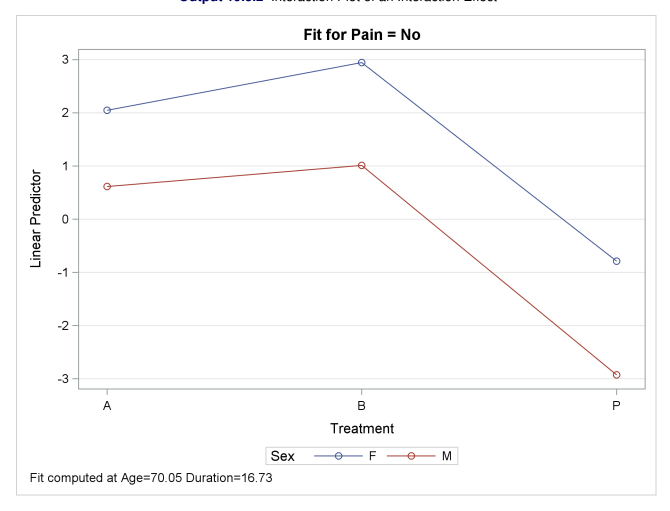


Output 19.3.1 Default Fit Plot Sliced by Treatment

In the following statements, an INTERACTION *plot-type* is specified for the Treatment variable, with the Sex effect chosen for grouping the fits. The Age and Duration variables are set to their mean values for computing the predicted values. The NOOBS option suppresses the display of the binary observations on this plot. The LINK option is specified to display the fit on the LOGIT scale; if there is no interaction between Treatment and Sex, then the resulting curves shown in Output 19.3.2 will have similar slopes across the treatments.

```
ods graphics on;
proc logistic data=Neuralgia;
   class Treatment Sex / param=ref;
   model Pain= Treatment|Sex Age Duration;
   effectplot interaction(x=Treatment sliceby=Sex) / noobs link;
run;
ods graphics off;
```

In Output 19.3.2, the slopes of the lines seem "parallel" across the treatments, corroborating the nonsignificance of the interaction terms.

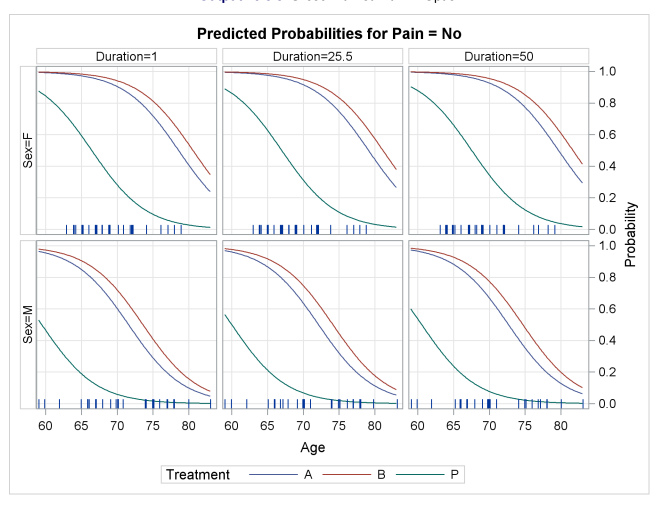


Output 19.3.2 Interaction Plot of an Interaction Effect

In the following statements, the interaction effect is removed, and the Duration variable is investigated further. The PLOTBY(ROWS)= option displays the Sex levels in the rows of a panel of plots, and the AT option computes the fits for several values of the Duration main effect in the columns of the panel. The OBS(FRINGE) option moves the observations to a fringe (rug) plot at the bottom of the plot, the observations are subsetted and displayed according to the value of the PLOTBY= variable, and the JITTER option makes overlaid fringes more visible. A STORE statement is also specified to save the model information for a later display. These statements produce Output 19.3.3.

```
ods graphics on;
proc logistic data=Neuralgia;
   class Treatment Sex / param=ref;
   model Pain= Treatment Sex Age Duration;
   effectplot slicefit(sliceby=Treatment plotby(rows)=Sex)
      / at(Duration=min midrange max) obs(fringe jitter(seed=39393));
   store logimodel;
run;
ods graphics off;
```

The predicted probability curves in Output 19.3.3 look very similar across the different values of the Duration variable, which agrees with the nonsignificance of Duration in this model. The fringe plot displays only female patients in the SEX=F row of the panel and displays only male patients in the SEX=M row, because the PLOTBY=SEX option subsets the observations.



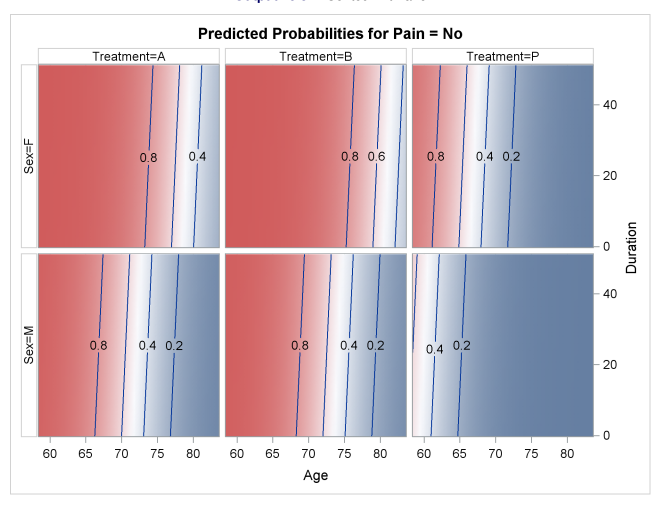
Output 19.3.3 Sliced-Fit Plot with AT Option

The following statements use the stored model and the PLM procedure to display a panel of contour plots:

```
ods graphics on;
proc plm restore=logimodel;
   effectplot contour(plotby=Treatment) / at(Sex=all);
run;
ods graphics off;
```

Output 19.3.4 again confirms that Duration is not significant.

Output 19.3.4 Contour Fit Panel



This statement documentation applies to the following SAS/STAT procedures: LIFEREG, LOGISTIC, ORTHOREG, PHREG, PLM, PROBIT, QUANTREG, SURVEYLOGISTIC, SURVEYPHREG, and SURVEYREG. It also applies to the RELIABILITY procedure in SAS/QC software. The ESTIMATE statement in the GENMOD, GLIMMIX, GLM, and MIXED procedures are documented in the respective procedure chapters.

The ESTIMATE statement provides a mechanism for obtaining custom hypothesis tests. Estimates are formed as linear estimable functions of the form  $L\beta$ . You can perform hypothesis tests for the estimable functions, construct confidence limits, and obtain specific nonlinear transformations.

# Syntax: ESTIMATE Statement

```
ESTIMATE <'label' > estimate-specification < (divisor=n) > <, <'label' > estimate-specification < (divisor=n) > > <, ... > </options > ;
```

The basic element of the ESTIMATE statement is the *estimate-specification*, which consists of model effects and their coefficients. An *estimate-specification* takes the general form

```
effect name < effect values ... >
```

The following variables can appear in the ESTIMATE statement:

is an optional label that identifies the particular row of the estimate in the output.

effect identifies an effect that appears in the MODEL statement. The keyword INTERCEPT can

be used as an effect when an intercept is fitted in the model. You do not need to include

all effects that are in the MODEL statement.

values are constants that are elements of the L matrix and are associated with the fixed and

random effects. There are two basic methods of specifying the entries of the  $\bf L$  matrix. The traditional representation—also known as the positional syntax—relies on entering coefficients in the position they assume in the  $\bf L$  matrix. For example, in the following statements the elements of  $\bf L$  that are associated with the  $\bf b$  main effect receive a 1 in the

first position and a-1 in the second position:

```
class a b;
model y = a b a*b;
estimate 'B at A2' b 1 -1 a*b 0 0 1 -1;
```

The elements that are associated with the interaction receive a 1 in the third position and a -1 in the fourth position. In order to specify coefficients correctly for the interaction term, you need to know how the levels of a and b vary in the interaction, which is governed by the order of the variables in the CLASS statement. The nonpositional syntax is designed to make it easier to enter coefficients for interactions and is necessary to enter coefficients for effects that are constructed with the EFFECT statement. In square brackets you enter the coefficient followed by the associated levels of the CLASS variables. If B has two levels and A has three levels, the previous ESTIMATE statement, by using nonpositional syntax for the interaction term, becomes the following statement:

```
estimate 'B at A2' b 1 -1 a*b [1, 2 1] [-1, 2 2];
```

The previous statement assigns value 1 to the interaction where A is at level 2 and B is at level 1, and it assigns –1 to the interaction where both classification variables are at level 2. The comma that separates the entry for the **L** matrix from the level indicators is optional. Further details about the nonpositional contrast syntax and its use with constructed effects can be found in the section "Positional and Nonpositional Syntax for Coefficients in Linear Functions" on page 455.

Based on the *estimate-specifications* in your ESTIMATE statement, the procedure constructs the matrix  $\mathbf{L}$  to test the hypothesis  $H: \mathbf{L}\boldsymbol{\beta} = \mathbf{0}$ . The procedure supports nonpositional syntax for the coefficients of model effects in the ESTIMATE statement. For details see the section "Positional and Nonpositional Syntax for Coefficients in Linear Functions" on page 455.

The procedure then produces for each row  $\mathbf{l}$  of  $\mathbf{L}$  an approximate t test of the hypothesis  $H: \mathbf{l}\boldsymbol{\beta} = 0$ . You can also obtain multiplicity-adjusted p-values and confidence limits for multirow estimates with the ADJUST= option.

Note that multirow estimates are permitted. Unlike releases prior to SAS 9.22, you do not need to specify a 'label' for every row of the estimate; the procedure constructs a default label if a label is not specified.

If the procedure finds the estimate to be nonestimable, then it displays "Non-est" for the estimate entry.

Table 19.18 summarizes important options in the ESTIMATE statement. All ESTIMATE options are subsequently discussed in alphabetical order.

 Option
 Description

 Construction and Computation of Estimable Functions

 DIVISOR=
 Specifies a list of values to divide the coefficients

 NOFILL
 Suppresses the automatic fill-in of coefficients for higher-order effects

 SINGULAR=
 Tunes the estimability checking difference

Table 19.18 ESTIMATE Statement Options

Table 19.18 continued

Option	Description
Degrees of Freedom and	p-values
ADJUST=	Determines the method for multiple comparison adjustment of estimates
ALPHA=α	Determines the confidence level $(1 - \alpha)$
LOWER	Performs one-sided, lower-tailed inference
STEPDOWN	Adjusts multiplicity-corrected <i>p</i> -values further in a step-down fashion
TESTVALUE=	Specifies values under the null hypothesis for tests
UPPER	Performs one-sided, upper-tailed inference
Statistical Output	
CL	Constructs confidence limits
CORR	Displays the correlation matrix of estimates
COV	Displays the covariance matrix of estimates
E	Prints the L matrix
JOINT	Produces a joint $F$ or chi-square test for the estimable functions
PLOTS=	Requests ODS statistical graphics if the analysis is sampling-based
SEED=	Specifies the seed for computations that depend on random numbers
Generalized Linear Mod	leling
CATEGORY=	Specifies how to construct estimable functions with multinomial data
EXP	Exponentiates and displays estimates
ILINK	Computes and displays estimates and standard errors on the inverse linked scale

You can specify the following options in the ESTIMATE statement after a slash (/).

# ADJDFE=SOURCE

#### ADJDFE=ROW

specifies how denominator degrees of freedom are determined when p-values and confidence limits are adjusted for multiple comparisons with the ADJUST= option. When you do not specify the ADJDFE= option, or when you specify ADJDFE=SOURCE, the denominator degrees of freedom for multiplicity-adjusted results are the denominator degrees of freedom for the final effect that is listed in the ESTIMATE statement from the "Type III" table.

The ADJDFE=ROW setting is useful if you want multiplicity adjustments to take into account that denominator degrees of freedom are not constant across estimates. For example, this can be the case when the denominator degrees of freedom are computed by the Satterthwaite method or according to Kenward and Roger (1997).

The ADJDFE= option has an effect only in mixed models that use these degree-of-freedom methods. It is not supported by the procedures that perform chi-square-based inference (LOGISTIC, PHREG, and SURVEYLOGISTIC).

ADJUST=BON
ADJUST=SCHEFFE
ADJUST=SIDAK
ADJUST=SIMULATE<(simoptions)>

# ADJUST=T

requests a multiple comparison adjustment for the *p*-values and confidence limits for the estimates. The adjusted quantities are produced in addition to the unadjusted quantities. Adjusted confidence limits are produced if the CL or ALPHA= option is in effect. For a description of the adjustments, see Chapter 45, "The GLM Procedure," and Chapter 67, "The MULTTEST Procedure," and the documentation for the ADJUST= option in the LSMEANS statement.

If the STEPDOWN option is in effect, the *p*-values are further adjusted in a step-down fashion.

## **ALPHA**=number

requests that a t type confidence interval be constructed with confidence level 1 - number. The value of *number* must be between 0 and 1; the default is 0.05. If the "Estimates" table shows infinite degrees of freedom, then the confidence interval is a z type interval.

# CATEGORY=category-options

specifies how to construct estimates and multiplicity corrections for models with multinomial data (ordinal or nominal). This option is also important for constructing sets of estimable functions for *F* or chi-square tests with the JOINT option.

The *category-options* are used to indicate how response variable levels are treated in constructing the estimable functions. Possible values for the *category-options* are the following:

#### **JOINT**

computes the estimable functions for every nonredundant category and treats them as a set. For example, a three-row ESTIMATE statement in a model with three response categories leads to six estimable functions.

#### **SEPARATE**

computes the estimable functions for every nonredundant category in turn. For example, a three-row ESTIMATE statement in a model with three response categories leads to two sets of three estimable functions.

# quoted-value-list

computes the estimable functions only for the specified list of values. The list must consist of formatted values of the response categories, and you must specify an *estimate-specification* for each response category in the list.

Consider the following ESTIMATE statements in the LOGISTIC procedure for an ordinal model with response categories 'vg', 'g', 'm', 'b', and 'vb'. Because there are five response categories, there are four nonredundant categories for the cumulative link model.

```
proc logistic data=icecream;
   class brand / param=glm;
   model taste(order=data) = brand / link=logit;
   freq count;
   estimate brand 1 -1,
            intercept 1 brand 0 1 / category='m','vg';
   estimate intercept 1 brand 1
                                   / category=joint
                                     adjust=simulate(seed=1);
   estimate brand 1 -1,
            brand 1 1 -2
                                   / category=separate
                                     adjust=bon;
run;
```

The first ESTIMATE statement requests a two-row estimable function. The result is produced for two of the four nonredundant response categories. The second ESTIMATE statement produces four t tests, one for each nonredundant category. The multiplicity adjustment with p-value computation by simulation treats the four estimable functions as a unit for family-wise Type I error protection. The third ESTIMATE statement computes a two-row estimable function and reports its results separately for all nonredundant categories. The Bonferroni adjustment in this statement applies to a family of two tests that correspond to the two-row estimable function. Four Bonferroni adjustments for sets of size two are performed.

The CATEGORY= option is supported only by the procedures that support generalized linear modeling (LOGISTIC and SURVEYLOGISTIC) and by PROC PLM when it is used to perform statistical analyses on item stores created by these procedures.

# **CHISQ**

requests that chi-square tests be performed in addition to F tests, when you request an F test with the JOINT option. This option has no effect in procedures that produce chi-square statistics by default.

#### CL

requests that t type confidence limits be constructed. If the procedure shows the degrees of freedom in the "Estimates" table as infinite, then the confidence limits are z intervals. The confidence level is 0.95 by default, and you can change the confidence level with the ALPHA= option. The confidence intervals are adjusted for multiplicity when you specify the ADJUST= option. However, if a step-down p-value adjustment is requested with the STEPDOWN option, only the p-values are adjusted for multiplicity.

#### **CORR**

displays the estimated correlation matrix of the linear combination of the parameter estimates.

#### COV

displays the estimated covariance matrix of the linear combination of the parameter estimates.

# **DF**=number

specifies the degrees of freedom for the t test and confidence limits. This option is not supported by the procedures that perform chi-square-based inference (LOGISTIC, PHREG, and SUVEYLOGISTIC).

#### **DIVISOR**=*value*-list

specifies a list of values by which to divide the coefficients so that fractional coefficients can be entered as integer numerators. If you do not specify *value-list*, a default value of 1.0 is assumed. Missing values in the *value-list* are converted to 1.0.

If the number of elements in *value-list* exceeds the number of rows of the estimate, the extra values are ignored. If the number of elements in *value-list* is less than the number of rows of the estimate, the last value in *value-list* is copied forward.

If you specify a row-specific divisor as part of the specification of the estimate row, this value multiplies the corresponding divisor that is implied by the *value-list*. For example, the following statement divides the coefficients in the first row by 8, and the coefficients in the third and fourth row by 3:

Coefficients in the second row are not altered.

Ε

requests that the L matrix coefficients be displayed.

#### **EXP**

requests exponentiation of the estimate. When you model data with the logit, cumulative logit, or generalized logit link functions, and the estimate represents a log odds ratio or log cumulative odds ratio, the EXP option produces an odds ratio. In proportional hazards model, this option produces estimates of hazard ratios. If you specify the CL or ALPHA= option, the (adjusted) confidence bounds are also exponentiated.

The EXP option is supported only by PROC PHREG, PROC SURVEYPHREG, the procedures that support generalized linear modeling (LOGISTIC and SURVEYLOGISTIC), and by PROC PLM when it is used to perform statistical analyses on item stores created by these procedures.

### **ILINK**

requests that the estimate and its standard error also be reported on the scale of the mean (the inverse linked scale). The computation of the inverse linked estimate depends on the estimation mode. For example, if the analysis is based on a posterior sample when a BAYES statement is present, the inversely linked estimate is the average of the inversely linked values across the sample of posterior parameter estimates. If the analysis is not based on a sample of parameter estimates, the procedure computes the value on the mean scale by applying the inverse link to the estimate. The interpretation of this quantity depends on the *effect values* specified in your ESTIMATE statement and on the link function. For example, in a model for binary data with logit link the following statements compute

$$\frac{1}{1+\exp\{-(\alpha_1-\alpha_2)\}}$$

where  $\alpha_1$  and  $\alpha_2$  are the fixed-effects solutions that are associated with the first two levels of the classification effect A:

```
class A;
model y = A / dist=binary link=logit;
estimate 'A one vs. two' A 1 -1 / ilink;
```

This quantity is not the difference of the probabilities that are associated with the two levels,

$$\pi_1 - \pi_2 = \frac{1}{1 + \exp\{-\beta_0 - \alpha_1\}} - \frac{1}{1 + \exp\{-\beta_0 - \alpha_2\}}$$

The standard error of the inversely linked estimate is based on the delta method. If you also specify the CL option, the procedure computes confidence limits for the estimate on the mean scale. In multinomial models for nominal data, the limits are obtained by the delta method. In other models they are obtained from the inverse link transformation of the confidence limits for the estimate. The ILINK option is specific to an ESTIMATE statement.

The ILINK option is supported only by the procedures that support generalized linear modeling (LOGISTIC and SURVEYLOGISTIC) and by PROC PLM when it is used to perform statistical analyses on item stores created by these procedures.

# JOINT < (joint-test-options) >

requests that a joint F or chi-square test be produced for the rows of the estimate. The JOINT option in the ESTIMATE statement essentially replaces the CONTRAST statement.

When the LOWERTAILED or the UPPERTAILED options are in effect, or if the BOUNDS option described below is in effect, the JOINT option produces the chi-bar-square statistic according to Silvapulle and Sen (2004). This statistic uses a simulation-based approach to compute p-values in situations where the alternative hypotheses of the estimable functions are not simple two-sided hypotheses. See the section "Joint Hypothesis Tests with Complex Alternatives, the Chi-Bar-Square Statistic" on page 457 for more information about this test statistic.

You can specify the following joint-test-options in parentheses:

### $ACC = \gamma$

specifies the accuracy radius for determining the necessary sample size in the simulation-based approach of Silvapulle and Sen (2004) for tests with order restrictions. The value of  $\gamma$  must be strictly between 0 and 1; the default value is 0.005.

