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SAS/STAT® 9.22 User's Guide

The LOESS Procedure

(Book Excerpt)



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Chapter 50

The LOESS Procedure

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Overview: LOESS Procedure

The LOESS procedure implements a nonparametric method for estimating regression surfaces pioneered by Cleveland, Devlin, and Grosse (1988), Cleveland and Grosse (1991), and Cleveland, Grosse, and Shyu (1992). The LOESS procedure allows great flexibility because no assumptions about the parametric form of the regression surface are needed.

The SAS System provides many regression procedures such as the GLM, REG, and NLIN procedures for situations in which you can specify a reasonable parametric model for the regression surface. You can use the LOESS procedure for situations in which you do not know a suitable parametric form of the regression surface. Furthermore, the LOESS procedure is suitable when there are outliers in the data and a robust fitting method is necessary.

The main features of the LOESS procedure are as follows:

- fits nonparametric models
- supports the use of multidimensional data
- supports multiple dependent variables
- supports both direct and interpolated fitting that uses kd trees
- performs statistical inference
- performs automatic smoothing parameter selection
- performs iterative reweighting to provide robust fitting when there are outliers in the data
- supports graphical displays produced through ODS Graphics

Local Regression and the Loess Method

Assume that for $i = 1$ to n , the i th measurement y_i of the response y and the corresponding measurement x_i of the vector x of p predictors are related by

$$y_i = g(x_i) + \epsilon_i$$

where g is the regression function and ϵ_i is a random error. The idea of local regression is that at a predictor x , the regression function $g(x)$ can be locally approximated by the value of a function in some specified parametric class. Such a local approximation is obtained by fitting a regression surface to the data points within a chosen neighborhood of the point x .

In the loess method, weighted least squares is used to fit linear or quadratic functions of the predictors at the centers of neighborhoods. The radius of each neighborhood is chosen so that the neighborhood contains a specified percentage of the data points. The fraction of the data, called the *smoothing parameter*, in each local neighborhood controls the smoothness of the estimated surface. Data points

in a given local neighborhood are weighted by a smooth decreasing function of their distance from the center of the neighborhood.

In a direct implementation, such fitting is done at each point at which the regression surface is to be estimated. A much faster computational procedure is to perform such local fitting at a selected sample of points in predictor space and then to blend these local polynomials to obtain a regression surface.

You can use the LOESS procedure to perform statistical inference provided that the error distribution satisfies some basic assumptions. In particular, such analysis is appropriate when the ϵ_i are i.i.d. normal random variables with mean 0. By using the iterative reweighting, the LOESS procedure can also provide statistical inference when the error distribution is symmetric but not necessarily normal. Furthermore, by doing iterative reweighting, you can use the LOESS procedure to perform robust fitting in the presence of outliers in the data.

While all output of the LOESS procedure can be optionally displayed, most often the LOESS procedure is used to produce output data sets that will be viewed and manipulated by other SAS procedures. PROC LOESS uses the Output Delivery System (ODS) to place results in output data sets. This is a departure from older SAS procedures that provide OUTPUT statements to create SAS data sets from analysis results.

Getting Started: LOESS Procedure

Scatter Plot Smoothing

The following data from the Connecticut Tumor Registry presents age-adjusted numbers of melanoma incidences per 100,000 people for the 37 years from 1936 to 1972 (Houghton, Flannery, and Viola 1980).

```
data Melanoma;
  input Year Incidences @@;
  format Year d4.0;
datalines;
1936    0.9   1937    0.8   1938    0.8   1939    1.3
1940    1.4   1941    1.2   1942    1.7   1943    1.8
1944    1.6   1945    1.5   1946    1.5   1947    2.0
1948    2.5   1949    2.7   1950    2.9   1951    2.5
1952    3.1   1953    2.4   1954    2.2   1955    2.9
1956    2.5   1957    2.6   1958    3.2   1959    3.8
1960    4.2   1961    3.9   1962    3.7   1963    3.3
1964    3.7   1965    3.9   1966    4.1   1967    3.8
1968    4.7   1969    4.4   1970    4.8   1971    4.8
1972    4.8
;
```