#### EPS=€

specifies the accuracy confidence level for determining the necessary sample size in the simulationbased approach of Silvapulle and Sen (2004) for tests with order restrictions. The value of  $\epsilon$  must be strictly between 0 and 1; the default value is 0.01.

#### LABEL='label'

assigns an identifying label to the joint test. If you do not specify a label, the first non-default label for the ESTIMATE rows is used to label the joint test.

## **NOEST**

#### **ONLY**

performs only the F or chi-square test and suppresses other results from the ESTIMATE statement. This option is useful for emulating the CONTRAST statement that is available in other procedures.

#### NSAMP=n

specifies the number of samples for the simulation-based method of Silvapulle and Sen (2004). If n is not specified, it is constructed from the values of the ALPHA= $\alpha$ , the ACC= $\gamma$ , and the EPS= $\epsilon$  options. With the default values for  $\gamma$ ,  $\epsilon$ , and  $\alpha$  (0.005, 0.01, and 0.05, respectively), NSAMP=12,604 by default.

#### **CHISQ**

adds a chi-square test if the procedure produces an F test by default.

#### **BOUNDS**=value-list

specifies boundary values for the estimable linear function. The null value of the hypothesis is always zero. If you specify a positive boundary value z, the hypotheses are  $H:\theta=0$ ,  $H_a::\theta>0$  with the added constraint that  $\theta< z$ . The same is true for negative boundary values. The alternative hypothesis is then  $H_a:\theta<0$  subject to the constraint  $\theta>-|z|$ . If you specify a missing value, the hypothesis is assumed to be two-sided. The BOUNDS option enables you to specify sets of one- and two-sided joint hypotheses. If all values in *value-list* are set to missing, the procedure performs a simulation-based p-value calculation for a two-sided test.

#### **LOWER**

#### LOWERTAILED

requests that the *p*-value for the *t* test be based only on values that are less than the test statistic. A two-tailed test is the default. A lower-tailed confidence limit is also produced if you specify the CL or ALPHA= option.

Note that for ADJUST=SCHEFFE the one-sided adjusted confidence intervals and one-sided adjusted *p*-values are the same as the corresponding two-sided statistics, because this adjustment is based on only the right tail of the *F* distribution.

If you request a joint test with the JOINT option, then a one-sided left-tailed order restriction is applied to all estimable functions, and the corresponding chi-bar-square statistic of Silvapulle and Sen (2004) is computed in addition to the two-sided, standard, F or chi-square statistic. See the JOINT option for how to control the computation of the simulation-based chi-bar-square statistic.

#### **NOFILL**

suppresses the automatic fill-in of coefficients of higher-order effects.

## **PLOTS**=plot-options

produces ODS statistical graphics of the distribution of estimable functions if the procedure performs the analysis in a sampling-based mode. For example, this is the case when procedures support a BAYES statement and perform a Bayesian analysis. The estimable functions are then computed for each of the posterior parameter estimates, and the "Estimates" table reports simple descriptive statistics for the evaluated functions. The PLOTS= option enables you in this situation to visualize the distribution of the estimable function. The following *plot-options* are available:

# **ALL**

produces all possible plots with their default settings.

# BOXPLOT<(boxplot-options)>

produces box plots of the distribution of the estimable function across the posterior sample. A separate box is generated for each estimable function, and all boxes appear on a single graph by default. You can affect the appearance of the box plot graph with the following options:

# ORIENTATION=VERTICAL | HORIZONTAL

# ORIENT=VERT | HORIZ

specifies the orientation of the boxes. The default is vertical orientation of the box plots.

#### **NPANELPOS**=number

specifies how to break the series of box plots across multiple panels. If the NPANELPOS option is not specified, or if number equals zero, then all box plots are displayed in a single graph; this is the default. If a negative number is specified, then exactly up to |number| of box plots are displayed per panel. If *number* is positive, then the number of boxes per panel is balanced to achieve small variation in the number of box plots per graph.

# **DISTPLOT**<(distplot-options)>

# DIST<(distplot-options)>

generates panels of histograms with a kernel density overlaid. A separate plot in each panel contains the results for each estimable function. You can specify the following distplot-options in parentheses:

#### **BOX | NOBOX**

controls the display of a horizontal box plot of the estimable function's distribution across the posterior sample below the graph. The BOX option is enabled by default.

#### **HIST | NOHIST**

controls the display of the histogram of the estimable function's distribution across the posterior sample. The HIST option is enabled by default.

## **NORMAL | NONORMAL**

controls the display of a normal density estimate on the graph. The NONORMAL option is enabled by default.

# KERNEL | NOKERNEL

controls the display of a kernel density estimate on the graph. The KERNEL option is enabled by default.

# NROWS=number

specifies the highest number of rows in a panel. The default is 3.

#### NCOLS=number

specifies the highest number of columns in a panel. The default is 3.

#### **UNPACK**

unpacks the panel into separate graphics.

#### NONE

does not produce any plots.

# SEED=number

specifies the seed for the sampling-based components of the computations for the ESTIMATE statement (for example, chi-bar-square statistics and simulated p-values). The value of number must be an integer. The seed is used to start the pseudo-random number generator for the simulation. If you do not specify a seed, or if you specify a value less than or equal to zero, the seed is generated from reading the time of day from the computer clock. There could be multiple ESTIMATE statements with SEED= specifications and there could be other statements that can supply a random number seed. Since the procedure has only one random number stream, the initial seed is shown in the SAS log.

#### SINGULAR=number

tunes the estimability checking. If  $\mathbf{v}$  is a vector, define ABS( $\mathbf{v}$ ) to be the largest absolute value of the elements of  $\mathbf{v}$ . If ABS( $\mathbf{L} - \mathbf{L}\mathbf{T}$ ) is greater than  $c^*$ number for any row of  $\mathbf{L}$  in the contrast, then  $\mathbf{L}\boldsymbol{\beta}$  is declared nonestimable. Here,  $\mathbf{T}$  is the Hermite form matrix ( $\mathbf{X}'\mathbf{X}$ ) $^-\mathbf{X}'\mathbf{X}$ , and c is ABS( $\mathbf{L}$ ), except when it equals 0, and then c is 1. The value for *number* must be between 0 and 1; the default is 1E–4.

# STEPDOWN< (step-down-options)>

requests that multiplicity adjustments for the *p*-values of estimates be further adjusted in a step-down fashion. Step-down methods increase the power of multiple testing procedures by taking advantage of the fact that a *p*-value is never declared significant unless all smaller *p*-values are also declared significant. The STEPDOWN adjustment combined with ADJUST=BON corresponds to the methods of Holm (1979) and "Method 2" of Shaffer (1986); this is the default. Using step-down-adjusted *p*-values combined with ADJUST=SIMULATE corresponds to the method of Westfall (1997).

If the ESTIMATE statement is applied with a STEPDOWN option in a mixed model where the degrees-of-freedom method is that of Kenward and Roger (1997) or of Satterthwaite, then step-down-adjusted *p*-values are produced only if the ADJDFE=ROW option is in effect.

Also, the STEPDOWN option affects only *p*-values, not confidence limits. For ADJUST=SIMULATE, the generalized least squares hybrid approach of Westfall (1997) is used to increase Monte Carlo accuracy. You can specify the following *step-down-options* in parentheses after the STEPDOWN option:

#### MAXTIME=n

specifies the time (in seconds) to be spent computing the maximal logically consistent sequential subsets of equality hypotheses for TYPE=LOGICAL. The default is MAXTIME=60. If the MAXTIME value is exceeded, the adjusted tests are not computed. When this occurs, you can try increasing the MAXTIME value. However, note that there are common multiple comparisons problems for which this computation requires a huge amount of time—for example, all pairwise comparisons between more than 10 groups. In such cases, try to use TYPE=FREE (the default) or TYPE=LOGICAL(n) for small n.

# ORDER=PVALUE

## **ORDER=ROWS**

specifies the order in which the step-down tests to be performed. ORDER=PVALUE is the default, with estimates being declared significant only if all estimates with smaller (unadjusted) *p*-values are significant. If you specify ORDER=ROWS, then significances are evaluated in the order in which they are specified in the syntax.

# **REPORT**

specifies that a report on the step-down adjustment be displayed, including a listing of the sequential subsets (Westfall 1997) and, for ADJUST=SIMULATE, the step-down simulation results.

# TYPE=LOGICAL<(n)>

#### TYPE=FREE

specifies how step-down adjustment are made. If you specify TYPE=LOGICAL, the step-down adjustments are computed by using maximal logically consistent sequential subsets of equality hypotheses (Shaffer 1986; Westfall 1997). Alternatively, for TYPE=FREE, sequential subsets are computed ignoring logical constraints. The TYPE=FREE results are more conservative than those for TYPE=LOGICAL, but they can be much more efficient to produce for many estimates. For example, it is not feasible to take logical constraints between all pairwise comparisons of more than about 10 groups. For this reason, TYPE=FREE is the default.

However, you can reduce the computational complexity of taking logical constraints into account by limiting the depth of the search tree used to compute them, specifying the optional depth parameter as a number n in parentheses after TYPE=LOGICAL. As with TYPE=FREE, results for TYPE=LOGICAL(n) are conservative relative to the true TYPE=LOGICAL results. But even for TYPE=LOGICAL(0) they can be appreciably less conservative than TYPE=FREE, and they are computationally feasible for much larger numbers of estimates. If you do not specify n or if n = -1, the full search tree is used.

#### **TESTVALUE**=*value*-list

#### **TESTMEAN=**value-list

specifies the value under the null hypothesis for testing the estimable functions in the ESTIMATE statement. The rules for specifying the *value-list* are very similar to those for specifying the divisor list in the DIVISOR= option. If no TESTVALUE= is specified, all tests are performed as  $H: \mathbf{L}\boldsymbol{\beta} = 0$ . Missing values in the *value-list* also are translated to zeros. If you specify fewer values than rows in the ESTIMATE statement, the last value in *value-list* is carried forward.

The TESTVALUE= option affects only *p*-values from individual, joint, and multiplicity-adjusted tests. It does not affect confidence intervals.

The TESTVALUE option is not available for the multinomial distribution, and the values are ignored when you perform a sampling-based (Bayesian) analysis.

#### **UPPER**

# **UPPERTAILED**

requests that the *p*-value for the *t* test be based only on values that are greater than the test statistic. A two-tailed test is the default. An upper-tailed confidence limit is also produced if you specify the CL or ALPHA= option.

Note that for ADJUST=SCHEFFE the one-sided adjusted confidence intervals and one-sided adjusted *p*-values are the same as the corresponding two-sided statistics, because this adjustment is based on only the right tail of the *F* distribution.

If you request a joint test with the JOINT option, then a one-sided right-tailed order restriction is applied to all estimable functions, and the corresponding chi-bar-square statistic of Silvapulle and Sen (2004) is computed in addition to the two-sided, standard, *F* or chi-square statistic. See the JOINT option for how to control the computation of the simulation-based chi-bar-square statistic.

# Positional and Nonpositional Syntax for Coefficients in Linear Functions

When you define custom linear hypotheses with the ESTIMATE statement, the procedure sets up an **L** vector or matrix that conforms to the model effect solutions. (Note that the following remarks also apply to the LSMESTIMATE statement, where you specify coefficients of the matrix **K** which is then converted into a coefficient matrix that conforms to the model effects solutions.)

There are two methods for specifying the entries in a coefficient matrix (hereafter simply referred to as the L matrix); they are called the positional and nonpositional methods. In the positional form, which is the traditional method, you provide a list of values that occupy the elements of the L matrix that is associated with the effect in question in the order in which the values are listed. For traditional model effects that consist of continuous and classification variables, the positional syntax is simpler in some cases (main effects) and more cumbersome in others (interactions). When you work with effects that are constructed through the EFFECT statement, the nonpositional syntax is essential.

For example, consider the following two-way model with interactions where factors A and B have three and two levels, respectively:

```
proc logistic;
   class a b;
   model y = a b a*b;
run;
```

To test the difference of the B levels at the second level of A with an ESTIMATE statement (a slice), you need to assign coefficients 1 and -1 to the levels of B and to the levels of the interaction where A is at the second level. Two examples of equivalent ESTIMATE statements that use positional and nonpositional syntax are as follows:

```
estimate 'B at A2' b 1 -1 a*b 0 0 1 -1 ; estimate 'B at A2' b 1 -1 a*b [1 \ 2 \ 1] [-1 \ 2 \ 2];
```

Because A precedes B in the CLASS statement, the levels of the interaction are formed as  $\alpha_1\beta_1,\alpha_1\beta_2,\alpha_2\beta_1,\alpha_2\beta_2,\cdots$ . If B precedes A in the CLASS statement, you need to modify the coefficients accordingly:

```
proc logistic;
  class b a;
  model y = a b a*b;
  estimate 'B at A2' b 1 -1 a*b 0 1 0 0 -1 ;
  estimate 'B at A2' b 1 -1 a*b [1 1 2] [-1 2 2];
  estimate 'B at A2' b 1 -1 a*b [1, 1 2] [-1, 2 2];
run:
```

You can optionally separate the **L** value entry from the level indicators with a comma, as in the last ESTIMATE statement.

The general syntax for defining coefficients with the nonpositional syntax is as follows:

```
effect-name [multiplier <, > level-values] ... < [multiplier <, > level-values] >
```

The first entry in square brackets is the multiplier that is applied to the elements of **L** for the effect after the *level-values* have been resolved and any necessary action that forms **L** has been taken.

The *level-values* are organized in a specific form:

- The number of entries should equal the number of terms that are needed to construct the effect. For effects that do not contain any constructed effects, this number is simply the number of terms in the name of the effect.
- Values of continuous variables that are needed for the construction of the L matrix precede the level indicators of CLASS variables.
- If the effect involves constructed effects, then you need to provide as many continuous and classification variables as are needed for the effect formation. For example, if a collection effect is defined as

```
class c;
effect v = collection(x1 x2 c);
```

then a proper nonpositional syntax would be

```
v [0.5, 0.2 0.3 3]
```

• If an effect contains both regular terms (old-style effects) and constructed effects, then the order of the coefficients is as follows: continuous values for old-style effects, class levels for classification variables in old-style effects, continuous values for constructed effects, and finally class levels that are needed for constructed effects. Assume that C has four levels so that effect v contributes six elements to the L matrix. When the procedure resolves this syntax, the values 0.2 and 0.3 are assigned to the positions for x1 and x2 and a 1 is associated with the third level of C. The resulting vector is then multiplied by 0.5 to produce

```
[0.1 \quad 0.15 \quad 0 \quad 0 \quad 0.5 \quad 0]
```

Note that you enter the **levels** of the classification variables in the square brackets, not their formatted values. The ordering of the levels of classification variables can be gleaned from the "Class Level Information" table.

To specify values for continuous variables, simply give their value as one of the terms in the effect. The nonpositional syntax in the following ESTIMATE statement is read as "1 times the value 0.4 in the column that is associated with level 2 of A"

```
proc phreg;
  class a / param=glm;
  model y = a a*x / s;
  lsmeans a / e at x=0.4;
  estimate 'A2 at x=0.4' intercept 1 a 0 1 a*x [1,0.4 2] / e;
run;
```

Because the value before the comma serves as a multiplier, the same estimable function could also be constructed with the following statements:

```
estimate 'A2 at x=0.4' intercept 1 a 0 1 a*x [ 4, 0.1 2]; estimate 'A2 at x=0.4' intercept 1 a 0 1 a*x [ 2, 0.2 2]; estimate 'A2 at x=0.4' intercept 1 a 0 1 a*x [-1, -0.4 2];
```

Note that continuous variables that are needed to construct an effect are always listed before any CLASS variables.

When you work with constructed effects, the nonpositional syntax works in the same way. For example, the following model contains a classification effect and a B-spline. The first two ESTIMATE statements produce predicted values for level 1 of C when the continuous variable x takes on the values 20 and 10, respectively.

In this example, the ORTHOREG procedure computes the spline coefficients for the first ESTIMATE statement based on x = 20, and similarly in the second statement for x = 10. The third ESTIMATE statement computes the difference of the predicted values. Because the spline effect does not interact with the classification variable, this difference does not depend on the level of C. If such an interaction is present, you can estimate the difference in predicted values for a given level of C by using the nonpositional syntax. Because the effect C\*spl contains both old-style terms (C) and a constructed effect, you specify the values for the old-style terms before assigning values to constructed effects.

```
proc orthoreg;
  class c;
  effect spl = spline(x / knotmethod=equal(5));
  model y = spl*c;
  estimate 'C2 = 1, x=20' intercept 1 c*spl [1,1 20];
  estimate 'C2 = 2, x=20' intercept 1 c*spl [1,2 20];
  estimate 'C diff at x=20' c*spl [1,1 20] [-1,2 20];
run;
```

It is recommended that you add the E option to the ESTIMATE or LSMESTIMATE statement to verify that the L matrix is formed according to your expectations.

In any row of an ESTIMATE statement you can choose positional and nonpositional syntax separately for each effect. However, you cannot mix the two forms of syntax for coefficients of a single effect. For example, the following statement is not proper because both forms of syntax are used for the interaction effect:

```
estimate 'A1B1 - A1B2' b 1 -1 a*b 0 1 [-1, 1 2];
```

# Joint Hypothesis Tests with Complex Alternatives, the Chi-Bar-Square Statistic

Silvapulle and Sen (2004) propose a test statistic for testing hypotheses where the null or the alternative hypothesis or both involve inequalities. You can test special cases of these hypotheses with the JOINT option in the ESTIMATE and the LSMESTIMATE statement. Consider the k estimable functions  $\mathbf{L}\boldsymbol{\beta}$  and the hypotheses  $H_0$ :  $\mathbf{L}\boldsymbol{\beta} = \mathbf{0}$  and  $H_a$ :  $\mathbf{L}\boldsymbol{\beta} \geq \mathbf{0}$ . The alternative hypothesis defines a convex cone  $\mathcal C$  at the origin. Suppose that under the null hypothesis  $\mathbf{L}\widehat{\boldsymbol{\beta}}$  follows a multivariate normal distribution with mean  $\mathbf{0}$  and variance  $\mathbf{V}$ . The restricted alternative prevents you from using the usual F or chi-square test machinery,

p-values are obtained by simulation; see, in particular, Chapter 3.4 in Silvapulle and Sen (2004).

Briefly, let U be a multivariate normal random variable with mean 0 and variance matrix V. The chi-bar-square statistic is the random variable

$$\overline{\chi}^{2} = \mathbf{U}'\mathbf{V}^{-1}\mathbf{U} - Q$$

$$Q = \min_{\boldsymbol{\theta} \in C} (\mathbf{U} - \boldsymbol{\theta})'\mathbf{V}^{-1}(\mathbf{U} - \boldsymbol{\theta})$$

and it can be motivated by a geometric argument. The quadratic form in Q is the V-projection of U onto the cone C. Suppose that this projected point is  $\tilde{U}$ . If  $U \in C$ , then Q = 0 and  $\tilde{U} = U$ . If U is completely outside of the cone C, then  $\tilde{U}$  is a point on the surface of the cone. Similarly,  $U'V^{-1}U$  is the length of the segment from the origin to U in the V-space with norm  $||x|| = (x'V^{-1}x)^{1/2}$ . If you apply the Pythagorean theorem, you can see that the chi-bar-square statistic measures the length of the segment from the origin to the projected point  $\tilde{U}$  in C.

To calculate p-values for chi-bar-square statistics, a simulation-based approach is taken. Consider again the set of k estimable functions  $\mathbf{L}\boldsymbol{\beta}$  with estimate  $\mathbf{L}\widehat{\boldsymbol{\beta}} = \mathbf{U}$  and variance  $\mathbf{L}\mathrm{Var}[\hat{\boldsymbol{\beta}}]\mathbf{L}' = \mathbf{V}$ .

First, the observed value of the statistic is computed as

$$\overline{\chi}_{obs}^2 = \mathbf{U}'\mathbf{V}^{-1}\mathbf{U} - Q$$

Then, n independent random samples  $\mathbb{Z}_1, \dots, \mathbb{Z}_n$  are drawn from an  $N(0, \mathbb{V})$  distribution and the following chi-bar-statistics are computed for the sample:

$$\overline{\chi}_{1}^{2} = \mathbf{Z}_{1}'\mathbf{V}^{-1}\mathbf{Z}_{1} - \min_{\boldsymbol{\theta} \in C}(\mathbf{Z}_{1} - \boldsymbol{\theta})'\mathbf{V}^{-1}(\mathbf{Z}_{1} - \boldsymbol{\theta})$$

$$\vdots$$

$$\overline{\chi}_{n}^{2} = \mathbf{Z}_{n}'\mathbf{V}^{-1}\mathbf{Z}_{n} - \min_{\boldsymbol{\theta} \in C}(\mathbf{Z}_{n} - \boldsymbol{\theta})'\mathbf{V}^{-1}(\mathbf{Z}_{n} - \boldsymbol{\theta})$$

The *p*-value is estimated by the fraction of simulated statistics that are greater than or equal to the observed value  $\overline{\chi}_{obs}^2$ .

Notice that unless U is interior to the cone C, finding the value of Q requires the solution to a quadratic optimization problem. When k is large, or when many simulations are requested, the computation of p-values for chi-bar-square statistics might require considerable computing time.

# **ODS Table Names: ESTIMATE Statement**

Each table created by the ESTIMATE statement has a name associated with it, and you can use this name to refer to the table when you use the Output Delivery System (ODS) to select tables and create output data sets. These names are listed in Table 19.19. For more information about ODS, see Chapter 20, "Using the Output Delivery System."

Table 19.19 ODS Tables Produced by the ESTIMATE statement

Table Name	Description	Required Option
Coef	L matrix coefficients	E
Estimates	ESTIMATE statement results	Default
Contrasts	Joint test results	JOINT

# **ODS Graphics: ESTIMATE Statement**

This section describes the use of ODS Graphics for creating statistical graphs of the distribution of estimable functions with the ESTIMATE statement. The plots can be produced only in association with the LIFEREG and PHREG procedures, which can perform Bayesian analysis. The plots are available via these procedures directly, and also via PROC PLM when it is run using an item store that was created by these procedures.

To request these graphs you must do the following:

- ensure that ODS Graphics is enabled
- use a BAYES statement with PROC LIFEREG or PROC PHREG, or use PROC PLM to perform statistical analysis on an item store that was saved from a Bayesian analysis
- request plots with the PLOTS= option in the ESTIMATE statement

For more information about ODS Graphics, see Chapter 21, "Statistical Graphics Using ODS." The available graphs are summarized in Table 19.20.

**Table 19.20** Graphs Produced by the ESTIMATE statement

ODS Graph Name	Plot Description	Required Option
BoxPlot	Displays box plots of estimable functions across a posterior sample.	PLOTS=BOXPLOT
DistPanel	Displays panels of histograms with kernel density curves overlaid. Each plot contains the results for the posterior sample of each estimable function.	PLOTS=DISTPLOT
DistPlot	Displays a histogram with a kernel density curve overlaid. The plot contains the results for the posterior sample of the estimable function.	PLOTS=DISTPLOT(UNPACK)

For details about the *plot-options* of the ESTIMATE statement, see the PLOTS= option in the section "ESTIMATE Statement" on page 444.

# **LSMEANS Statement**

This statement documentation applies to the following procedures:

GENMOD, LIFEREG, LOGISTIC, ORTHOREG, PHREG, PLM, PROBIT, SURVEYLOGISTIC, SUR-VEYPHREG, and SURVEYREG. It also applies to the RELIABILITY procedure in SAS/QC software. The GLIMMIX, GLM, and MIXED procedures also support LSMEANS statements. The relevant statement documentation for these procedures can be found in the specific procedure chapter.

The LSMEANS statement computes least squares means (LS-means) of fixed effects. In the GLM, MIXED, and GLIMMIX procedures, LS-means are predicted population margins—that is, they estimate the marginal means over a balanced population. In a sense, LS-means are to unbalanced designs as class and subclass arithmetic means are to balanced designs.

Thus it is important not to interpret the name with a strict association with least squares estimation. Least squares is the predominant estimation technique for the type of models in which LS-means were first applied. Their interpretation and importance reaches beyond the least squares principle, however. A more appropriate approach to LS-means views them as linear combinations of the parameter estimates that are constructed in such a way that they correspond to average predicted values in a population where the levels of classification variables are balanced.

This contemporary—and historically correct—interpretation of the concept of least squares means underlines their importance in all classes of models where predicted values are reasonably formed as linear combinations of the parameter estimates. LS-means distinguish themselves from general estimable functions in that they take the structure for the model and data into account through the structure of the X and X'X matrix in your model. For example, in a generalized linear model the structure of the X matrix informs the analysis about the possible levels of classification variables and predictions on the linear (the linked) scale are computed as  $x'\beta$ . LS-means are thus meaningful quantities in such models when the linear estimable function that corresponds to an averaged prediction is constructed on the linked scale. For example, in a binomial model with logit link, the least squares means are predicted population margins of the logits. You can then transform the least squares means to the data scale with the ILINK option, and you can display differences of least squares means in terms of odds ratios with the ODDSRATIO option. The underlying principle—unless you perform a Bayesian analysis—is to construct the estimates or their differences on the linked scale and to apply appropriate transformations in a second step.

Least squares means computations are also supported for multinomial models.

LS-means are computed as  $L\beta$  where the L matrix that is constructed to compute the predicted values is the same as the L matrix that is formed in PROC GLM.

Each LS-mean is computed as  $L\hat{\beta}$ , where L is the coefficient matrix that is associated with the least squares mean and  $\hat{\beta}$  is the estimate of the fixed-effects parameter vector. The approximate standard error for the LS-mean is computed as the square root of  $\widehat{\mathbf{LVar}}[\widehat{\boldsymbol{\beta}}]\mathbf{L}'$ . The approximate variance matrix of the fixed-effects estimates depends on the estimation method.

# **Syntax: LSMEANS Statement**

LSMEANS < model-effects > < / options > ;

LS-means can be computed for any effect in the statistical model that involves only CLASS variables. You can specify multiple effects in one LSMEANS statement or in multiple LSMEANS statements, and all LSMEANS statements must appear after the MODEL statement. If you do not specify *model-effects*, the options in the LSMEANS statement are applied to all suitable model effects.

As in the ESTIMATE statement, the L matrix is tested for estimability; if this test fails, the procedure displays "Non-est" for the LS-means entries. Note that linear functions of LS-means, such as differences, can be estimable, even if the means themselves are not estimable. Estimability checks for differences are thus applied separately from checks for the means.

Assuming the LS-mean is estimable, the procedure constructs an approximate *t* test to test the null hypothesis that the associated population quantity equals zero.

Table 19.21 summarizes important options in the LSMEANS statement. All LSMEANS options are subsequently discussed in alphabetical order.

Table 19.21 LSMEANS Statement Options

Option	Description
	mputation of LS-Means
AT	Modifies the covariate value in computing LS-means
BYLEVEL	Computes separate margins
DIFF	Requests differences of LS-means
OM=	Specifies the weighting scheme for LS-means computation as de-
	termined by the input data set
SINGULAR=	Tunes estimability checking
Degrees of Freedom a	and p-values
ADJUST=	Determines the method for multiple-comparison adjustment of LS-
	means differences
$ALPHA=\alpha$	Determines the confidence level $(1 - \alpha)$
STEPDOWN	Adjusts multiple-comparison p-values further in a step-down
	fashion
Statistical Output	
CL	Constructs confidence limits for means and mean differences
CORR	Displays the correlation matrix of LS-means
COV	Displays the covariance matrix of LS-means
E	Prints the L matrix
LINES	Produces a "Lines" display for pairwise LS-means differences
MEANS	Prints the LS-means
PLOTS=	Requests graphs of means and mean comparisons
SEED=	Specifies the seed for computations that depend on random numbers

**Table 19.21** continued **Option Description Generalized Linear Modeling** Exponentiates and displays estimates of LS-means or LS-means differences **ILINK** Computes and displays estimates and standard errors of LS-means (but not differences) on the inverse linked scale **ODDSRATIO** Reports (simple) differences of least squares means in terms of odds ratios if permitted by the link function

You can specify the following options in the LSMEANS statement after a slash (/):

# ADJDFE=ROW ADJDFE=SOURCE

specifies how denominator degrees of freedom are determined when p-values and confidence limits are adjusted for multiple comparisons with the ADJUST= option. When you do not specify the ADJDFE= option or when you specify ADJDFE=SOURCE, the denominator degrees of freedom for multiplicity-adjusted results are the denominator degrees of freedom for the LS-mean effect in the "Type III Tests of Fixed Effects" table. When you specify ADJDFE=ROW, the denominator degrees of freedom for multiplicity-adjusted results correspond to the degrees of freedom that are displayed in the DF column of the "Differences of Least Squares Means" table.

The ADJDFE=ROW setting is particularly useful if you want multiplicity adjustments to take into account that denominator degrees of freedom are not constant across LS-mean differences.

In one-way models with heterogeneous variance, combining certain ADJUST= options with the ADJDFE=ROW option corresponds to particular methods of performing multiplicity adjustments in the presence of heteroscedasticity. For example, the following statements fit a heteroscedastic one-way model and perform Dunnett's T3 method (Dunnett 1980), which is based on the studentized maximum modulus (ADJUST=SMM):

```
proc glimmix;
   class A;
   model y = A / ddfm=satterth;
   random _residual_ / group=A;
   lsmeans A / adjust=smm adjdfe=row;
run:
```

If you combine the ADJDFE=ROW option with ADJUST=SIDAK, the multiplicity adjustment corresponds to the T2 method of Tamhane (1979), and ADJUST=TUKEY corresponds to the method of Games-Howell (Games and Howell 1976). Note that ADJUST=TUKEY gives the exact results for the case of fractional degrees of freedom in the one-way model, but it does not take into account that the degrees of freedom are subject to variability. A more conservative method, such as ADJUST=SMM, might protect the overall error rate better.

Unless the ADJUST= option is specified in the LSMEANS statement, the ADJDFE= option has no effect. The option is not supported by the procedures that perform chi-square-based inference (GENMOD, LOGISTIC, PHREG, and SURVEYLOGISTIC).

ADJUST=BON

ADJUST=DUNNETT

ADJUST=NELSON

ADJUST=SCHEFFE

ADJUST=SIDAK

ADJUST=SIMULATE<(simoptions)>

ADJUST=SMM | GT2

ADJUST=TUKEY

requests a multiple comparison adjustment for the *p*-values and confidence limits for the differences of LS-means. The adjusted quantities are produced in addition to the unadjusted quantities. By default, the procedure performs all pairwise differences. If you specify ADJUST=DUNNETT, the procedure analyzes all differences with a control level. If you specify ADJUST=NELSON, ANOM differences are taken. The ADJUST= option implies the DIFF option.

The BON (Bonferroni) and SIDAK adjustments involve correction factors described in Chapter 45, "The GLM Procedure," and Chapter 67, "The MULTTEST Procedure"; also see Westfall and Young (1993) and Westfall et al. (1999). When you specify ADJUST=TUKEY and your data are unbalanced, the procedure uses the approximation described in Kramer (1956) and identifies the adjustment as "Tukey-Kramer" in the results. Similarly, when you specify ADJUST=DUNNETT or ADJUST=NELSON and the LS-means are correlated, the procedure uses the factor-analytic covariance approximation described in Hsu (1992) and identifies the adjustment in the results as "Dunnett-Hsu" or "Nelson-Hsu," respectively. The approximation derives an approximate "effective sample sizes" for which exact critical values are computed. Computing the exact adjusted *p*-values and critical values for unbalanced designs can be computationally intensive, in particular for ADJUST=NELSON. A simulation-based approach, as specified by the ADJUST=SIM option, while nondeterministic, can provide inferences that are sufficiently accurate in much less time. The preceding references also describe the SCHEFFE and SMM adjustments.

Nelson's adjustment applies only to the analysis of means (Ott 1967; Nelson 1982, 1991, 1993), where LS-means are compared against an average LS-mean. It does not apply to all pairwise differences of least squares means. See the DIFF=ANOM option for more details regarding the analysis of means with the procedure.

The SIMULATE adjustment computes adjusted p-values and confidence limits from the simulated distribution of the maximum or maximum absolute value of a multivariate t random vector. All covariance parameters, except the residual scale parameter, are fixed at their estimated values throughout the simulation, potentially resulting in some underdispersion. The simulation estimates q, the true  $(1-\alpha)$  quantile, where  $1-\alpha$  is the confidence coefficient. The default  $\alpha$  is 0.05, and you can change this value with the ALPHA= option in the LSMEANS statement.

The number of samples is set so that the tail area for the simulated q is within  $\gamma$  of  $1 - \alpha$  with  $100(1 - \epsilon)\%$  confidence. In equation form,

$$\Pr(|F(\widehat{q}) - (1 - \alpha)| \le \gamma) = 1 - \epsilon$$

where  $\hat{q}$  is the simulated q and F is the true distribution function of the maximum; see Edwards and Berry (1987) for details. By default,  $\gamma = 0.005$  and  $\epsilon = 0.01$ , placing the tail area of  $\hat{q}$  within 0.005 of 0.95 with 99% confidence. You can specify the following *simoptions* in parentheses after the ADJUST=SIMULATE option:

#### ACC=value

specifies the target accuracy radius  $\gamma$  of a  $100(1-\epsilon)\%$  confidence interval for the true probability content of the estimated  $(1-\alpha)$  quantile. The default value is ACC=0.005.

#### EPS=value

specifies the value  $\epsilon$  for a  $100 \times (1 - \epsilon)\%$  confidence interval for the true probability content of the estimated  $(1 - \alpha)$  quantile. The default value for the accuracy confidence is 99%, which corresponds to EPS=0.01.

#### NSAMP=n

specifies the sample size for the simulation. By default, n is set based on the values of the target accuracy radius  $\gamma$  and accuracy confidence  $100 \times (1 - \epsilon)\%$  for an interval for the true probability content of the estimated  $(1 - \alpha)$  quantile. With the default values for  $\gamma$ ,  $\epsilon$ , and  $\alpha$  (0.005, 0.01, and 0.05, respectively), NSAMP=12,604 by default.

# SEED=number

specifies an integer that is used to start the pseudo-random number generator for the simulation. If you do not specify a seed, or specify a value less than or equal to zero, the seed is by default generated from reading the time of day from the computer's clock.

#### **THREADS**

specifies that the computational work for the simulation be divided into parallel threads, where the number of threads is the value of the SAS system option CPUCOUNT=. For large simulations (as specified directly using the NSAMP= *simoption* or indirectly using the ACC= or EPS= *simoptions*), parallel processing can markedly speed up the computation of adjusted *p*-values and confidence intervals. However, because the parallel processing has different pseudo-random number streams, the precise results are different from the default ones, which are computed in sequence rather than in parallel. This option overrides the SAS system option THREADS | NOTHREADS.

## **NOTHREADS**

specifies that the computational work for the simulation be performed in sequence rather than in parallel. NOTHREADS is the default. This option overrides the SAS system option THREADS | NOTHREADS.

If the STEPDOWN option is in effect, the *p*-values are further adjusted in a step-down fashion. For certain options and data, this adjustment is exact under an iid  $N(0, \sigma^2)$  model for the dependent variable, in particular for the following:

- for ADJUST=DUNNETT when the means are uncorrelated
- for ADJUST=TUKEY with STEPDOWN(TYPE=LOGICAL) when the means are balanced and uncorrelated.

The first case is a consequence of the nature of the successive step-down hypotheses for comparisons with a control; the second uses an extension of the maximum studentized range distribution appropriate for partition hypotheses (Royen 1989). Finally, for STEPDOWN(TYPE=FREE), ADJUST=TUKEY employs the Royen (1989) extension in such a way that the resulting *p*-values are conservative.

#### **ALPHA**=number

requests that a t type confidence interval be constructed for each of the LS-means with confidence level 1 - number. The value of number must be between 0 and 1; the default is 0.05.

AT variable=value

AT (variable-list)=(value-list)

#### AT MEANS

modifies the values of the covariates that are used in computing LS-means. By default, all covariate effects are set equal to their mean values for computation of standard LS-means. The AT option enables you to assign arbitrary values to the covariates. Additional columns in the output table indicate the values of the covariates.

If there is an effect that contains two or more covariates, the AT option sets the effect equal to the product of the individual means rather than the mean of the product (as with standard LS-means calculations). The AT MEANS option sets covariates equal to their mean values (as with standard LS-means) and incorporates this adjustment to crossproducts of covariates.

As an example, consider the following statements:

```
class A;
model Y = A x1 x2 x1*x2;
lsmeans A;
lsmeans A / at means;
lsmeans A / at x1=1.2;
lsmeans A / at (x1 x2) = (1.2 0.3);
```

For the first two LSMEANS statements, the LS-means coefficient for x1 is  $\overline{x}_1$  (the mean of x1) and for x2 is  $\overline{x}_2$  (the mean of x2). However, for the first LSMEANS statement, the coefficient for x1\*x2 is  $\overline{x}_1x_2$ , but for the second LSMEANS statement, the coefficient is  $\overline{x}_1 \times \overline{x}_2$ . The third LSMEANS statement sets the coefficient for x1 equal to 1.2 and leaves it at  $\overline{x}_2$  for x2, and the final LSMEANS statement sets these values to 1.2 and 0.3, respectively.

Even if you specify a WEIGHT variable, the unweighted covariate means are used for the covariate coefficients if there is no AT specification. If you specify the AT option, WEIGHT or FREQ variables are taken into account as follows. The weighted covariate means are then used for the covariate coefficients for which no explicit AT values are given, or if you specify AT MEANS. Observations that do not contribute to the analysis because of a missing dependent variable are included in computing the covariate means. Use the E option in conjunction with the AT option to check that the modified LS-means coefficients are the ones you want.

The AT option is disabled if you specify the BYLEVEL option.

## **BYLEVEL**

requests that separate margins be computed for each level of the LSMEANS effect.

The standard LS-means have equal coefficients across classification effects. The BYLEVEL option changes these coefficients to be proportional to the observed margins. This adjustment is reasonable when you want your inferences to apply to a population that is not necessarily balanced but has the margins observed in the input data set. In this case, the resulting LS-means are actually equal to raw means for fixed-effects models and certain balanced random-effects models, but their estimated

standard errors account for the covariance structure that you have specified. If a WEIGHT statement is specified, the procedure uses weighted margins to construct the LS-means coefficients.

If the AT option is specified, the BYLEVEL option disables it.

#### CL

requests that t type confidence limits be constructed for each of the LS-means. The confidence level is 0.95 by default; this can be changed with the ALPHA= option. If you specify an ADJUST= option, then the confidence limits are adjusted for multiplicity. But if you also specify STEPDOWN, then only p-values are step-down adjusted, not the confidence limits.

#### **CORR**

displays the estimated correlation matrix of the least squares means as part of the "Least Squares Means" table.

#### COV

displays the estimated covariance matrix of the least squares means as part of the "Least Squares Means" table.

#### **DF**=number

specifies the degrees of freedom for the t test and confidence limits. The default is the denominator degrees of freedom taken from the "Type III Tests" table that corresponds to the LS-means effect. The option is not supported by the procedures that perform chi-square-based inference (GENMOD, LOGISTIC, PHREG and SURVEYLOGISTIC).

# DIFF<=difftype>

# PDIFF<=difftype>

requests that differences of the LS-means be displayed. You can use one of the following optional difftype values to specify which differences to produce:

# **ALL**

requests all pairwise differences; this is the default.

## **ANOM**

requests differences between each LS-mean and the average LS-mean, as in the analysis of means (Ott 1967). The average is computed as a weighted mean of the LS-means, the weights being inversely proportional to the diagonal entries of the  $L(X'X)^{-}L'$  matrix. If LS-means are nonestimable, this design-based weighted mean is replaced with an equally weighted mean. Note that the ANOM procedure in SAS/QC software implements both tables and graphics for the analysis of means with a variety of response types. For one-way designs and normal data with identity link, the DIFF=ANOM computations are equivalent to the results of PROC ANOM. If the LS-means being compared are uncorrelated, exact adjusted p-values and critical values for confidence limits can be computed in the analysis of means; see Nelson (1982, 1991, 1993) and Guirguis and Tobias (2004) in addition to the documentation for the ADJUST=NELSON option.