The following PROC SGPLOT statements produce the simple scatter plot of these data displayed in Figure 50.1.

```
proc sgplot data=Melanoma;
  scatter y=Incidences x=Year;
run;
```

Figure 50.1 Scatter Plot of the Melanoma Data

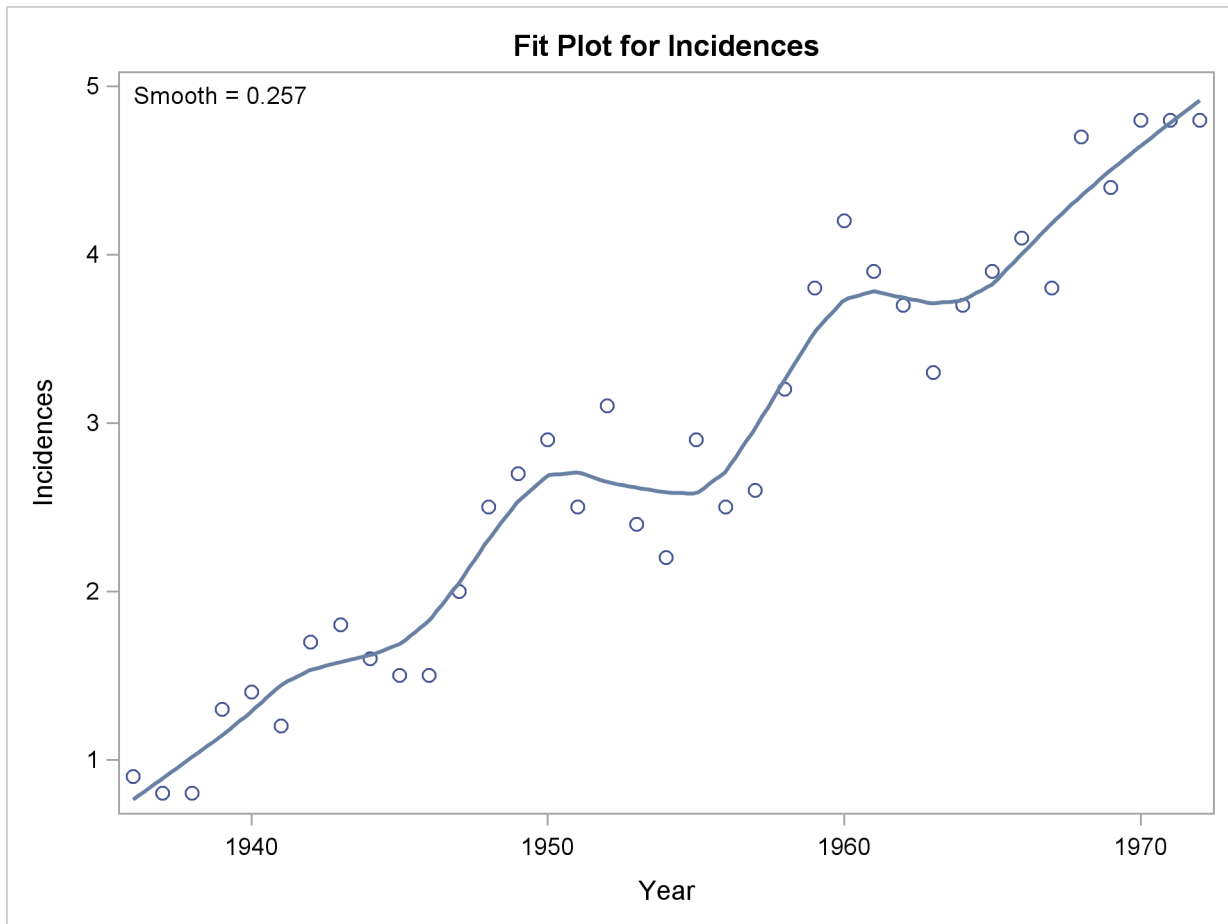


Suppose that you want to smooth the response variable *Incidences* as a function of the variable *Year*. The following PROC LOESS statements request this analysis with the default settings:

```
ods graphics on;

proc loess data=Melanoma;
  model Incidences=Year;
run;
```

You use the PROC LOESS statement to invoke the procedure and specify the data set. The MODEL statement names the dependent and independent variables.