# **CONTROL**

requests differences with a control, which, by default, is the first valid level of each of the specified LSMEANS effects. For example, suppose the effects A and B are classification variables, both of them have two levels 1 and 2, and the A=1, B=1 cell is missing. Unless the procedure supports a MISSING option in the CLASS statement and the option is in effect, the following LSMEANS statement uses the level (1,2) of A\*B as the control:

## lsmeans A\*B / diff=control;

Nevertheless, you can still specify a valid level as the control—for example, (2,1) of A\*B. To specify which levels of the effects are the controls, list the quoted formatted values in parentheses after the CONTROL keyword. For example, if the effects A, B, and C are classification variables, each having two levels, 1 and 2, the following LSMEANS statement specifies the (1,2) level of A\*B and the (2,1) level of B\*C as controls:

```
lsmeans A*B B*C / diff=control('1' '2' '2' '1');
```

For multiple effects, the results depend upon the order of the list, and so you should check the output to make sure that the controls are correct.

Two-tailed tests and confidence limits are associated with the CONTROL difftype. For one-tailed results, use either the CONTROLL or CONTROLU difftype.

## **CONTROLL**

tests whether the noncontrol levels are significantly smaller than the control; the upper confidence limits for the control minus the noncontrol levels are considered to be infinity and are displayed as missing.

## **CONTROLU**

tests whether the noncontrol levels are significantly larger than the control; the upper confidence limits for the noncontrol levels minus the control are considered to be infinity and are displayed as missing.

If you want to perform multiple comparison adjustments on the differences of LS-means, you must specify the ADJUST= option.

The differences of the LS-means are displayed in a table titled "Differences of Least Squares Means."

Ε

requests that the L matrix coefficients for the LSMEANS effects be displayed.

## **EXP**

requests exponentiation of the LS-means or LS-mean differences. When you model data with the logit, cumulative logit, or generalized logit link functions, and the estimate represents a log odds ratio or log cumulative odds ratio, the EXP option produces an odds ratio. In proportional hazards model, the exponentiation of the LS-mean differences produces estimates of hazard ratios. If you specify the CL or ALPHA= option, the (adjusted) confidence bounds are also exponentiated.

The EXP option is supported only by PROC PHREG, PROC SURVEYPHREG, the procedures that support generalized linear modeling (GENMOD, LOGISTIC, and SURVEYLOGISTIC), and PROC PLM when it is used to perform statistical analyses on item stores that are created by these procedures.

# **ILINK**

requests that estimates and their standard errors in the "Least Squares Means" table also be reported on the scale of the mean (the inverse linked scale). This enables you to obtain estimates of predicted probabilities and their standard errors in logistic models, for example. The option is specific to an LSMEANS statement. If you also specify the CL option, the procedure computes confidence intervals

for the predicted means by applying the inverse link transform to the confidence limits on the linked (linear) scale. Standard errors on the inverse linked scale are computed by the delta method.

The ILINK option is supported only by the procedures that support generalized linear modeling (GENMOD, LOGISTIC and SURVEYLOGISTIC) and by PROC PLM when it is used to perform statistical analyses on item stores that are created by these procedures.

#### **LINES**

presents results of comparisons between all pairs of least squares means by listing the means in descending order and indicating nonsignificant subsets by line segments beside the corresponding LS-means. When all differences have the same variance, these comparison lines are guaranteed to accurately reflect the inferences that are based on the corresponding tests, which are made by comparing the respective p-values to the value of the ALPHA= option (0.05 by default). However, equal variances might not be the case for differences between LS-means. If the variances are not all the same, then the comparison lines might be conservative, in the sense that if you base your inferences on the lines alone, you will detect fewer significant differences than the tests indicate. If there are any such differences, the procedure lists the pairs of means that are inferred to be significantly different by the tests but not by the comparison lines. However, even though the variances in many cases are unequal, they are similar enough that the comparison lines accurately reflect the test inferences.

# **MEANS | NOMEANS**

determines whether to print the least squares means themselves. For most procedure, MEANS is the default behavior. For example, the NOMEANS option is the default for the PHREG procedure. You can then use the MEANS option to produce the table of least squares means, if desired.

#### **ODDSRATIO**

#### OR

requests that LS-mean differences (DIFF, ADJUST= options) are also reported in terms of odds ratios. The ODDSRATIO option is ignored unless you use either the logit, cumulative logit, or generalized logit link function. If you specify the CL or ALPHA= option, confidence intervals for the odds ratios are also computed. These intervals are adjusted for multiplicity when you specify the ADJUST= option.

The ODDSRATIO option is supported only by the procedures that support generalized linear modeling (GENMOD, LOGISTIC and SURVEYLOGISTIC) and by PROC PLM when it is used to perform statistical analyses on item stores created by these procedures.

## OBSMARGINS<=OM-data-set>

#### **OM**<=*OM*-data-set>

specifies a potentially different weighting scheme for the computation of LS-means coefficients. The standard LS-means have equal coefficients across classification effects; however, the OM option changes these coefficients to be proportional to those found in the *OM-data-set*. This adjustment is reasonable when you want your inferences to apply to a population that is not necessarily balanced but has the margins that are observed in *OM-data-set*.

By default, *OM-data-set* is the same as the analysis data set. You can optionally specify another data set that describes the population for which you want to make inferences. This data set must contain all model variables except for the dependent variable (which is ignored if it is present). In addition, the levels of all CLASS variables must be the same as those that occur in the analysis data set. If a level of a classification effect in the original data set is not present in the *OM-data-set*, the LS-means

for that level are undefined. The corresponding rows of the LSMeans table are displayed as missing. Specifying an *OM-data-set* enables you to construct arbitrarily weighted LS-means.

In computing the observed margins, the procedure uses all observations for which there are no missing or invalid independent variables, including those for which there are missing dependent variables. Also, if you use a WEIGHT statement, the procedure computes weighted margins to construct the LS-means coefficients. If your data are balanced, the LS-means are unchanged by the OM option.

The BYLEVEL option modifies the observed-margins LS-means. Instead of computing the margins across all of the *OM-data-set*, the procedure computes separate margins for each level of the LSMEANS effect in question. In this case the resulting LS-means are actually equal to raw means for fixed-effects models and certain balanced random-effects models, but their estimated standard errors account for the covariance structure that you have specified.

You can use the E option in conjunction with either the OM or BYLEVEL option to verify that the modified LS-means coefficients are the ones you want. It is possible that the modified LS-means are not estimable when the standard ones are estimable, or vice versa.

#### **PDIFF**

is the same as the DIFF option.

```
PLOT | PLOTS< = plot-request< (options) >>
```

```
PLOT | PLOTS< =(plot-request< (options) > < ... plot-request< (options) > >) >
```

requests that graphics related to least squares means be produced via ODS Graphics, provided that ODS Graphics is enabled and the *plot-request* does not conflict with other options in the LSMEANS statement. For general information about ODS Graphics, see Chapter 21, "Statistical Graphics Using ODS."

The available options and suboptions are as follows:

#### **ALL**

requests that the default plots that correspond to this LSMEANS statement be produced. The default plot depends on the options in the statement.

#### **ANOMPLOT**

## **ANOM**

requests an analysis-of-means display in which least squares means are compared to an average least squares mean. Least squares mean ANOM plots are produced only for those model effects that are listed in LSMEANS statements and have options that do not contradict with the display. For example, the following statements produce analysis-of-mean plots for effects A and C:

```
lsmeans A / diff=anom plot=anom;
lsmeans B / diff plot=anom;
lsmeans C / plot=anom;
```

The DIFF option in the second LSMEANS statement implies all pairwise differences.

## **BOXPLOT**< boxplot-options>

produces box plots of the distribution of the least squares mean or least squares mean differences across a posterior sample. For example, this plot is available in procedures that support a Bayesian analysis through the BAYES statement.

A separate box is generated for each estimable function, and all boxes appear on a single graph by default. You can affect the appearance of the box plot graph with the following options:

# ORIENTATION=VERTICAL | HORIZONTAL

# ORIENT=VERT | HORIZ

specifies the orientation of the boxes. The default is vertical orientation of the box plots.

#### **NPANELPOS**=number

specifies how to break the series of box plots across multiple panels. If the NPANELPOS option is not specified, or if *number* equals zero, then all box plots are displayed in a single graph; this is the default. If a negative number is specified, then exactly up to |number| of box plots are displayed per panel. If *number* is positive, then the number of boxes per panel is balanced to achieve small variation in the number of box plots per graph.

#### CONTROLPLOT

#### CONTROL

requests a display in which least squares means are visually compared against a reference level. These plots are produced only for statements with options that are compatible with control differences. For example, the following statements produce control plots for effects A and C:

```
lsmeans A / diff=control('1') plot=control;
lsmeans B / diff
                           plot=control;
1smeans C
                           plot=control;
```

The DIFF option in the second LSMEANS statement implies all pairwise differences.

```
DIFFPLOT< (diffplot-options) >
DIFFOGRAM< (diffplot-options) >
DIFF<(diffplot-options)>
```

requests a display of all pairwise least squares mean differences and their significance. The display is also known as a "mean-mean scatter plot" when it is based on arithmetic means (Hsu 1996; Hsu and Peruggia 1994). For each comparison a line segment, centered at the LS-means in the pair, is drawn. The length of the segment corresponds to the projected width of a confidence interval for the least squares mean difference. Segments that fail to cross the 45-degree reference line correspond to significant least squares mean differences.

LS-mean difference plots are produced only for statements with options that are compatible with the display. For example, the following statements request differences against a control level for the A effect, all pairwise differences for the B effect, and the least squares means for the C effect:

```
lsmeans A / diff=control('1') plot=diff;
1smeans B / diff
                              plot=diff;
1smeans C
                              plot=diff;
```

The DIFF= type in the first statement is incompatible with a display of all pairwise differences.

You can specify the following diffplot-options:

#### **ABS**

determines the positioning of the line segments in the plot. This is the default *diffplot-options*. When the ABS option is in effect, all line segments are shown on the same side of the reference line.

#### **NOABS**

determines the positioning of the line segments in the plot. The NOABS option separates comparisons according to the sign of the difference.

## **CENTER**

marks the center point for each comparison. This point corresponds to the intersection of two least squares means.

## **NOLINES**

suppresses the display of the line segments that represent the confidence bounds for the differences of the least squares means. The NOLINES option implies the CENTER option. The default is to draw line segments in the upper portion of the plot area without marking the center point.

# **DISTPLOT**< distplot-options>

# **DIST**< distplot-options >

generates panels of histograms with a kernel density overlaid if the analysis has access to a set of posterior parameter estimates. For example, this plot is available in procedures that support a Bayesian analysis through the BAYES statement. A separate plot in each panel contains the results for each least squares mean or least squares mean differences. You can specify the following *distplot-options* in parentheses:

#### **BOX | NOBOX**

controls the display of a horizontal box plot of the estimable function's distribution across the posterior sample below the graph. The BOX option is enabled by default.

#### **HIST | NOHIST**

controls the display of the histogram of the estimable function's distribution across the posterior sample. The HIST option is enabled by default.

# **NORMAL | NONORMAL**

controls the display of a normal density estimate on the graph. The NONORMAL option is enabled by default.

# KERNEL | NOKERNEL

controls the display of a kernel density estimate on the graph. The KERNEL option is enabled by default.

#### NROWS=number

specifies the highest number of rows in a panel. The default is 3.

# **NCOLS**=number

specifies the highest number of columns in a panel. The default is 3.

#### **UNPACK**

unpacks the panel into separate graphics.

# MEANPLOT< (meanplot-options) >

requests displays of the least squares means.

The following *meanplot-options* control the display of the least squares means.

## **ASCENDING**

displays the least squares means in ascending order. This option has no effect if means are displayed in separate plots.

## CL

displays upper and lower confidence limits for the least squares means. By default, 95% limits are drawn. You can change the confidence level with the ALPHA= option. Confidence limits are drawn by default if the CL option is specified in the LSMEANS statement.

#### **CLBAND**

displays confidence limits as bands. This option implies the JOIN option.

## **DESCENDING**

displays the least squares means in descending order. This option has no effect if means are displayed in separate plots.

#### **ILINK**

requests that means (and confidence limits) be displayed on the inverse linked scale.

#### **JOIN**

# CONNECT

connects the least squares means with lines. This option is implied by the CLBAND option. If the effect contains nested variables and a SLICEBY= effect contains classification variables that appear as crossed effects, this option is ignored.

#### SLICEBY=fixed-effect

specifies an effect by which to group the means in a single plot. For example, the following statement requests a plot in which the levels of A are placed on the horizontal axis and the means that belong to the same level of B are joined by lines:

# lsmeans A\*B / plot=meanplot(sliceby=b join);

Unless the LS-mean effect contains at least two classification variables, the SLICEBY= option has no effect. The *fixed-effect* does not have to be an effect in your MODEL statement, but it must consist entirely of classification variables and it must be contained in the LS-mean effect.

#### PLOTBY=fixed-effect

specifies an effect by which to break interaction plots into separate displays. For example, the following statement requests for each level of C one plot of the A\*B cell means that are associated with that level of C:

#### lsmeans A\*B\*C / plot=meanplot(sliceby=b plotby=c clband);

In each plot, levels of A are displayed on the horizontal axis, and confidence bands are drawn around the means that share the same level of B.

The PLOTBY= option has no effect unless the LS-mean effect contains at least three classification variables. The *fixed-effect* does not have to be an effect in the MODEL statement, but it must consist entirely of classification variables and it must be contained in the LS-mean effect.

## **NONE**

requests that no plots be produced.

When LS-mean calculations are adjusted for multiplicity by using the ADJUST= option, the plots are adjusted accordingly.

## SEED=number

specifies the seed for the sampling-based components of the computations for the LSMEANS statement (for example, chi-bar-square statistics and simulated *p*-values). The value of *number* must be an integer. The seed is used to start the pseudo-random-number generator for the simulation. If you do not specify a seed, or if you specify a value less than or equal to zero, the seed is generated from reading the time of day from the computer clock. Note that there could be multiple LSMEANS statements with SEED= specifications and there could be other statements that can supply a random number seed. Since the procedure has only one random number stream, the initial seed is shown in the SAS log.

# SINGULAR=number

tunes the estimability checking. If  $\mathbf{v}$  is a vector, define ABS( $\mathbf{v}$ ) to be the largest absolute value of the elements of  $\mathbf{v}$ . If ABS( $\mathbf{K}' - \mathbf{K}'\mathbf{T}$ ) is greater than  $c^*$ number for any row of  $\mathbf{K}'$  in the contrast, then  $\mathbf{K}'\boldsymbol{\beta}$  is declared nonestimable. Here,  $\mathbf{T}$  is the Hermite form matrix ( $\mathbf{X}'\mathbf{X}$ ) $^-\mathbf{X}'\mathbf{X}$ , and c is ABS( $\mathbf{K}'$ ), except when it equals 0, and then c is 1. The value for *number* must be between 0 and 1; the default is 1E–4.

# STEPDOWN<(step-down-options)>

requests that multiple comparison adjustments for the *p*-values of LS-mean differences be further adjusted in a step-down fashion. Step-down methods increase the power of multiple comparisons by taking advantage of the fact that a *p*-value is never declared significant unless all smaller *p*-values are also declared significant. The STEPDOWN adjustment combined with ADJUST=BON corresponds to the methods of Holm (1979) "Method 2" of Shaffer (1986); this is the default. Using step-down-adjusted *p*-values combined with ADJUST=SIMULATE corresponds to the method of Westfall (1997).

If the denominator degrees of freedom are computed by the Kenward-Roger (Kenward and Roger 1997) or Satterthwaite method in a mixed model, then step-down-adjusted *p*-values are produced only if the ADJDFE=ROW option is in effect.

Also, STEPDOWN affects only p-values, not confidence limits. For ADJUST=SIMULATE, the generalized least squares hybrid approach of Westfall (1997) is used to increase Monte Carlo accuracy.

You can specify the following *step-down-options* in parentheses:

# MAXTIME=n

specifies the time (in seconds) to be spent computing the maximal logically consistent sequential subsets of equality hypotheses for TYPE=LOGICAL. The default is MAXTIME=60. If the MAXTIME value is exceeded, the adjusted tests are not computed. When this occurs, you can try increasing the MAXTIME value. However, note that there are common multiple comparisons problems for which this computation requires a huge amount of time—for example, all pairwise comparisons between more than 10 groups. In such cases, try to use TYPE=FREE (the default) or TYPE=LOGICAL(n) for small n.

## **REPORT**

specifies that a report on the step-down adjustment be displayed, including a listing of the sequential subsets (Westfall 1997) and, for ADJUST=SIMULATE, the step-down simulation results.

# TYPE=LOGICAL<(n)>

#### TYPE=FREE

specifies how step-down adjustment are made. If you specify TYPE=LOGICAL, the step-down adjustments are computed by using maximal logically consistent sequential subsets of equality hypotheses (Shaffer 1986; Westfall 1997). Alternatively, for TYPE=FREE, sequential subsets are computed ignoring logical constraints. The TYPE=FREE results are more conservative than those for TYPE=LOGICAL, but they can be much more efficient to produce for many comparisons. For example, it is not feasible to take logical constraints between all pairwise comparisons of more than 10 groups. For this reason, TYPE=FREE is the default.

However, you can reduce the computational complexity of taking logical constraints into account by limiting the depth of the search tree used to compute them, specifying the optional depth parameter as a number n in parentheses after TYPE=LOGICAL. As with TYPE=FREE, results for TYPE=LOGICAL(n) are conservative relative to the true TYPE=LOGICAL results. But even for TYPE=LOGICAL(0) they can be appreciably less conservative than TYPE=FREE, and they are computationally feasible for much larger numbers of comparisons. If you do not specify n or if n = -1, the full search tree is used.

# **ODS Table Names: LSMEANS Statement**

Each table created by the LSMEANS statement has a name associated with it, and you can use this name to refer to the table when using the Output Delivery System (ODS) to select tables and create output data sets. These names are listed in Table 19.22. For more information about ODS, see Chapter 20, "Using the Output Delivery System."

Table 1912 See Tables 1 Teadest Sy the Zelliz tive statement		
<b>Table Name</b>	Description	Required Option
Coef	L matrix coefficients	Е
Diffs	Differences of LS-means	DIFF or ADJUST= or
		STEPDOWN
LSMeans	LS-means	Default
LSMLines	Lines display for LS-means	LINES

**Table 19.22** ODS Tables Produced by the LSMEANS statement

# **ODS Graphics: LSMEANS Statement**

This section describes the use of ODS Graphics for creating graphics that are related to LS-means in procedures that support the common LSMEANS or SLICE statement. There are two groups of available plots: those that can be produced by all procedures that support these two statements, and those that can be produced only in association with the three procedures that can perform Bayesian analysis (PROC GENMOD, PROC LIFEREG, and PROC PHREG). Plots that are associated with the Bayesian analysis are available via these procedures directly, and also by using PROC PLM with an item store that was created by these procedures.

Plots in the first group depict the LS-means and their differences; when LS-mean comparisons are adjusted for multiplicity by using the ADJUST= option, the plots are adjusted accordingly. To request plots in this group, ODS Graphics must be enabled and you must request plots with the appropriate PLOTS= option in the LSMEANS or SLICE statement. Plots in the second group depict the posterior sample distribution of LS-means and their differences. To request plots in this group, you must also use a BAYES statement with PROC GENMOD, PROC PHREG, or PROC LIFEREG, or you must use PROC PLM to perform statistical analysis on an item store that was saved from a Bayesian analysis.

For more information about ODS Graphics, see Chapter 21, "Statistical Graphics Using ODS." The available graphs are summarized in Table 19.23 and Table 19.24.