Figure 50.2 Default Loess Fit for the Melanoma data

When you enable graphics with an ODS GRAPHICS statement, PROC LOESS produces several default plots. [Figure 50.2](#) shows the “Fit Plot” that overlays the loess fit on a scatter plot of the data. You can see that the loess fit captures the increasing trend in the data as well as the periodic pattern in the data, which is related to an 11-year sunspot activity cycle.

Figure 50.3 Fit Summary

The LOESS Procedure	
Selected Smoothing Parameter: 0.257	
Dependent Variable: Incidences	
Fit Summary	
Fit Method	kd Tree
Blending	Linear
Number of Observations	37
Number of Fitting Points	37
kd Tree Bucket Size	1
Degree of Local Polynomials	1
Smoothing Parameter	0.25676
Points in Local Neighborhood	9
Residual Sum of Squares	2.03105
Trace[L]	8.62243
GCV	0.00252
AICC	-1.17277

Figure 50.3 shows the “Fit Summary” table. This table details the settings used and provides statistics about the fit that is produced. You can see that smoothing parameter value for this loess fit is 0.257. This smoothing parameter determines the fraction of the data in each local neighborhood. In this example, there are 37 data points and so the smoothing parameter value of 0.257 yields local neighborhoods containing 9 observations.

Figure 50.4 Smoothing Parameter Selection

Optimal Smoothing Criterion	
AICC	Smoothing Parameter
-1.17277	0.25676

The “Smoothing Criterion” table provides information about how this smoothing parameter value is selected. The default method implemented in PROC LOESS chooses the smoothing parameter that minimizes the AICC criterion (Hurvich, Simonoff, and Tsai 1998) that strikes a balance between the residual sum of squares and the complexity of the fit.

You use options in the MODEL statement to change the default settings and request optionally displayed tables. For example, the following statements request that the “Model Summary” and “Output Statistics” tables be included in the displayed output. By default, these tables are not displayed.

```
proc loess data=Melanoma;
  model Incidences=Year / details(ModelSummary OutputStatistics);
run;
```

Figure 50.5 Model Summary Table

The LOESS Procedure				
Dependent Variable: Incidences				
Model Summary				
Smoothing Parameter	Local Points	Residual SS	GCV	AICC
0.41892	15	3.42229	0.00339	-0.96252
0.68919	25	4.05838	0.00359	-0.93459
0.31081	11	2.51054	0.00279	-1.12034
0.20270	7	1.58513	0.00239	-1.12221
0.17568	6	1.56896	0.00241	-1.09706
0.28378	10	2.50487	0.00282	-1.10402
0.20270	7	1.58513	0.00239	-1.12221
0.25676	9	2.03105	0.00252	-1.17277
0.22973	8	2.02965	0.00256	-1.15145
0.25676	9	2.03105	0.00252	-1.17277

The “Model Summary” table shown in [Figure 50.5](#) provides information about all the models that PROC LOESS evaluated in choosing the smoothing parameter value.

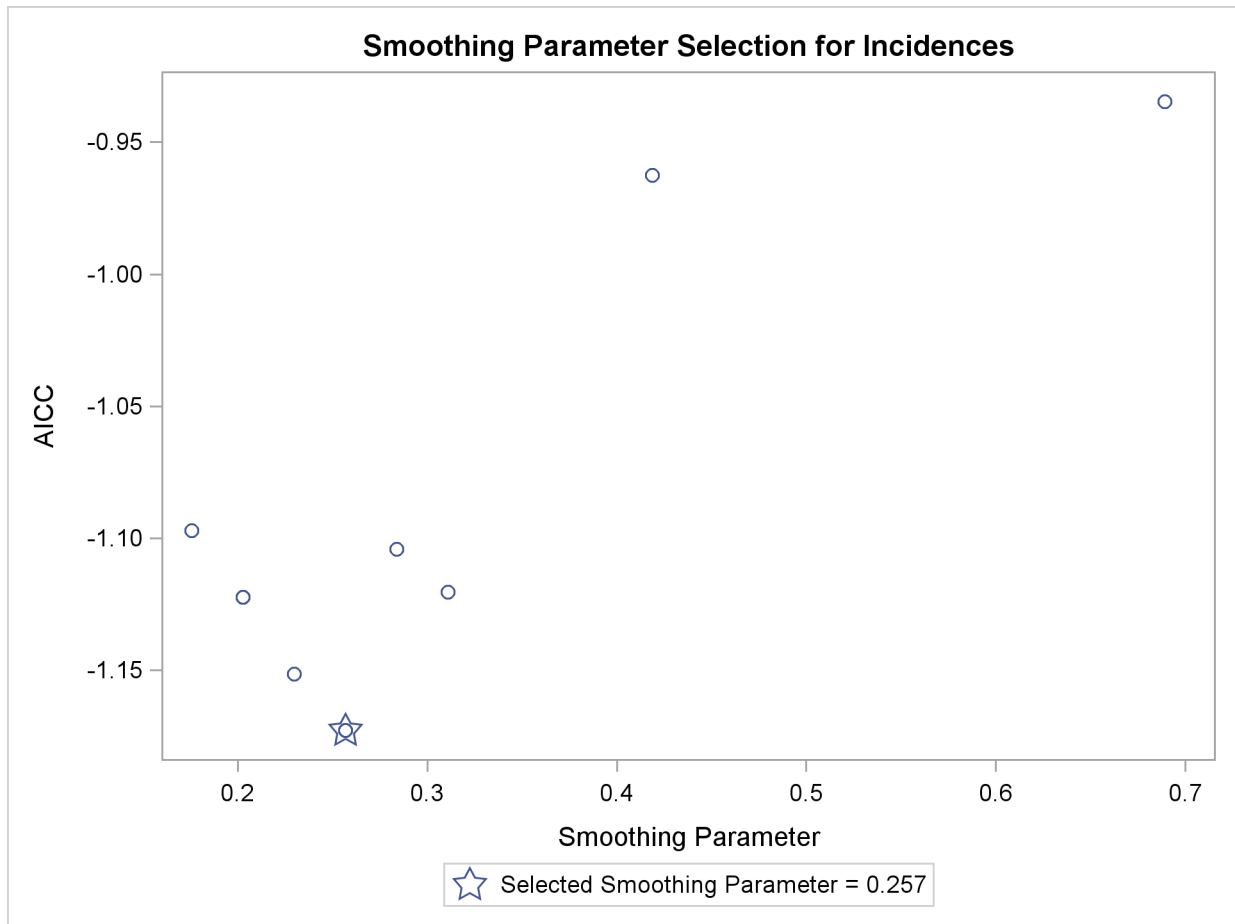
Figure 50.6 AICC Criterion by Smoothing Parameter

Figure 50.6 shows the “Criterion Plot” that provides a graphical display of the smoothing parameter selection process.