**Table 19.23** Graphs Produced by All Procedures That Support the Common LSMEANS or SLICE Statement

ODS Graph Name	<b>Plot Description</b>	Required Option
AnomPlot	Requests an analysis of means display in which least squares means are com- pared to an average least squares mean.	PLOTS=ANOM
ControlPlot	Requests a display in which least squares means are compared to a reference level.	PLOTS=CONTROL
DiffPlot	Displays all pairwise least squares mean differences and their significance. This plot is also known as a "meanmean scatter plot" when based on arithmetic means.	PLOTS=DIFF
MeanPlot	Displays least squares means.	PLOTS=MEANPLOT

**Table 19.24** Graphs Produced by Procedures That Support the LSMEANS or SLICE Statement and Bayesian Analysis

<b>ODS Graph Name</b>	<b>Plot Description</b>	Required Option
BoxPlot	Displays box plots of LS-means or LS-mean differences across a posterior sample.	PLOTS=BOXPLOT

Table 19.24 continued

Table Fell Committee		
<b>ODS Graph Name</b>	Plot Description	Required Option
DistPanel	Displays panels of histograms with kernel density curves overlaid. Each plot contains the results for the posterior sample of each LS-mean or LS-mean difference.	PLOTS=DISTPLOT
DistPlot	Displays a histogram with a kernel density curve overlaid. The plot contains the results for the posterior sample of the LS-mean or LS-mean difference.	PLOTS=DISTPLOT(UNPACK)

You can supply the same *plot-options* to the SLICE statement to produce these graphs. For details about the plot-options of the LSMEANS or SLICE statement, see the PLOTS= option in the section "LSMEANS Statement" on page 460. For more details about the DIFFPLOT in particular, see the section "Graphics for LS-Mean Comparisons" on page 3256 in Chapter 44, "The GLIMMIX Procedure."

# LSMESTIMATE Statement

This statement documentation applies to the following SAS/STAT procedures: GENMOD, LIFEREG, LOGISTIC, MIXED, ORTHOREG, PHREG, PLM, PROBIT, SURVEYLOGISTIC, SURVEYPHREG, and SURVEYREG. It also applies to the RELIABILITY procedure in SAS/QC software. The LSMESTIMATE statement in the GLIMMIX procedure is documented in Chapter 44, "The GLIMMIX Procedure."

The LSMESTIMATE statement provides a mechanism for obtaining custom hypothesis tests among least squares means. In contrast to the LSMEANS statement, the LSMESTIMATE statement does not produce the least squares means or their differences; instead, you can estimate any linear function of the least squares means (including the means themselves or their differences). In contrast to the linear functions that are constructed with the ESTIMATE statement, you do not specify coefficients for the individual parameter estimates. Instead, with the LSMESTIMATE statement you specify coefficients for the least squares means; these are then converted for you into estimable functions for the parameter estimates.

The LSMESTIMATE statement thus combines important and convenient features of the LSMEANS and the ESTIMATE statement. As with the LSMEANS statement, the following conditions are true:

- You need to specify only a single effect; the mapping into linear estimable functions in terms of the parameter estimates is performed by the procedure.
- You can use the AT=, BYLEVEL, and OBSMARGINS options to affect the computation of the underlying least squares means.

As with the ESTIMATE statement you can do the following:

- specify multiple-row linear combinations.
- perform multiplicity corrections to control the familywise Type I error probability with the ADJUST= option.
- construct general linear functions of the least squares means.
- perform joint *F* or chi-square tests with or without order restrictions through the JOINT option.
- rely on positional or nonpositional syntax to specify coefficients for linear functions. For details about using nonpositional syntax, see the section "Positional and Nonpositional Syntax for Coefficients in Linear Functions" on page 455.

The computation of an LSMESTIMATE involves two coefficient matrices. Suppose that there are  $n_l$  levels for a valid least squares means effect (an effect that is part of your model and consists of classification variables only). Then the LS-means are formed as  $\mathbf{L}_1 \widehat{\boldsymbol{\beta}}$ , where  $\mathbf{L}_1$  is a  $(n_l \times p)$  coefficient matrix. The  $(k \times n_l)$  coefficient matrix  $\mathbf{K}$  is formed from the *values* that you supply in the k rows of the LSMESTIMATE statement. The least squares means estimates then represent the  $(k \times 1)$  vector

$$KL_1\beta = L\beta$$

Because the analytic features and capabilities of the LSMESTIMATE statement are an amalgam of the LSMEANS and the ESTIMATE statement, the syntax of the statement follows the same pattern.

# **Syntax: LSMESTIMATE Statement**

```
LSMESTIMATE model-effect < 'label' > values < divisor=n> <, < 'label' > values < divisor=n> > <, ... > </ options>;
```

In contrast to a multirow estimate in the ESTIMATE statement, you specify only a single effect in the LSMESTIMATE statement. The row labels are optional and follow the *model-effect* specification. For example, the following statements fit a split-split-plot design and compare the average of the third and fourth LS-mean of the whole-plot factor A to the first LS-mean of the factor:

```
proc glimmix;
   class a b block;
   model y = a b a*b / s;
   random int a / sub=block;
   lsmestimate A 'al vs avg(a3,a4)' 2 0 -1 -1 divisor=2;
run;
```

The order in which coefficients are assigned to the least squares means corresponds to the order in which they are displayed in the "Least Squares Means" table. You can use the ELSM option to see how coefficients are matched to levels of the fixed effect.

The optional *divisor=n* specification enables you to assign a separate divisor to each row of the LSMES-TIMATE. You can also assign divisor values through the DIVISOR= option. See the description of the DIVISOR= option that follows for the interaction between the two ways of specifying divisors.

Table 19.25 summarizes important options in the LSMESTIMATE statement. All LSMESTIMATE options are subsequently discussed in alphabetical order.

 Table 19.25
 LSMESTIMATE Statement Options

Option	Description	
Construction and Co	omputation of LS-Means	
AT	Modifies covariate values in computing LS-means	
BYLEVEL	Computes separate margins	
DIVISOR=	Specifies a list of values to divide the coefficients	
OM=	Specifies the weighting scheme for LS-means computation as determined by a data set	
SINGULAR=	Tunes estimability checking	
Degrees of Freedom	and p-values	
ADJUST=	Determines the method for multiple-comparison adjustment of LS-means differences	
$ALPHA=\alpha$	Determines the confidence level $(1 - \alpha)$	
LOWER	Performs one-sided, lower-tailed inference	
STEPDOWN	Adjusts multiple-comparison <i>p</i> -values further in a step-down fashion	
TESTVALUE=	Specifies values under the null hypothesis for tests	
UPPER	Performs one-sided, upper-tailed inference	
Statistical Output		
CL	Constructs confidence limits for means and mean differences	
CORR	Displays the correlation matrix of LS-means	
COV	Displays the covariance matrix of LS-means	
E	Prints the L matrix	
ELSM	Prints the <b>K</b> matrix	
JOINT	Produces a joint <i>F</i> or chi-square test for the LS-means and LS-means differences	
PLOTS=	Requests graphs of means and mean comparisons	
SEED=	Specifies the seed for computations that depend on random numbers	
Generalized Linear I	Modeling	
CATEGORY=	Specifies how to construct estimable functions with multinomial data	
EXP	Exponentiates and displays LS-means estimates	
ILINK	Computes and displays estimates and standard errors of LS-means (but not differences) on the inverse linked scale	

You can specify the following options in the LSMESTIMATE statement after a slash (/):

# ADJDFE=SOURCE

# ADJDFE=ROW

specifies how denominator degrees of freedom are determined when *p*-values and confidence limits are adjusted for multiple comparisons with the ADJUST= option. When you do not specify the ADJDFE= option or when you specify ADJDFE=SOURCE, the denominator degrees of freedom for

multiplicity-adjusted results are the denominator degrees of freedom for the LS-mean effect in the "Type III Tests of Fixed Effects" table.

The ADJDFE=ROW setting is useful if you want multiplicity adjustments to take into account that denominator degrees of freedom are not constant across estimates. For example, this can be the case when the denominator degrees of freedom are computed by the Satterthwaite or Kenward-Roger method (Kenward and Roger 1997) in a mixed model.

The ADJDFE= option is not supported by the procedures that perform chi-square-based inference (GENMOD, LOGISTIC, PHREG and SURVEYLOGISTIC).

ADJUST=BON
ADJUST=SCHEFFE
ADJUST=SIDAK
ADJUST=SIMULATE<(simoptions)>

## ADJUST=T

requests a multiple comparison adjustment for the *p*-values and confidence limits for the LS-mean estimates. The adjusted quantities are produced in addition to the unadjusted *p*-values and confidence limits. Adjusted confidence limits are produced if the CL or ALPHA= option is in effect. For a description of the adjustments, see Chapter 45, "The GLM Procedure," and Chapter 67, "The MULTTEST Procedure," in addition to the documentation for the ADJUST= option in the LSMEANS statement.

Not all adjustment methods of the LSMEANS statement are available for the LSMESTIMATE statement. Multiplicity adjustments in the LSMEANS statement are designed specifically for differences of least squares means.

If you specify the STEPDOWN option, the p-values are further adjusted in a step-down fashion.

# **ALPHA**=number

requests that a t type confidence interval be constructed for each of the LS-means with confidence level 1 - number. The value of *number* must be between 0 and 1; the default is 0.05.

AT variable=value

AT (variable-list)=(value-list)

#### AT MEANS

modifies the values of the covariates used in computing LS-means. See the AT option in the LSMEANS statement for details.

# **BYLEVEL**

requests that the procedure compute separate margins for each level of the LSMEANS effect.

The standard LS-means have equal coefficients across classification effects. The BYLEVEL option changes these coefficients to be proportional to the observed margins. This adjustment is reasonable when you want your inferences to apply to a population that is not necessarily balanced but has the margins observed in the input data set. In this case, the resulting LS-means are actually equal to raw means for fixed-effects models and certain balanced random-effects models, but their estimated standard errors account for the covariance structure that you have specified. If a WEIGHT statement is specified, the procedure uses weighted margins to construct the LS-means coefficients.

If the AT option is specified, the BYLEVEL option disables it.

# CATEGORY=category-options

specifies how to construct estimates and multiplicity corrections for models with multinomial data (ordinal or nominal). This option is also important for constructing sets of estimable functions for F tests with the JOINT option.

The *category-options* indicate how response variable levels are treated in constructing the estimable functions. Possible value for the category-options are the following:

## **JOINT**

computes the estimable functions for every nonredundant category and treats them as a set. For example, a three-row LSMESTIMATE statement in a model with three response categories leads to six estimable functions.

## **SEPARATE**

computes the estimable functions for every nonredundant category in turn. For example, a three-row LSMESTIMATE statement in a model with three response categories leads to two sets of three estimable functions.

# quoted-value-list

computes the estimable functions only for the specified list of values. The list must consist of formatted values of the response categories.

For further details about using the CATEGORY= option in models for multinomial data, see the documentation for the CATEGORY= option in the ESTIMATE statement.

The CATEGORY= option is supported only by the procedures that support generalized linear modeling (GENMOD, LOGISTIC, and SURVEYLOGISTIC) and by PROC PLM when it is used to perform statistical analyses on item stores that were created by these procedures.

## **CHISQ**

requests that chi-square tests be performed in addition to F tests, when you request an F test with the JOINT option. This option has no effect in procedures that produce chi-square statistics by default.

# CL

requests that t type confidence limits be constructed for each of the LS-means. The confidence level is 0.95 by default; this can be changed with the ALPHA= option. If you specify an ADJUST= option, then the confidence limits are adjusted for multiplicity. But if you also specify STEPDOWN, then only p-values are step-down adjusted, not the confidence limits.

# **CORR**

displays the estimated correlation matrix of the linear combination of the least squares means.

#### COV

displays the estimated covariance matrix of the linear combination of the least squares means.

## **DF**=number

specifies the degrees of freedom for the tests and confidence limits. The option is not supported by the procedures that perform chi-square-based inference (GENMOD, LOGISTIC, PHREG, and SURVEYLOGISTIC).

#### **DIVISOR**=value-list

specifies a list of values by which to divide the coefficients so that fractional coefficients can be entered as integer numerators. If you do not specify *value-list*, a default value of 1.0 is assumed. Missing values in the *value-list* are converted to 1.0.

If the number of elements in *value-list* exceeds the number of rows of the estimate, the extra values are ignored. If the number of elements in *value-list* is less than the number of rows of the estimate, the last value in *value-list* is carried forward.

If you specify a row-specific divisor as part of the specification of the estimate row, this value multiplies the corresponding value in the *value-list*. For example, the following statement divides the coefficients in the first row by 8, and the coefficients in the third and fourth row by 3:

Coefficients in the second row are not altered.

Ε

requests that the L coefficients of the estimable function be displayed. These are the coefficients that apply to the fixed-effect parameter estimates. The E option displays the coefficients that you would need to enter in an equivalent ESTIMATE statement.

#### **ELSM**

requests that the K matrix coefficients be displayed. These are the coefficients that apply to the LS-means. This option is useful to ensure that you assigned the coefficients correctly to the LS-means.

# **EXP**

requests exponentiation of the least squares means estimate. When you model data with the logit link function and the estimate represents a log odds ratio, the EXP option produces an odds ratio. If you specify the CL or ALPHA= option, the (adjusted) confidence limits for the estimate are also exponentiated.

The EXP option is supported only by PROC PHREG, PROC SURVEYPHREG, the procedures that support generalized linear modeling (GENMOD, LOGISTIC, and SURVEYLOGISTIC), and by PROC PLM when it is used to perform statistical analyses on item stores that were created by these procedures.

#### ILINK

requests that the estimate and its standard error also be reported on the scale of the mean (the inverse linked scale). The computation of the inverse linked estimate depends on the estimation mode. For example, if the analysis is based on a posterior sample when a BAYES statement is present, the inversely linked estimate is the average of the inversely linked values across the sample of posterior parameter estimates. If the analysis is not based on a sample of parameter estimates, the procedure computes the value on the mean scale by applying the inverse link to the estimate.

The interpretation of the inversely linked quantity depends on the coefficients that are specified in your LSMESTIMATE statement and the link function. For example, in a model for binary data with logit

$$q = \frac{1}{1 + \exp\{-(\tau_1 - \tau_2)\}}$$

where  $\tau_1$  and  $\tau_2$  are the least squares means that are associated with the first two levels of the classification effect A:

```
proc logistic;
   class A / param=glm;
   model y = A / dist=binary link=logit;
   lsmestimate A 1 -1 / ilink;
run;
```

The quantity q is not the difference of the probabilities associated with the two levels,

$$\pi_1 - \pi_2 = \frac{1}{1 + \exp\{-\tau_1\}} - \frac{1}{1 + \exp\{-\tau_2\}}$$

The standard error of the inversely linked estimate is based on the delta method. If you also specify the CL or ALPHA= option, the procedure computes confidence intervals for the inversely linked estimate. These intervals are obtained by applying the inverse link to the confidence intervals on the linked scale.

The ILINK option is supported only by the procedures that support generalized linear modeling (GENMOD, LOGISTIC, and SURVEYLOGISTIC) and by PROC PLM when it is used to perform statistical analyses on item stores that were created by these procedures.

# JOINT<(joint-test-options)>

requests that a joint *F* or chi-square test be produced for the rows of the estimate. For more information about the simulation-based *p*-value calculation, see the section "Joint Hypothesis Tests with Complex Alternatives, the Chi-Bar-Square Statistic" on page 457. You can specify the following *joint-test-options* in parentheses:

#### $ACC = \gamma$

specifies the accuracy radius for determining the necessary sample size in the simulation-based approach of Silvapulle and Sen (2004) for tests with order restrictions. The value of  $\gamma$  must be strictly between 0 and 1; the default value is 0.005.

# EPS=€

specifies the accuracy confidence level for determining the necessary sample size in the simulation-based approach of Silvapulle and Sen (2004) for F tests with order restrictions. The value of  $\epsilon$  must be strictly between 0 and 1; the default value is 0.01.

#### LABEL='label'

assigns an identifying label to the joint test. If you do not specify a label, the first non-default label for the ESTIMATE rows is used to label the joint test.

# **NOEST**

# **ONLY**

performs only the joint test and suppresses other results from the ESTIMATE statement. This option is useful for emulating the CONTRAST statement that is available in other procedures.

#### NSAMP=n

specifies the number of samples for the simulation-based method of Silvapulle and Sen (2004). If n is not specified, it is constructed from the values of the ALPHA= $\alpha$ , the ACC= $\gamma$ , and the EPS= $\epsilon$  options. With the default values for  $\gamma$ ,  $\epsilon$ , and  $\alpha$  (0.005, 0.01, and 0.05, respectively), NSAMP=12,604 by default.

#### **CHISQ**

adds a chi-square test if the procedure produces an F test by default.

## **BOUNDS**=value-list

specifies boundary values for the estimable linear function. The null value of the hypothesis is always zero. If you specify a positive boundary value z, the hypotheses are  $H:\theta=0$ ,  $H_a:\theta>0$  with the added constraint that  $\theta< z$ . The same is true for negative boundary values. The alternative hypothesis is then  $H_a:\theta<0$  subject to the constraint  $\theta>-|z|$ . If you specify a missing value, the hypothesis is assumed to be two-sided. The BOUNDS option enables you to specify sets of one- and two-sided joint hypotheses. If all values in *value-list* are set to missing, the procedure performs a simulation-based p-value calculation for a two-sided test.

## **LOWER**

#### **LOWERTAILED**

requests that the *p*-value for the *t* test be based only on values that are less than the test statistic. A two-tailed test is the default. A lower-tailed confidence limit is also produced if you specify the CL or ALPHA= option.

Note that for ADJUST=SCHEFFE the one-sided adjusted confidence intervals and one-sided adjusted *p*-values are the same as the corresponding two-sided statistics, because this adjustment is based on only the right tail of the *F* distribution.

If you request an *F* test with the JOINT option, then a one-sided left-tailed order restriction is applied to all estimable functions, and the corresponding chi-bar-square statistic of Silvapulle and Sen (2004) is computed in addition to the two-sided, standard, *F* or chi-square statistic. See the JOINT option for how to control the computation of the simulation-based chi-bar-square statistic.

#### OBSMARGINS<=OM-data-set>

# OM<=OM-data-set>

specifies a potentially different weighting scheme for the computation of LS-means coefficients. The standard LS-means have equal coefficients across classification effects; however, the OM option changes these coefficients to be proportional to those found in the *OM-data-set*. This adjustment is reasonable when you want your inferences to apply to a population that is not necessarily balanced but has the margins observed in *OM-data-set*. See the OBSMARGINS option in the LSMEANS statement for further details.

## **PLOTS**=plot-options

produces ODS statistical graphics of the distribution of estimable functions if the procedure performs the analysis in a sampling-based mode. For example, this is the case when procedures support a BAYES statement and perform a Bayesian analysis. The estimable functions are then computed for each of the posterior parameter estimates, and the "Least Squares Means Estimates" table reports simple descriptive statistics for the evaluated functions. In this situation, the PLOTS= option enables you to visualize the distribution of the estimable function. The following *plot-options* are available:

#### **ALL**

produces all possible plots with their default settings.

# BOXPLOT<(boxplot-options)>

produces box plots of the distribution of the estimable function across the posterior sample. A separate box plot is generated for each estimable function and all box plots appear on a single graph by default. You can affect the appearance of the box plot graph with the following options:

# ORIENTATION=VERTICAL | HORIZONTAL

# ORIENT=VERT | HORIZ

specifies the orientation of the boxes. The default is vertical orientation of the box plots.

#### **NPANELPOS**=number

specifies how to break the series of box plots across multiple panels. If the NPANELPOS option is not specified, or if *number* equals zero, then all box plots are displayed in a single graph; this is the default. If a negative number is specified, then exactly up to |number| of box plots are displayed per panel. If *number* is positive, then the number of boxes per panel is balanced to achieve small variation in the number of box plots per graph.

# **DISTPLOT**<(distplot-options)>

# DIST<(distplot-options)>

generates panels of histograms with a kernel density overlaid. A separate plot in each panel contains the results for each estimable function. You can specify the following *distplot-options* in parentheses:

## **BOX | NOBOX**

controls the display of a horizontal box plot below the histogram. The BOX option is enabled by default.

# HIST | NOHIST

controls the display of the histogram of the estimable function's distribution across the posterior sample. The HIST option is enabled by default.

# NORMAL | NONORMAL

controls the display of a normal density estimate on the graph. The NONORMAL option is enabled by default.

# **KERNEL | NOKERNEL**

controls the display of a kernel density estimate on the graph. The KERNEL option is enabled by default.

#### **NROWS**=number

specifies the highest number of rows in a panel. The default is 3.

# NCOLS=number

specifies the highest number of columns in a panel. The default is 3.

# **UNPACK**

unpacks the panel into separate graphics.

#### NONE

does not produce any plots.

#### SEED=number

specifies the seed for the sampling-based components of the computations for the LSMESTIMATE statement (for example, chi-bar-square statistics and simulated *p*-values). The value of *number* must be an integer. The seed is used to start the pseudo-random-number generator for the simulation. If you do not specify a seed, or if you specify a value less than or equal to zero, the seed is generated from reading the time of day from the computer clock. Note that there could be multiple LSMESTIMATE statements with SEED= specifications and there could be other statements that can supply a random number seed. Since the procedure has only one random number stream, the initial seed is shown in the SAS log.

# SINGULAR=number

tunes the estimability checking as documented for the SINGULAR= option in the ESTIMATE statement.

# STEPDOWN<(step-down-options)>

requests that multiplicity adjustments for the *p*-values of estimable functions be further adjusted in a step-down fashion. Step-down methods increase the power of multiple testing procedures by taking advantage of the fact that a *p*-value is never declared significant unless all smaller *p*-values are also declared significant. The STEPDOWN adjustment combined with ADJUST=BON corresponds to the methods of Holm (1979) and "Method 2" of Shaffer (1986); this is the default. Using step-down-adjusted *p*-values combined with ADJUST=SIMULATE corresponds to the method of Westfall (1997).

If the ESTIMATE statement is applied with a STEPDOWN option in a mixed model where the degrees-of-freedom method is that of Kenward and Roger (1997) or of Satterthwaite, then step-down-adjusted *p*-values are produced only if the ADJDFE=ROW option is in effect.

Also, the STEPDOWN option affects only *p*-values, not confidence limits. For ADJUST=SIMULATE, the generalized least squares hybrid approach of Westfall (1997) is used to increase Monte Carlo accuracy.

You can specify the following *step-down-options* in parentheses:

## **MAXTIME**=*n*

specifies the time (in seconds) to be spent computing the maximal logically consistent sequential subsets of equality hypotheses for TYPE=LOGICAL. The default is MAXTIME=60. If the MAXTIME value is exceeded, the adjusted tests are not computed. When this occurs, you can try increasing the MAXTIME value. However, note that there are common multiple comparisons problems for which this computation requires a huge amount of time—for example, all pairwise comparisons between more than 10 groups. In such cases, try to use TYPE=FREE (the default) or TYPE=LOGICAL(n) for small n.

# ORDER=PVALUE

# **ORDER=ROWS**

specifies the order in which the step-down tests are performed. ORDER=PVALUE is the default, with LS-mean estimates being declared significant only if all LS-mean estimates with smaller (unadjusted) *p*-values are significant. If you specify ORDER=ROWS, then significances are evaluated in the order in which they are specified.

#### **REPORT**

specifies that a report on the step-down adjustment be displayed, including a listing of the sequential subsets (Westfall 1997) and, for ADJUST=SIMULATE, the step-down simulation results.

# TYPE=LOGICAL<(n)> TYPE=FREE

# specifies how step-down adjustment are made. If you specify TYPE=LOGICAL, the step-down adjustments are computed by using maximal logically consistent sequential subsets of equality hypotheses (Shaffer 1986; Westfall 1997). Alternatively, for TYPE=FREE, sequential subsets

are computed ignoring logical constraints. The TYPE=FREE results are more conservative than those for TYPE=LOGICAL, but they can be much more efficient to produce for many estimates. For example, it is not feasible to take logical constraints between all pairwise comparisons of more than about 10 groups. For this reason, TYPE=FREE is the default. However, you can reduce the computational complexity of taking logical constraints into account

by limiting the depth of the search tree used to compute them, specifying the optional depth parameter as a number n in parentheses after TYPE=LOGICAL. As with TYPE=FREE, results for TYPE=LOGICAL(n) are conservative relative to the true TYPE=LOGICAL results. But even for TYPE=LOGICAL(0), they can be appreciably less conservative than TYPE=FREE, and they are computationally feasible for much larger numbers of estimates. If you do not specify n or if n =-1, the full search tree is used.

## TESTVALUE=value-list

#### TESTMEAN=value-list

specifies the value under the null hypothesis for testing the estimable functions in the LSMESTIMATE statement. The rules for specifying the *value-list* are very similar to those for specifying the divisor list in the DIVISOR= option. If no TESTVALUE= is specified, all tests are performed as  $H: L\beta = 0$ . Missing values in the *value-list* also are translated to zeros. If you specify fewer values than rows in the LSMESTIMATE statement, the last value in *value-list* is carried forward.

The TESTVALUE= option affects only p-values from individual, joint, and multiplicity-adjusted tests. It does not affect confidence intervals.

The TESTVALUE option is not available for the multinomial distribution, and the values are ignored when you perform a sampling-based (Bayesian) analysis.

## **UPPER**

#### **UPPERTAILED**

requests that the p-value for the t test be based only on values that are greater than the test statistic. A two-tailed test is the default. An upper-tailed confidence limit is also produced if you specify the CL or ALPHA= option.

Note that for ADJUST=SCHEFFE the one-sided adjusted confidence intervals and one-sided adjusted p-values are the same as the corresponding two-sided statistics, because this adjustment is based on only the right tail of the F distribution.

If you request a joint test with the JOINT option, then a one-sided right-tailed order restriction is applied to all estimable functions, and the corresponding chi-bar-square statistic of Silvapulle and Sen (2004) is computed in addition to the two-sided, standard, F or chi-square statistic. See the JOINT option for how to control the computation of the simulation-based chi-bar-square statistic.

#### **ODS Table Names: LSMESTIMATE Statement**

Each table created by the LSMESTIMATE statement has a name associated with it, and you can use this name to refer to the table when using the Output Delivery System (ODS) to select tables and create output data sets. These names are listed in Table 19.26. For more information about ODS, see Chapter 20, "Using the Output Delivery System."

**Table 19.26** ODS Tables Produced by the LSMESTIMATE statement

Table Name	Description	Required Option
Coef	L matrix coefficients or K matrix coefficients	E or ELSM
LSMEstimates Contrasts	Estimates among LS-means Joint test results for LS-means esti- mates	Default JOINT

## **ODS Graphics: LSMESTIMATE Statement**

This section describes the use of ODS for creating statistical graphs of the distribution of LS-means and LS-mean differences with the LSMESTIMATE statement. The plots can be produced only in association with the three procedures that can perform Bayesian analysis (PROC GENMOD, PROC LIFEREG, and PROC PHREG). The plots are available via these procedures directly, and also via PROC PLM when run using an item store that was created by these procedures. To request these graphs, you must do the following:

- · ensure that ODS Graphics is enabled
- use a BAYES statement with PROC GENMOD, PROC LIFEREG, or PROC PHREG, or use PROC PLM to perform statistical analysis on an item store that was saved from a Bayesian analysis
- request plots with the PLOTS= option in the LSMESTIMATE statement

For more information about ODS Graphics, see Chapter 21, "Statistical Graphics Using ODS." The available graphs are summarized in Table 19.27.

**Table 19.27** Graphs Produced by the LSMESTIMATE statement

ODS Graph Name	Plot Description	Required Option
BoxPlot	Displays box plots of LS-means or LS-mean differences across a posterior sample.	PLOTS=BOXPLOT

Table 19.27 continued

Table 19121 Contained		
<b>ODS Graph Name</b>	Plot Description	Required Option
DistPanel	Displays panels of histograms with kernel density curves overlaid. Each plot contains the results for the posterior sample of each LS-mean or LS-mean difference.	PLOTS=DISTPLOT
DistPlot	Displays a histogram with a kernel density curve overlaid. The plot contains the results for the posterior sample of the LS-mean or LS-mean difference.	PLOTS=DISTPLOT(UNPACK)

For details about the *plot-options* of the LSMESTIMATE statement, see the PLOTS= option in the section "LSMESTIMATE Statement" on page 476.

## **NLOPTIONS Statement**

This section applies to the following procedures:

CALIS, GLIMMIX, HPMIXED, PHREG, SURVEYPHREG, and VARIOGRAM. See the individual procedure chapters for deviations from the common syntax and defaults shown here.

## Syntax: NLOPTIONS Statement

The NLOPTIONS statement provides you with syntax to control aspects of the nonlinear optimizations in the CALIS, GLIMMIX, HPMIXED, PHREG, SURVEYPHREG, and VARIOGRAM procedures.

#### **NLOPTIONS** < options>;

The nonlinear optimization options are described in alphabetical order after Table 19.28, which summarizes the options by category. The notation used in describing the options is generic in the sense that  $\psi$  denotes the  $p \times 1$  vector of parameters for the optimization and  $\psi_i$  is its ith element. The objective function being minimized, its  $p \times 1$  gradient vector, and its  $p \times p$  Hessian matrix are denoted as  $f(\psi)$ ,  $g(\psi)$ , and  $H(\psi)$ , respectively. The gradient with respect to the *i*th parameter is denoted as  $g_i(\psi)$ . Superscripts in parentheses denote the iteration count; for example,  $f(\psi)^{(k)}$  is the value of the objective function at iteration k. In the mixed model procedures, the parameter vector  $\psi$  might consist of fixed effects only, covariance parameters only, or fixed effects and covariance parameters. In the CALIS procedure,  $\psi$  consists of all independent parameters that are defined in the models and in the PARAMETERS statement.

 Table 19.28
 Options to Control Aspects of the Optimization

	Described Control Aspects of the Optimization	
Option	Description	
Optimization		
HESCAL=	Determines the type of Hessian scaling	
INHESSIAN=	Specifies the start for approximated Hessian	
LINESEARCH=	Specifies the line-search method	
LSPRECISION=	Specifies the line-search precision	
RESTART=	Specifies the iteration number for update restart	
TECHNIQUE=	Determines the minimization technique	
UPDATE=	Determines the update technique	
Termination Criteria		
ABSCONV=	Tunes an absolute function convergence criterion	
ABSFCONV=	Tunes an absolute function difference convergence criterion	
ABSGCONV=	Tunes the absolute gradient convergence criterion	
ABSXCONV=	Tunes the absolute parameter convergence criterion	
FCONV=	Tunes the relative function convergence criterion	
FCONV2=	Tunes another relative function convergence criterion	
FSIZE=	Specifies the value used in the FCONV and GCONV criteria	
GCONV=	Tunes the relative gradient convergence criterion	
GCONV2=	Tunes another relative gradient convergence criterion	
MAXFUNC=	Specifies the maximum number of function calls	
MAXITER=	Specifies the maximum number of iterations	
MAXTIME=	Specifies the upper limit for seconds of CPU time	
MINITER=	Specifies the minimum number of iterations	
XCONV=	Specifies the relative parameter convergence criterion	
XSIZE=	Specifies the value used in the XCONV criterion	
Step Length		
DAMPSTEP=	Dampens steps in a line search	
INSTEP=	Specifies the initial trust region radius	
MAXSTEP=	Specifies the maximum trust region radius	
Printed Output		
PALL	Displays (almost) all printed output	
PHISTORY	Displays optimization history	
NOPRINT	Suppresses all printed output	
Covariance Matrix Toler	rances	
ASINGULAR=	Specifies the absolute singularity for inertia	
MSINGULAR=	Specifies the relative M singularity for inertia	
VSINGULAR=	Specifies the relative V singularity for inertia	

lable 19.2	28 continuea
Option	Description
<b>Constraint Specification</b>	ons
LCEPSILON=	Specifies the range for active constraints
LCDEACT=	Specifies the LM tolerance for deactivating
LCSINGULAR=	Specifies the tolerance for dependent constraints
Remote Monitoring	
SOCKET=	Specifies the fileref for remote monitoring

## ABSCONV=r

#### ABSTOL=r

specifies an absolute function convergence criterion: for minimization, termination requires  $f(\psi^{(k)}) \le$ r. The default value of r is the negative square root of the largest double-precision value, which serves only as a protection against overflows.

#### ABSFCONV = r < n >

#### ABSFTOL=r<n>

specifies an absolute function difference convergence criterion:

• For all techniques except NMSIMP (specified by the TECHNIQUE= option), termination requires a small change of the function value in successive iterations,

$$|f(\boldsymbol{\psi}^{(k-1)}) - f(\boldsymbol{\psi}^{(k)})| \le r$$

• The same formula is used for the NMSIMP technique, but  $\psi(k)$  is defined as the vertex with the lowest function value, and  $\psi^{(k-1)}$  is defined as the vertex with the highest function value in the simplex.

The default value is r = 0. The optional integer value n specifies the number of successive iterations for which the criterion must be satisfied before the process can be terminated.

#### ABSGCONV = r < n >

#### ABSGTOL=r < n >

specifies an absolute gradient convergence criterion:

• For all techniques except NMSIMP (specified by the TECHNIQUE= option), termination requires the maximum absolute gradient element to be small:

$$\max_{j} |g_{j}(\boldsymbol{\psi}^{(k)})| \leq r$$

• This criterion is not used by the NMSIMP technique.

The default value is r = 1E-5. The optional integer value n specifies the number of successive iterations for which the criterion must be satisfied before the process can be terminated.

#### ABSXCONV = r < n >

#### ABSXTOL=r<n>

specifies an absolute parameter convergence criterion:

• For all techniques except NMSIMP, termination requires a small Euclidean distance between successive parameter vectors,

$$\| \psi^{(k)} - \psi^{(k-1)} \|_2 \le r$$

• For the NMSIMP technique, termination requires either a small length  $\alpha^{(k)}$  of the vertices of a restart simplex,

$$\alpha^{(k)} \leq r$$

or a small simplex size,

$$\delta^{(k)} \leq r$$

where the simplex size  $\delta^{(k)}$  is defined as the L1 distance from the simplex vertex  $\boldsymbol{\xi}^{(k)}$  with the smallest function value to the other p simplex points  $\boldsymbol{\psi}_{l}^{(k)} \neq \boldsymbol{\xi}^{(k)}$ :

$$\delta^{(k)} = \sum_{\boldsymbol{\psi}_l \neq y} \| \boldsymbol{\psi}_l^{(k)} - \boldsymbol{\xi}^{(k)} \|_1$$

The default is r = 1E-8 for the NMSIMP technique and r = 0 otherwise. The optional integer value n specifies the number of successive iterations for which the criterion must be satisfied before the process can terminate.

#### ASINGULAR=r

#### ASING=r

specifies an absolute singularity criterion for the computation of the inertia (number of positive, negative, and zero eigenvalues) of the Hessian and its projected forms. The default value is the square root of the smallest positive double-precision value.

#### DAMPSTEP<=r>

specifies that the initial step length value  $\alpha^{(0)}$  for each line search (used by the QUANEW, CONGRA, or NEWRAP technique) cannot be larger than r times the step length value used in the former iteration. If the DAMPSTEP option is specified but r is not specified, the default is r=2. The DAMPSTEP= option can prevent the line-search algorithm from repeatedly stepping into regions where some objective functions are difficult to compute or where they could lead to floating-point overflows during the computation of objective functions and their derivatives. The DAMPSTEP= option can save time-consuming function calls during the line searches of objective functions that result in very small steps.

#### FCONV=r<n>

#### **FTOL**=*r*<*n*>

specifies a relative function convergence criterion:

 For all techniques except NMSIMP, termination requires a small relative change of the function value in successive iterations,

$$\frac{|f(\boldsymbol{\psi}^{(k)}) - f(\boldsymbol{\psi}^{(k-1)})|}{\max(|f(\boldsymbol{\psi}^{(k-1)})|, \text{FSIZE})} \le r$$

where FSIZE is defined by the FSIZE= option.

• The same formula is used for the NMSIMP technique, but  $\psi^{(k)}$  is defined as the vertex with the lowest function value and  $\psi^{(k-1)}$  is defined as the vertex with the highest function value in the simplex.

The default is  $r = 10^{-\text{FDIGITS}}$ , where FDIGITS is by default  $-\log_{10}\{\epsilon\}$  and  $\epsilon$  is the machine precision. Some procedures, such as the GLIMMIX procedure, enable you to change the value with the FDIGITS= option in the PROC statement. The optional integer value n specifies the number of successive iterations for which the criterion must be satisfied before the process can terminate.

#### **FCONV2=***r*<*n*>

#### **FTOL2**=*r*<*n*>

specifies a second function convergence criterion:

• For all techniques except NMSIMP, termination requires a small predicted reduction,

$$df^{(k)} \approx f(\boldsymbol{\psi}^{(k)}) - f(\boldsymbol{\psi}^{(k)} + \mathbf{s}^{(k)})$$

of the objective function. The predicted reduction

$$df^{(k)} = -\mathbf{g}^{(k)'}\mathbf{s}^{(k)} - \frac{1}{2}\mathbf{s}^{(k)'}\mathbf{H}^{(k)}\mathbf{s}^{(k)}$$
$$= -\frac{1}{2}\mathbf{s}^{(k)'}\mathbf{g}^{(k)} \le r$$

is computed by approximating the objective function f by the first two terms of the Taylor series and substituting the Newton step,

$$\mathbf{s}^{(k)} = -[\mathbf{H}^{(k)}]^{-1}\mathbf{g}^{(k)}$$

• For the NMSIMP technique, termination requires a small standard deviation of the function values of the p+1 simplex vertices  $\psi_l^{(k)}$ ,  $l=0,\ldots,p$ ,

$$\sqrt{\frac{1}{n+1} \sum_{l} \left[ f(\boldsymbol{\psi}_{l}^{(k)}) - \overline{f}(\boldsymbol{\psi}^{(k)}) \right]^{2}} \leq r$$

where  $\overline{f}(\psi^{(k)}) = \frac{1}{p+1} \sum_{l} f(\psi_{l}^{(k)})$ . If there are  $p_{act}$  boundary constraints active at  $\psi^{(k)}$ , the mean and standard deviation are computed only for the  $n+1-p_{act}$  unconstrained vertices.

The default value is r = 1E-6 for the NMSIMP technique and r = 0 otherwise. The optional integer value n specifies the number of successive iterations for which the criterion must be satisfied before the process can terminate.

#### FSIZE=r

specifies the FSIZE parameter of the relative function and relative gradient termination criteria. The default value is r = 0. For more details, see the FCONV= and GCONV= options.

#### GCONV=r<n>

#### GTOL=r < n >

specifies a relative gradient convergence criterion:

• For all techniques except CONGRA and NMSIMP, termination requires that the normalized predicted function reduction be small,

$$\frac{\mathbf{g}(\boldsymbol{\psi}^{(k)})'[\mathbf{H}^{(k)}]^{-1}\mathbf{g}(\boldsymbol{\psi}^{(k)})}{\max(|f(\boldsymbol{\psi}^{(k)})|, \text{FSIZE})} \leq r$$

where FSIZE is defined by the FSIZE= option. For the CONGRA technique (where a reliable Hessian estimate **H** is not available), the following criterion is used:

$$\frac{\parallel \mathbf{g}(\boldsymbol{\psi}^{(k)}) \parallel_2^2 \ \parallel \mathbf{s}(\boldsymbol{\psi}^{(k)}) \parallel_2}{\parallel \mathbf{g}(\boldsymbol{\psi}^{(k)}) - \mathbf{g}(\boldsymbol{\psi}^{(k-1)}) \parallel_2 \max(|f(\boldsymbol{\psi}^{(k)})|, \text{FSIZE})} \leq r$$

• This criterion is not used by the NMSIMP technique.

The default value is r = 1E-8. The optional integer value n specifies the number of successive iterations for which the criterion must be satisfied before the process can terminate.

#### GCONV2=r<n>

#### GTOL2=r < n >

specifies another relative gradient convergence criterion:

• For least squares problems and the TRUREG, LEVMAR, NRRIDG, and NEWRAP techniques, the following criterion of Browne (1982) is used:

$$\max_{j} \frac{|\mathbf{g}_{j}(\boldsymbol{\psi}^{(k)})|}{\sqrt{f(\boldsymbol{\psi}^{(k)})\mathbf{H}_{i,j}^{(k)}}} \leq r$$

• This criterion is not used by the other techniques.

The default value is r = 0. The optional integer value n specifies the number of successive iterations for which the criterion must be satisfied before the process can terminate.

#### HESCAL=0 | 1 | 2 | 3

#### HS=0 | 1 | 2 | 3

specifies the scaling version of the Hessian (or crossproduct Jacobian) matrix used in NRRIDG, TRUREG, LEVMAR, NEWRAP, or DBLDOG optimization.

If HS is not equal to 0, the first iteration and each restart iteration set the diagonal scaling matrix  $D^{(0)} = \text{diag}(d_i^{(0)})$ :

$$d_i^{(0)} = \sqrt{\max(|H_{i,i}^{(0)}|, \epsilon)}$$

where  $H_{i,i}^{(0)}$  are the diagonal elements of the Hessian (or crossproduct Jacobian). In every other iteration, the diagonal scaling matrix  $D^{(0)} = \operatorname{diag}(d_i^{(0)})$  is updated depending on the HS option:

**0** specifies that no scaling be done.

specifies the Moré (1978) scaling update:

$$d_i^{(k+1)} = \max \left\lceil d_i^{(k)}, \sqrt{\max(|H_{i,i}^{(k)}|, \epsilon)} \right\rceil$$

2 specifies the Dennis, Gay, and Welsch (1981) scaling update:

$$d_i^{(k+1)} = \max \left[ 0.6 * d_i^{(k)}, \sqrt{\max(|H_{i,i}^{(k)}|, \epsilon)} \right]$$

**3** specifies that  $d_i$  be reset in each iteration:

$$d_i^{(k+1)} = \sqrt{\max(|H_{i,i}^{(k)}|, \epsilon)}$$

In each scaling update,  $\epsilon$  is the relative machine precision. The default value is HS=0. Scaling of the Hessian can be time-consuming in the case where general linear constraints are active.

#### INHESSIAN < = r >

#### INHESS < =r >

specifies how the initial estimate of the approximate Hessian is defined for the quasi-Newton techniques QUANEW and DBLDOG. There are two alternatives:

- If you do not use the r specification, the initial estimate of the approximate Hessian is set to the Hessian at  $\psi^{(0)}$ .
- If you do use the r specification, the initial estimate of the approximate Hessian is set to the multiple of the identity matrix r**I**.

By default (if you do not specify the option INHESSIAN=r), the initial estimate of the approximate Hessian is set to the multiple of the identity matrix r**I**, where the scalar r is computed from the magnitude of the initial gradient.

#### INSTEP=r

#### SALPHA=r

#### RADIUS=r

reduces the length of the first trial step during the line search of the first iterations. For highly nonlinear objective functions, such as the EXP function, the default initial radius of the trust-region algorithm TRUREG or DBLDOG or the default step length of the line-search algorithms can result in arithmetic overflows. If this occurs, you should specify decreasing values of 0 < r < 1 such as INSTEP=1E-1, INSTEP=1E-2, INSTEP=1E-4, and so on, until the iteration starts successfully.

- For trust-region algorithms (TRUREG or DBLDOG), the INSTEP= option specifies a factor r > 0 for the initial radius  $\Delta^{(0)}$  of the trust region. The default initial trust-region radius is the length of the scaled gradient. This step corresponds to the default radius factor of r = 1.
- For line-search algorithms (NEWRAP, CONGRA, or QUANEW), the INSTEP= option specifies an upper bound for the initial step length for the line search during the first five iterations. The default initial step length is r = 1.
- For the Nelder-Mead simplex algorithm, by using TECH=NMSIMP, the INSTEP=*r* option defines the size of the start simplex.

#### LCDEACT=r

#### LCD=r

specifies a threshold *r* for the Lagrange multiplier that determines whether an active inequality constraint remains active or can be deactivated. For maximization, *r* must be greater than zero; for minimization, *r* must be smaller than zero. An active inequality constraint can be deactivated only if its Lagrange multiplier is less than the threshold value. The default value is

$$r = \pm \min(0.01, \max(0.1 \times ABSGCONV, 0.001 \times \operatorname{gmax}^{(k)}))$$

where "+" is for maximization, "-" is for minimization, ABSGCONV is the value of the absolute gradient criterion, and  $gmax^{(k)}$  is the maximum absolute element of the gradient or the projected gradient.

#### LCEPSILON=r

#### LCEPS=r

#### LCE=r

specifies the range r for active and violated boundary constraints, where  $r \ge 0$ . If the point  $\psi^{(k)}$  satisfies the following condition, the constraint i is recognized as an active constraint:

$$|\sum_{j=1}^{k} a_{ij} \psi_j^{(k)} - b_i| \le r \times (|b_i| + 1)$$

Otherwise, the constraint i is either an inactive inequality or a violated inequality or equality constraint. The default value is r = 1E-8. During the optimization process, the introduction of rounding errors can force the optimization to increase the value of r by a factor of  $10^k$  for some k > 0. If this happens, it is indicated by a message displayed in the log.

#### LCSINGULAR=r

#### LCSING=r

#### LCS=r

specifies a criterion r, where  $r \ge 0$ , that is used in the update of the QR decomposition and that determines whether an active constraint is linearly dependent on a set of other active constraints. The default value is r = 1E-8. The larger r becomes, the more the active constraints are recognized as being linearly dependent. If the value of r is larger than 0.1, it is reset to 0.1.

#### LINESEARCH=i

#### LIS=i

specifies the line-search method for the CONGRA, QUANEW, and NEWRAP optimization techniques. See Fletcher (1987) for an introduction to line-search techniques. The value of i can be  $1, \ldots, 8$  as follows. The default is LIS=2.

- specifies a line-search method that needs the same number of function and gradient calls for cubic interpolation and cubic extrapolation; this method is similar to one used by the Harwell subroutine library.
- specifies a line-search method that needs more function than gradient calls for quadratic and cubic interpolation and cubic extrapolation; this method is implemented as shown in Fletcher (1987) and can be modified to an exact line search by using the LSPRECISION= option. This is the default.

3	specifies a line-search method that needs the same number of function and gradient calls for cubic interpolation and cubic extrapolation; this method is implemented as shown in Fletcher (1987) and can be modified to an exact line search by using the LSPRECISION= option.
4	specifies a line-search method that needs the same number of function and gradient calls for stepwise extrapolation and cubic interpolation.
5	specifies a line-search method that is a modified version of LIS=4.
6	specifies a golden-section line search (Polak 1971), which uses only function values for linear approximation.
7	specifies a bisection line search (Polak 1971), which uses only function values for linear approximation.
8	specifies the Armijo line-search technique (Polak 1971), which uses only function values for linear approximation.

#### LSPRECISION=r

#### LSP=r

specifies the degree of accuracy that should be obtained by the line-search algorithms LIS=2 and LIS=3. Usually an imprecise line search is inexpensive and successful. For more difficult optimization problems, a more precise and expensive line search might be necessary (Fletcher 1987). The LIS=2 line-search method (which is the default for the NEWRAP, QUANEW, and CONGRA techniques) and the LIS=3 line-search method approach exact line search for small LSPRECISION= values. If you have numerical problems, try to decrease the LSPRECISION= value to obtain a more precise line search. The default values are shown in Table 19.29.

TECH=	UPDATE=	LSP Default
QUANEW	DBFGS, BFGS	r = 0.4
QUANEW	DDFP, DFP	r = 0.06
CONGRA	All	r = 0.1
NEWRAP	No update	r = 0.9

 Table 19.29
 Default Values for Line-Search Precision

For more details, see Fletcher (1987).

#### MAXFUNC=i

#### MAXFU=i

specifies the maximum number *i* of function calls in the optimization process. The default values are as follows:

- 125 for the TRUREG, NRRIDG, NEWRAP, and LEVMAR techniques
- 500 for the QUANEW and DBLDOG techniques
- 1000 for the CONGRA technique
- 3000 for the NMSIMP technique

Optimization can terminate only after completing a full iteration. Therefore, the number of function calls that are actually performed can exceed the number that is specified by the MAXFUNC= option.

#### MAXITER=i

#### MAXIT=i

specifies the maximum number *i* of iterations in the optimization process. The default values are as follows:

- 50 for the TRUREG, NRRIDG, NEWRAP, and LEVMAR techniques
- 200 for the QUANEW and DBLDOG techniques
- 400 for the CONGRA technique
- 1000 for the NMSIMP technique

These default values are also valid when *i* is specified as a missing value.

#### MAXSTEP=r<n>

specifies an upper bound for the step length of the line-search algorithms during the first *n* iterations. By default, *r* is the largest double-precision value and *n* is the largest integer available. Setting this option can improve the speed of convergence for the CONGRA, QUANEW, and NEWRAP techniques.

#### MAXTIME=r

specifies an upper limit of *r* seconds of CPU time for the optimization process. The time specified by the MAXTIME= option is checked only once at the end of each iteration. Therefore, the actual running time can be much longer than that specified by the MAXTIME= option. The actual running time includes the rest of the time needed to finish the iteration and the time needed to generate the output of the results. By default, CPU time is not limited.

#### MINITER=i

#### MINIT=i

specifies the minimum number of iterations. The default value is 0. If you request more iterations than are actually needed for convergence to a stationary point, the optimization algorithms can behave strangely. For example, the effect of rounding errors can prevent the algorithm from continuing for the required number of iterations.

#### MSINGULAR=r

#### MSING=r

specifies a relative singularity criterion r, where r > 0, for the computation of the inertia (number of positive, negative, and zero eigenvalues) of the Hessian and its projected forms. The default value is 1E-12.

#### **NOPRINT**

suppresses output that is related to optimization, such as the iteration history. This option, along with all NLOPTIONS statement options for displayed output, are ignored by the GLIMMIX and HPMIXED procedures.

#### **PALL**

displays all optional output for optimization. This option is supported only by the CALIS and SURVEYPHREG procedures.

#### **PHISTORY**

#### **PHIST**

displays the optimization history. The PHISTORY option is implied if the PALL option is specified. The PHISTORY option is supported only by the CALIS and SURVEYPHREG procedures.

#### RESTART=i

#### REST=i

specifies that the QUANEW or CONGRA technique is restarted with a steepest search direction after at most i iterations, where i > 0. Default values are as follows:

- When TECHNIQUE=CONGRA and UPDATE=PB, restart is performed automatically; so i is not used.
- When TECHNIQUE=CONGRA and UPDATE  $\neq$  PB,  $i = \min(10p, 80)$ , where p is the number of parameters.
- When TECHNIQUE=QUANEW, *i* is the largest integer available.

#### SINGULAR=r

#### SING=r

specifies the singularity criterion r, 0r < 1, that is used for the inversion of the Hessian matrix. The default value is 1E-8.

#### SOCKET=fileref

specifies the fileref that contains the information needed for remote monitoring.

#### **TECHNIQUE**=value

#### TECH=value

#### OMETHOD=value

#### OM=value

specifies the optimization technique. You can find additional information about choosing an optimization technique in the section "Choosing an Optimization Algorithm" on page 501. Valid values for the TECHNIQUE= option are as follows:

#### CONGRA

performs a conjugate-gradient optimization, which can be more precisely specified with the UPDATE= option and modified with the LINESEARCH= option. When you specify this option, UPDATE=PB by default.

#### DBLDOG

performs a version of double-dogleg optimization, which can be more precisely specified with the UPDATE= option. When you specify this option, UPDATE=DBFGS by default.

#### LEVMAR

performs a highly stable, but for large problems memory- and time-consuming, Levenberg-Marquardt optimization technique, a slightly improved variant of the Moré (1978) implementation. You can also specify this technique with the alias LM or MARQUARDT. In the CALIS procedure, this is the default optimization technique if there are fewer than 40 parameters to estimate. The GLIMMIX and HPMIXED procedures do not support this optimization technique.

#### NMSIMP

performs a Nelder-Mead simplex optimization. The CALIS procedure does not support this optimization technique.

#### NONE

does not perform any optimization. This option can be used for the following:

- to perform a grid search without optimization
- to compute estimates and predictions that cannot be obtained efficiently with any of the optimization techniques
- to obtain inferences for known values of the covariance parameters

#### NEWRAP

performs a Newton-Raphson optimization that combines a line-search algorithm with ridging. The line-search algorithm LIS=2 is the default method.

#### NRRIDG

performs a Newton-Raphson optimization with ridging. This is the default optimization technique in the SURVEYPHREG procedure.

#### OUANEW

performs a quasi-Newton optimization, which can be defined more precisely with the UPDATE= option and modified with the LINESEARCH= option.

#### TRUREG

performs a trust-region optimization.

#### **UPDATE**=method

#### **UPD**=method

specifies the update method for the quasi-Newton, double-dogleg, or conjugate-gradient optimization technique. Not every update method can be used with each optimizer.

The following are the valid methods for the UPDATE= option:

#### BFGS

performs the original Broyden, Fletcher, Goldfarb, and Shanno (BFGS) update of the inverse Hessian matrix.

#### DBFGS

performs the dual BFGS update of the Cholesky factor of the Hessian matrix. This is the default update method.

#### DDFP

performs the dual Davidon, Fletcher, and Powell (DFP) update of the Cholesky factor of the Hessian matrix.

#### DFP

performs the original DFP update of the inverse Hessian matrix.

#### PB

performs the automatic restart update method of Powell (1977) and Beale (1972).

#### • FR

performs the Fletcher-Reeves update (Fletcher 1987).

- PR performs the Polak-Ribiere update (Fletcher 1987).
- CD performs a conjugate-descent update of Fletcher (1987).

#### VERSION=1 | 2

#### VS=1 | 2

specifies the version of the quasi-Newton optimization technique with nonlinear constraints.

VS=1 specifies the update of the  $\mu$  vector as in Powell (1978b, a) (update like VF02AD).

VS=2 specifies the update of the  $\mu$  vector as in Powell (1982b, a) (update like VMCWD).

The default is VERSION=2.

#### VSINGULAR=r

#### VSING=r

specifies a relative singularity criterion r, where r > 0, for the computation of the inertia (number of positive, negative, and zero eigenvalues) of the Hessian and its projected forms. The default value is r = 1E-8.

#### XCONV=r<n>

#### XTOL=r < n >

specifies the relative parameter convergence criterion:

• For all techniques except NMSIMP, termination requires a small relative parameter change in subsequent iterations:

$$\frac{\max_{j} |\psi_{j}^{(k)} - \psi_{j}^{(k-1)}|}{\max(|\psi_{j}^{(k)}|, |\psi_{j}^{(k-1)}|, \mathsf{XSIZE})} \leq r$$

• For the NMSIMP technique, the same formula is used, but  $\psi_j^{(k)}$  is defined as the vertex with the lowest function value and  $\psi_j^{(k-1)}$  is defined as the vertex with the highest function value in the simplex.

The default value is r = 1E-8 for the NMSIMP technique and r = 0 otherwise. The optional integer value n specifies the number of successive iterations for which the criterion must be satisfied before the process can be terminated.

#### XSIZE=r

specifies the XSIZE parameter r of the relative parameter termination criterion, where  $r \ge 0$ . The default value is r = 0. For more details, see the XCONV= option.

## **Choosing an Optimization Algorithm**

#### **First- or Second-Order Algorithms**

The factors that go into choosing a particular optimization technique for a particular problem are complex. Trial and error can be involved.

For many optimization problems, computing the gradient takes more computer time than computing the function value. Computing the Hessian sometimes takes much more computer time and memory than computing the gradient, especially when there are many decision variables. Unfortunately, optimization techniques that do not use some kind of Hessian approximation usually require many more iterations than techniques that do use a Hessian matrix, and, as a result, the total run time of these techniques is often longer. Techniques that do not use the Hessian also tend to be less reliable. For example, they can terminate more easily at stationary points than at global optima.

Table 19.30 shows which derivatives are required for each optimization technique.

Algorithm	First-Order	Second-Order
LEVMAR	X	X
TRUREG	X	X
NEWRAP	X	X
NRRIDG	X	X
QUANEW	X	-
DBLDOG	X	-
CONGRA	X	-
NMSIMP	-	-

**Table 19.30** Derivatives Required

The second-derivative methods TRUREG, NEWRAP, and NRRIDG are best for small problems where the Hessian matrix is not expensive to compute. Sometimes the NRRIDG algorithm can be faster than the TRUREG algorithm, but TRUREG can be more stable. The NRRIDG algorithm requires only one matrix with p(p+1)/2 double words; TRUREG and NEWRAP require two such matrices. Here, p denotes the number of parameters in the optimization.

The first-derivative methods QUANEW and DBLDOG are best for medium-sized problems where the objective function and the gradient are much faster to evaluate than the Hessian. In general, the QUANEW and DBLDOG algorithms require more iterations than TRUREG, NRRIDG, and NEWRAP, but each iteration can be much faster. The QUANEW and DBLDOG algorithms require only the gradient to update an approximate Hessian, and they require slightly less memory than TRUREG or NEWRAP (essentially one matrix with p(p+1)/2 double words).

The first-derivative method CONGRA is best for large problems where the objective function and the gradient can be computed much faster than the Hessian and where too much memory is required to store the (approximate) Hessian. In general, the CONGRA algorithm requires more iterations than QUANEW or DBLDOG, but each iteration can be much faster. Because CONGRA requires only a factor of p double-word memory, many large applications can be solved only by CONGRA.

Each optimization method uses one or more convergence criteria that determine when it has converged. An algorithm is considered to have converged when any one of the convergence criteria is satisfied. For example, under the default settings, the QUANEW algorithm will converge if ABSGCONV < 1E–5, FCONV <  $10^{-\mathrm{FDIGITS}}$ , or GCONV < 1E–8.

### **Algorithm Descriptions**

#### Trust Region Optimization (TRUREG)

The trust region method uses the gradient  $\mathbf{g}(\boldsymbol{\psi}^{(k)})$  and the Hessian matrix  $\mathbf{H}(\boldsymbol{\psi}^{(k)})$ ; thus, it requires that the objective function  $f(\boldsymbol{\psi})$  have continuous first- and second-order derivatives inside the feasible region.

The trust region method iteratively optimizes a quadratic approximation to the nonlinear objective function within a hyperelliptic trust region with radius  $\Delta$  that constrains the step size that corresponds to the quality of the quadratic approximation. The trust region method is implemented based on Dennis, Gay, and Welsch (1981); Gay (1983) and Moré and Sorensen (1983).

The trust region method performs well for small- to medium-sized problems, and it does not need many function, gradient, and Hessian calls. However, if the computation of the Hessian matrix is computationally expensive, one of the (dual) quasi-Newton or conjugate gradient algorithms might be more efficient.

#### Newton-Raphson Optimization with Line Search (NEWRAP)

The NEWRAP technique uses the gradient  $\mathbf{g}(\boldsymbol{\psi}^{(k)})$  and the Hessian matrix  $\mathbf{H}(\boldsymbol{\psi}^{(k)})$ ; thus, it requires that the objective function have continuous first- and second-order derivatives inside the feasible region. If second-order derivatives are computed efficiently and precisely, the NEWRAP method can perform well for medium-sized to large problems, and it does not need many function, gradient, and Hessian calls.

This algorithm uses a pure Newton step when the Hessian is positive definite and when the Newton step reduces the value of the objective function successfully. Otherwise, a combination of ridging and line search is performed to compute successful steps. If the Hessian is not positive definite, a multiple of the identity matrix is added to the Hessian matrix to make it positive definite (Eskow and Schnabel 1991).

In each iteration, a line search is performed along the search direction to find an approximate optimum of the objective function. The default line-search method uses quadratic interpolation and cubic extrapolation (LIS=2).

#### Newton-Raphson Ridge Optimization (NRRIDG)

The NRRIDG technique uses the gradient  $\mathbf{g}(\boldsymbol{\psi}^{(k)})$  and the Hessian matrix  $\mathbf{H}(\boldsymbol{\psi}^{(k)})$ ; thus, it requires that the objective function have continuous first- and second-order derivatives inside the feasible region.

This algorithm uses a pure Newton step when the Hessian is positive definite and when the Newton step reduces the value of the objective function successfully. If at least one of these two conditions is not satisfied, a multiple of the identity matrix is added to the Hessian matrix.

The NRRIDG method performs well for small- to medium-sized problems, and it does not require many function, gradient, and Hessian calls. However, if the computation of the Hessian matrix is computationally expensive, one of the (dual) quasi-Newton or conjugate gradient algorithms might be more efficient.

Because the NRRIDG technique uses an orthogonal decomposition of the approximate Hessian, each iteration of NRRIDG can be slower than that of the NEWRAP technique, which works with a Cholesky decomposition. Usually, however, NRRIDG requires fewer iterations than NEWRAP.

The (dual) quasi-Newton method uses the gradient  $g(\psi^{(k)})$ , and it does not need to compute second-order derivatives because they are approximated. It works well for medium-sized to moderately large optimization problems, where the objective function and the gradient are much faster to compute than the Hessian. However, in general, it requires more iterations than the TRUREG, NEWRAP, and NRRIDG techniques, which compute second-order derivatives. QUANEW is the default optimization algorithm because it provides an appropriate balance between the speed and stability required for most nonlinear mixed model applications.

The QUANEW technique is one of the following, depending upon the value of the UPDATE= option:

- the original quasi-Newton algorithm, which updates an approximation of the inverse Hessian
- the dual quasi-Newton algorithm, which updates the Cholesky factor of an approximate Hessian (this is the default)

You can specify four update formulas with the UPDATE= option:

- DBFGS performs the dual Broyden, Fletcher, Goldfarb, and Shanno (BFGS) update of the Cholesky factor of the Hessian matrix. This is the default.
- DDFP performs the dual Davidon, Fletcher, and Powell (DFP) update of the Cholesky factor of the Hessian matrix.
- BFGS performs the original BFGS update of the inverse Hessian matrix.
- DFP performs the original DFP update of the inverse Hessian matrix.

In each iteration, a line search is performed along the search direction to find an approximate optimum. The default line-search method uses quadratic interpolation and cubic extrapolation to obtain a step size  $\alpha$  that satisfies the Goldstein conditions. One of the Goldstein conditions can be violated if the feasible region defines an upper limit of the step size. Violating the left-side Goldstein condition can affect the positive definiteness of the quasi-Newton update. In that case, either the update is skipped or the iterations are restarted with an identity matrix, resulting in the steepest descent or ascent search direction. You can specify line-search algorithms other than the default with the LIS= option.

The QUANEW algorithm uses its own line-search technique. Of the options and parameters that control the line search for other algorithms, only the INSTEP= option applies here. In several applications, large steps in the first iterations are troublesome. You can use the INSTEP= option to impose an upper bound for the step size  $\alpha$  during the first five iterations. You can also use the INHESSIAN= option to specify a different starting approximation for the Hessian. If you specify only the INHESSIAN option, the Cholesky factor of a (possibly ridged) finite-difference approximation of the Hessian is used to initialize the quasi-Newton update process.

#### Double-Dogleg Optimization (DBLDOG)

The double-dogleg optimization method combines the ideas of the quasi-Newton and trust region methods. In each iteration, the double-dogleg algorithm computes the step  $\mathbf{s}^{(k)}$  as the linear combination of the steepest descent or ascent search direction  $\mathbf{s}_1^{(k)}$  and a quasi-Newton search direction  $\mathbf{s}_2^{(k)}$ ,

$$s^{(k)} = \alpha_1 s_1^{(k)} + \alpha_2 s_2^{(k)}$$

The step is requested to remain within a prespecified trust region radius; see Fletcher (1987, p, 107). Thus, the DBLDOG subroutine uses the dual quasi-Newton update but does not perform a line search. You can specify two update formulas with the UPDATE= option:

- DBFGS performs the dual Broyden, Fletcher, Goldfarb, and Shanno update of the Cholesky factor of the Hessian matrix. This is the default.
- DDFP performs the dual Davidon, Fletcher, and Powell update of the Cholesky factor of the Hessian matrix.

The double-dogleg optimization technique works well for medium-sized to moderately large optimization problems, where the objective function and the gradient are much faster to compute than the Hessian. The implementation is based on Dennis and Mei (1979); Gay (1983), but it is extended for dealing with boundary and linear constraints. The DBLDOG technique generally requires more iterations than the TRUREG, NEWRAP, and NRRIDG techniques, which require second-order derivatives; however, each of the DBLDOG iterations is computationally cheap. Furthermore, the DBLDOG technique requires only gradient calls for the update of the Cholesky factor of an approximate Hessian.

#### Conjugate Gradient Optimization (CONGRA)

Second-order derivatives are not required by the CONGRA algorithm and are not even approximated. The CONGRA algorithm can be expensive in function and gradient calls, but it requires only O(p) memory for unconstrained optimization. In general, many iterations are required to obtain a precise solution, but each of the CONGRA iterations is computationally cheap. You can specify four different update formulas for generating the conjugate directions by using the UPDATE= option:

- PB performs the automatic restart update method of Powell (1977) and Beale (1972). This is the
- FR performs the Fletcher-Reeves update (Fletcher 1987).
- PR performs the Polak-Ribiere update (Fletcher 1987).
- CD performs a conjugate-descent update of Fletcher (1987).

The default often behaves best for typical examples, whereas UPDATE=CD can perform poorly.

The CONGRA subroutine should be used for optimization problems with large p. For the unconstrained or boundary-constrained case, CONGRA requires only O(p) bytes of working memory, whereas all other optimization methods require order  $O(p^2)$  bytes of working memory. During p successive iterations, uninterrupted by restarts or changes in the working set, the conjugate gradient algorithm computes a cycle of p conjugate search directions. In each iteration, a line search is performed along the search direction to find an approximate optimum of the objective function. The default line-search method uses quadratic interpolation and cubic extrapolation to obtain a step size  $\alpha$  that satisfies the Goldstein conditions. One of the Goldstein conditions can be violated if the feasible region defines an upper limit for the step size. Other line-search algorithms can be specified with the LIS= option.

#### Nelder-Mead Simplex Optimization (NMSIMP)

The Nelder-Mead simplex method does not use any derivatives and does not assume that the objective function has continuous derivatives. The objective function itself needs to be continuous. This technique is quite expensive in the number of function calls, and it might be unable to generate precise results for  $p \gg 40$ .

The original Nelder-Mead simplex algorithm is implemented and extended to boundary constraints. This algorithm does not compute the objective for infeasible points, but it changes the shape of the simplex adapting to the nonlinearities of the objective function, which contributes to an increased speed of convergence. It uses a special termination criterion.

## **SLICE Statement**

This statement applies to the following SAS/STAT procedures:

GENMOD, GLIMMIX, LIFEREG, LOGISTIC, MIXED, ORTHOREG, PHREG, PLM, PROBIT, SUR-VEYLOGISTIC, SURVEYPHREG, and SURVEYREG. It also applies to the RELIABILITY procedure in SAS/QC software.

The SLICE statement is similar to the LSMEANS statement. You use it to perform inferences on model effects that consist entirely of classification variables. With the SLICE statement, these effects must be higher-order effects of at least two classification variables. The effect is then partitioned into subsets that correspond to variables used in forming the effect. You can use the same options as you use for the LSMEANS statement to perform an analysis for the partitions. This analysis is also known as an analysis of simple effects (Winer 1971).

By default, the interaction effect is partitioned by all main effects. For example, the following statements produce simple-effect differences among the A levels for each level of B and simple-effect differences among the B levels for each level of A:

```
class a b;
model y = a b a*b;
slice a*b / diff nof;
```

For example, if the *model-effect* is a three-way interaction effect, the default output includes comparisons of the two-way interaction means.

Suppose, for example, that the interaction effect A\*B is significant in your analysis and that you want to test the effect of A for each level of B. The appropriate statement is

```
slice A*B / sliceBy = B;
```

This produces an F test for each level of B that compares the equality of the levels of A.

For example, assume that in a balanced design factors A and B have a = 4 and b = 3 levels, respectively. Consider the following statements:

```
class a b;
model y = a b a*b;
slice a*b / sliceby=a diff;
```

The SLICE statement produces four F tests, one per level of A. The first of these tests is constructed by extracting the three rows that correspond to the first level of A from the coefficient matrix for the A\*B interaction. Call this matrix  $\mathbf{L}_{a1}$  and its rows  $\mathbf{l}_{a1}^{(1)}$ ,  $\mathbf{l}_{a1}^{(2)}$ , and  $\mathbf{l}_{a1}^{(3)}$ . The slice tests the two-degrees-of-freedom hypothesis

$$H: \left\{ \begin{array}{l} \left( \mathbf{l}_{a1}^{(1)} - \mathbf{l}_{a1}^{(2)} \right) \boldsymbol{\beta} = 0 \\ \left( \mathbf{l}_{a1}^{(1)} - \mathbf{l}_{a1}^{(3)} \right) \boldsymbol{\beta} = 0 \end{array} \right.$$

In a balanced design, where  $\mu_{ij}$  denotes the mean response if A is at level i and B is at level j, this hypothesis is equivalent to  $H: \mu_{11} = \mu_{12} = \mu_{13}$ . The DIFF option considers the three rows of  $L_{a1}$  in turn and performs tests of the difference between pairs of rows. By default, all pairwise differences within the subset of L are considered; in the example this corresponds to tests of the form

$$H: \left(\mathbf{l}_{a1}^{(1)} - \mathbf{l}_{a1}^{(2)}\right) \boldsymbol{\beta} = 0$$

$$H: \left(\mathbf{l}_{a1}^{(1)} - \mathbf{l}_{a1}^{(3)}\right) \boldsymbol{\beta} = 0$$

$$H: \left(\mathbf{l}_{a1}^{(2)} - \mathbf{l}_{a1}^{(3)}\right) \boldsymbol{\beta} = 0$$

In the example, with a = 4 and b = 3, this produces four sets of least squares means differences. Within each set, factor A is held fixed at a particular level and each set consists of three comparisons.

## Syntax: SLICE Statement

```
SLICE model-effect < / options > ;
```

You can specify all options of the LSMEANS statement in the SLICE statement. The philosophy of the SLICE statement is to apply the analysis according to the options to the subsets of the L matrix that correspond to chosen partitions.

The following behavior differences between the SLICE and the LSMEANS statement are noteworthy:

- The specification of the *model-effect* is optional in the LSMEANS statement and required in the SLICE statement.
- Only a single SLICE *model-effect* can be specified before the option slash (/). However, you can specify multiple partitioning rules with the SLICEBY option.
- The MEANS option is the default for most procedures in the LSMEANS statement. For the SLICE statement, the default is the NOMEANS option.

Also, the three generalized linear modeling options: EXP, ILINK, and ODDSRATIO in the SLICE statement are additionally supported by PROC GLIMMIX and by PROC PLM when it is used to perform statistical analyses on item stores that were created by PROC GLIMMIX.

In addition to the options in the LSMEANS statement, you can specify the following options in the SLICE statement after the slash (/):

```
SLICEBY <=> slice-specification

SIMPLE <=> slice-specification

SLICEBY(slice-specification <, slice-specification <, ...>>)

SIMPLE(slice-specification <, slice-specification <, ...>>)
```

determines how to construct the partition of the least squares means for the *model-effect*. A *slice-specification* consists of an effect name followed by an optional list of formatted values. For example, the following statements creates partitions of the A\*B interaction effect for all levels of variable A:

```
class a b;
model y = a b a*b;
slice a*b / sliceby=a;
```

The following statements produces two partitions of the interaction:

```
class a b;
model y = a b a*b;
slice a*b / sliceby(b='2' a='1') diff;
```

In the first partition the variable B takes on formatted value '2'. In the second partition the variable A takes on the formatted value '1'.

#### **NOF**

suppresses the F test for testing the mutual equality of the estimable functions in the partition.

#### **ODS Table Names: SLICE Statement**

Each table created by the SLICE statement has a name associated with it, and you can use this name to refer to the table when using the Output Delivery System (ODS) to select tables and create output data sets. These names are listed in Table 19.31. For more information about ODS, see Chapter 20, "Using the Output Delivery System."

Table Name	Description	Required Option
Coef	L matrix coefficients	Е
Slices	LS-means slices	MEANS
SliceDiffs	Simple differences of LS-means slices	DIFF or ADJUST= or
		STEPDOWN or NOF
SliceLines	Lines display for LS-means slices	LINES
SliceTests	Tests for LS-means slices	Default

Table 19.31 ODS Tables Produced by the SLICE statement

This statement applies to the following SAS/STAT procedures: GENMOD, GLIMMIX, GLM, GLMSELECT, LIFEREG, LOGISTIC, MIXED, ORTHOREG, PHREG, PROBIT, SURVEYLOGISTIC, SURVEYPHREG, and SURVEYREG. It also applies to the RELIABILITY procedure in SAS/QC software.

The STORE statement requests that the procedure save the context and results of the statistical analysis into an item store. An item store is a binary file format that cannot be modified by the user. The contents of the item store can be processed with the PLM procedure. One example of item store technology is to perform a time-consuming analysis and to store its results by using the STORE statement. At a later time you can then perform specific statistical analysis tasks based on the saved results of the previous analysis, without having to fit the model again. The following statements show an example in which a mixed model is fit with the MIXED procedure and the postprocessing analysis is performed with the PLM procedure:

```
proc mixed data=MyBigDataSet;
   class Env A B sub;
   model y = A B x / ddfm=KenwardRoger;
   random int A*B / sub=Env;
   repeated / subject=Env*A*B type=AR(1);
   store sasuser.mixed;
run;

proc plm restore=sasuser.mixed;
   show cov Parms;
   lsmeans A B / diff;
   score data=NewData out=ScoreResults;
run;
```

The STORE statement in the PROC MIXED step requests that the MIXED procedure save those results that are needed to perform statistical tasks with the PLM procedure. For example, the MIXED procedure saves the necessary pieces of information that relate to the Kenward-Roger degree-of-freedom method. The results from the LSMEANS statement in the PROC PLM step thus apply this technique for calculating denominator degrees of freedom. The SHOW statement in the PLM procedure reveals the contents of the item store in terms of ODS tables, and the SCORE statement computes predicted values in a new data set. For more information about postprocessing tasks based on item stores, see the documentation for the PLM procedure.

## **Syntax: STORE Statement**

```
STORE < OUT = > item-store-name < / LABEL = 'label' > ;
```

The *item-store-name* is a usual one- or two-level SAS name, like the names that are used for SAS data sets. If you specify a one-level name, then the item store resides in the WORK library and is deleted at the end of the SAS session. Since item stores usually are used to perform postprocessing tasks, typical usage specifies a two-level name of the form libname.membername.

If an item store by the same name as specified in the STORE statement already exists, the existing store is replaced.

You can add a custom label with the LABEL= option in the STORE statement after the slash (/). When the PLM procedure processes an item store, the label appears in the PROC PLM output along with other identifying information.

## **TEST Statement**

This statement documentation applies to the following procedures: ICPHREG, LIFEREG, ORTHOREG, PLM, PROBIT, SURVEYPHREG, and SURVEYREG. It also applies to the RELIABILITY procedure in SAS/QC software.

The TEST statement enables you to perform *F* tests for model effects that test Type I, II, or Type III hypotheses. See Chapter 15, "The Four Types of Estimable Functions," for details about the construction of Type I, II, and III estimable functions.

## **Syntax: TEST Statement**

**TEST** < model-effects > </ options > ;

Table 19.32 summarizes options in the TEST statement.

Option **Description CHISO** Requests chi-square tests DDF= Specifies denominator degrees of freedom for fixed effects E Requests Type I, Type II, and Type III coefficients E1 Requests Type I coefficients Requests Type II coefficients E2 Requests Type III coefficients E3 HTYPE= Indicates the type of hypothesis test to perform Adds a row that corresponds to the overall intercept INTERCEPT

 Table 19.32
 TEST Statement Options

You can specify the following options in the TEST statement after the slash (/):

#### **CHISQ**

requests that chi-square tests be performed for the relevant effects in addition to the F tests. Type III tests are the default; you can produce the Type I and Type II tests by using the HTYPE= option. This option has no effect when the procedure produces chi-square statistics by default.

#### DDF=value-list

#### DF=value-list

specifies the denominator degrees of freedom for the fixed effects. The *value-list* specification is a list of numbers or missing values (.) separated by commas. The order of degrees of freedom should match the order of the fixed effects that are specified in the TEST statement; otherwise it should match the order in which the effects appear in the "Type III Tests of Fixed Effects" table. If you want to retain the default degrees of freedom for a particular effect, use a missing value for its location in the list. In the following example, the first TEST statement assigns 3 denominator degrees of freedom to A and 4.7 to A\*B, while those for B remain the same, and the second TEST statement assigns 5 denominator degrees of freedom to A and uses the default degrees of freedom for B.

```
model Y = A B A*B;
test / ddf=3,.,4.7;
test B A / ddf=.,5;
```

Ε

requests that Type I, Type II, and Type III L matrix coefficients be displayed for all relevant effects.

#### E1 | EI

requests that Type I L matrix coefficients be displayed for all relevant effects.

#### **E2 | EII**

requests that Type II L matrix coefficients be displayed for all relevant effects.

#### E3 | EIII

requests that Type III L matrix coefficients be displayed for all relevant effects.

#### HTYPE=value-list

indicates the type of hypothesis test to perform on the fixed effects. Valid entries for values in the *value-list* are 1, 2, and 3, which correspond to Type I, Type II, and Type III tests, respectively. The default value is 3.

#### **INTERCEPT**

#### INT

adds a row to the tables for Type I, II, and III tests that correspond to the overall intercept.

#### **ODS Table Names: TEST Statement**

Each table created by the TEST statement has a name associated with it, and you can use this name to refer to the table when using the Output Delivery System (ODS) to select tables and create output data sets. These names are listed in Table 19.33. For more information about ODS, see Chapter 20, "Using the Output Delivery System."

Table Name	Description	Required Option
Coef	L matrix coefficients	E
Tests1	Type I tests of fixed effects	HTYPE=1
Tests2	Type II tests of fixed effects	HTYPE=2
Tests3	Type III tests of fixed effects	Default

**Table 19.33** ODS Tables Produced by the TEST statement

## **Programming Statements**

This section applies to the following procedures:

CALIS, GLIMMIX, MCMC, NLIN, NLMIXED, PHREG, and SURVEYPHREG.

The majority of the SAS/STAT modeling procedures can take advantage of the fact that the statistical model can easily be translated into programming syntax (statements and options). However, several procedures require additional flexibility in specifying models—for example, when the model contains general nonlinear functions, when it is necessary to specify complicated restrictions, or when user-supplied expressions need to be evaluated. Procedures that are listed at the beginning of the section support—in addition to the usual procedure statements and options—programming statements that can be used in the SAS DATA step.

The following are valid statements:

```
ABORT;
ARRAY arrayname < [ dimensions ] > < $ > < variables-and-constants >;
CALL name < (expression < , expression . . . >)>;
DO < variable = expression < TO expression > < BY expression >>
   <, expression < TO expression > < BY expression >> . . .
    < WHILE expression > < UNTIL expression >;
END;
GOTO statement-label;
IF expression;
IF expression THEN program-statement;
             ELSE program-statement;
variable = expression;
variable + expression;
LINK statement-label;
PUT < variable > < = > ...;
RETURN;
SELECT < (expression) >;
STOP;
SUBSTR(variable, index, length)= expression;
WHEN (expression)program-statement;
       OTHERWISE program-statement;
```

For the most part, these programming statements work the same as they do in the SAS DATA step, as documented in SAS Language Reference: Concepts. However, there are several differences:

- The ABORT statement does not allow any arguments.
- The DO statement does not allow a character index variable. Thus

```
do i = 1, 2, 3;
```

is supported, whereas the following statement is not supported:

```
do i = 'A', 'B', 'C';
```

• Not all procedures support LAG functionality. For example, the GLIMMIX procedure does not support lags.

- The PUT statement, used mostly for program debugging, supports only some of the features of the DATA step PUT statement, and it has some features that are not available with the DATA step PUT statement:
  - The PUT statement does not support line pointers, factored lists, iteration factors, overprinting, INFILE, the colon (:) format modifier, or "\$".
  - The PUT statement does support expressions, but the expression must be enclosed in parentheses. For example, the following statement displays the square root of x:

```
put (sqrt(x));
```

- The PUT statement supports the item PDV to display a formatted listing of all variables in the program. For example:

```
put _pdv_;
```

• The WHEN and OTHERWISE statements enable you to specify more than one target statement. That is, DO/END groups are not necessary for multiple-statement WHENs. For example, the following syntax is valid:

```
select:
   when (exp1) stmt1;
                stmt2;
   when (exp2) stmt3;
                stmt4;
end;
```

• The LINK statement is used in a program to jump immediately to the label statement\_label and to continue program execution at that point. It is not used to specify a link function in a generalized linear model.

Please consult the individual chapters for other, procedure-specific differences between programming statements and the SAS DATA step and for procedure-specific details, limitations, and rules.

When coding your programming statements, avoid defining variables that begin with an underscore ( ), because they might conflict with internal variables that are created by procedures that support programming statements.

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