Figure 50.7 Output Statistics

The LOESS Procedure				
Selected Smoothing Parameter: 0.257				
Dependent Variable: Incidences				
Output Statistics				
Obs	Year	Incidences	Predicted Incidences	Residual
1	1936	0.90000	0.76235	0.13765
2	1937	0.80000	0.88992	-0.08992
3	1938	0.80000	1.01764	-0.21764
4	1939	1.30000	1.14303	0.15697
5	1940	1.40000	1.28654	0.11346
6	1941	1.20000	1.44528	-0.24528
7	1942	1.70000	1.53482	0.16518
8	1943	1.80000	1.57895	0.22105
9	1944	1.60000	1.62058	-0.02058
10	1945	1.50000	1.68627	-0.18627
11	1946	1.50000	1.82449	-0.32449
12	1947	2.00000	2.04976	-0.04976
13	1948	2.50000	2.30981	0.19019
14	1949	2.70000	2.53653	0.16347
15	1950	2.90000	2.68921	0.21079
16	1951	2.50000	2.70779	-0.20779
17	1952	3.10000	2.64837	0.45163
18	1953	2.40000	2.61468	-0.21468
19	1954	2.20000	2.58792	-0.38792
20	1955	2.90000	2.57877	0.32123
21	1956	2.50000	2.71078	-0.21078
22	1957	2.60000	2.96981	-0.36981
23	1958	3.20000	3.26005	-0.06005
24	1959	3.80000	3.54143	0.25857
25	1960	4.20000	3.73482	0.46518
26	1961	3.90000	3.78186	0.11814
27	1962	3.70000	3.74362	-0.04362
28	1963	3.30000	3.70904	-0.40904
29	1964	3.70000	3.72917	-0.02917
30	1965	3.90000	3.82382	0.07618
31	1966	4.10000	4.00515	0.09485
32	1967	3.80000	4.18573	-0.38573
33	1968	4.70000	4.35152	0.34848
34	1969	4.40000	4.50284	-0.10284
35	1970	4.80000	4.64413	0.15587
36	1971	4.80000	4.78291	0.01709
37	1972	4.80000	4.91602	-0.11602

Figure 50.7 show the “Output Statistics” table that contains the predicted loess fit value at each observation in the input data set.

Although the default method for selecting the smoothing parameter value is often satisfactory, it is often a good practice to examine how the loess fit varies with the smoothing parameter. In some cases, fits with different smoothing parameters might reveal important features of the data that cannot be discerned by looking at a fit with just a single “best” smoothing parameter. [Example 50.4](#) provides

such an example. You can produce the loess fits for a range of smoothing parameters by using the **SMOOTH=** option in the **MODEL** statement as follows:

```
proc loess data=Melanoma;
  model Incidences=Year/smooth=0.1 0.25 0.4 0.6 residual;
  ods output OutputStatistics=Results;
run;
```

The **RESIDUAL** option causes the residuals to be added to the “Output Statistics” table. Note that, even if you do not specify the **DETAILS** option in the **MODEL** statement to request the display of the “Output Statistics” table, you can use an **ODS OUTPUT** statement to output this and other optionally displayed tables as data sets.

PROC PRINT displays the first five observations of the **Results** data set:

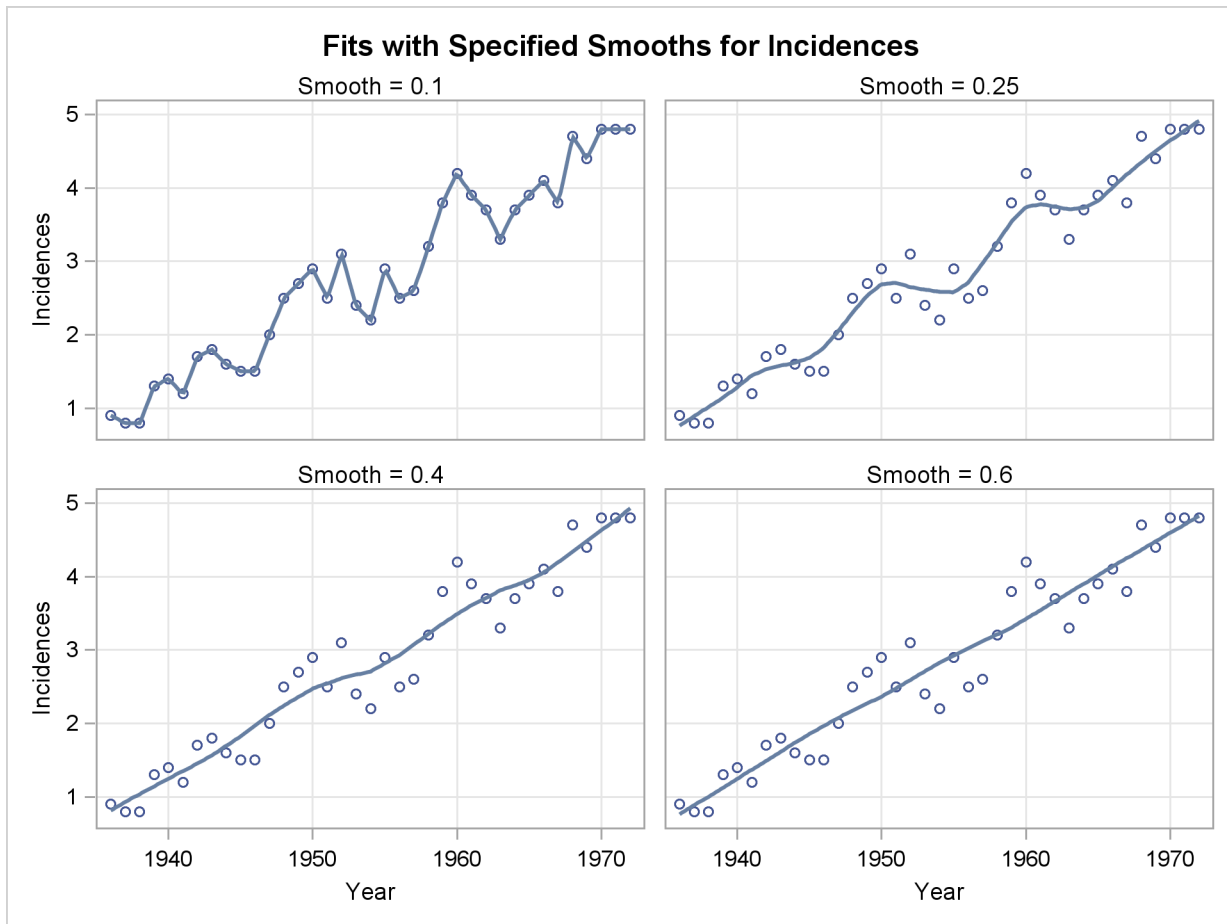
```
proc print data=Results(obs=5);
  id obs;
run;
```

Figure 50.8 PROC PRINT Output of the Results Data Set

Obs	Smoothing Parameter	Year	Dep Var	Pred	Residual
1	0.1	1936	0.9	0.90000	0
2	0.1	1937	0.8	0.80000	0
3	0.1	1938	0.8	0.80000	0
4	0.1	1939	1.3	1.30000	0
5	0.1	1940	1.4	1.40000	0

Note that the fits for all the smoothing parameters are placed in single data set. A variable named **SmoothingParameter** that you use to distinguish each fit is included in this data set.

When you specify a list of smoothing parameters for a model and you have enabled **ODS Graphics**, **PROC LOESS** produces a panel containing up to six plots that show the fit obtained for each value of the smoothing parameter that you specify. If you specify more than six smoothing values, then multiple panels are produced. For each regressor, **PROC LOESS** also produces panels of the residuals versus each regressor by the smoothing parameters that you specify.

Figure 50.9 Loess Fits for a Range of Smoothing Parameters

If you examine the plots in [Figure 50.9](#), you see that a visually reasonable fit is obtained with smoothing parameter values of 0.25. With smoothing parameter value 0.1, there is gross overfitting in the sense that the original data are exactly interpolated. When the smoothing parameter value is 0.4, you obtain an overly smooth fit where the contribution of the sunspot cycle has been mostly averaged away. At smoothing parameter value 0.6 the fit shows just the increasing trend in the data.

It is also instructive to look at scatter plots of the residuals for each of the fits. These are also produced by default by PROC LOESS when ODS Graphics is enabled.

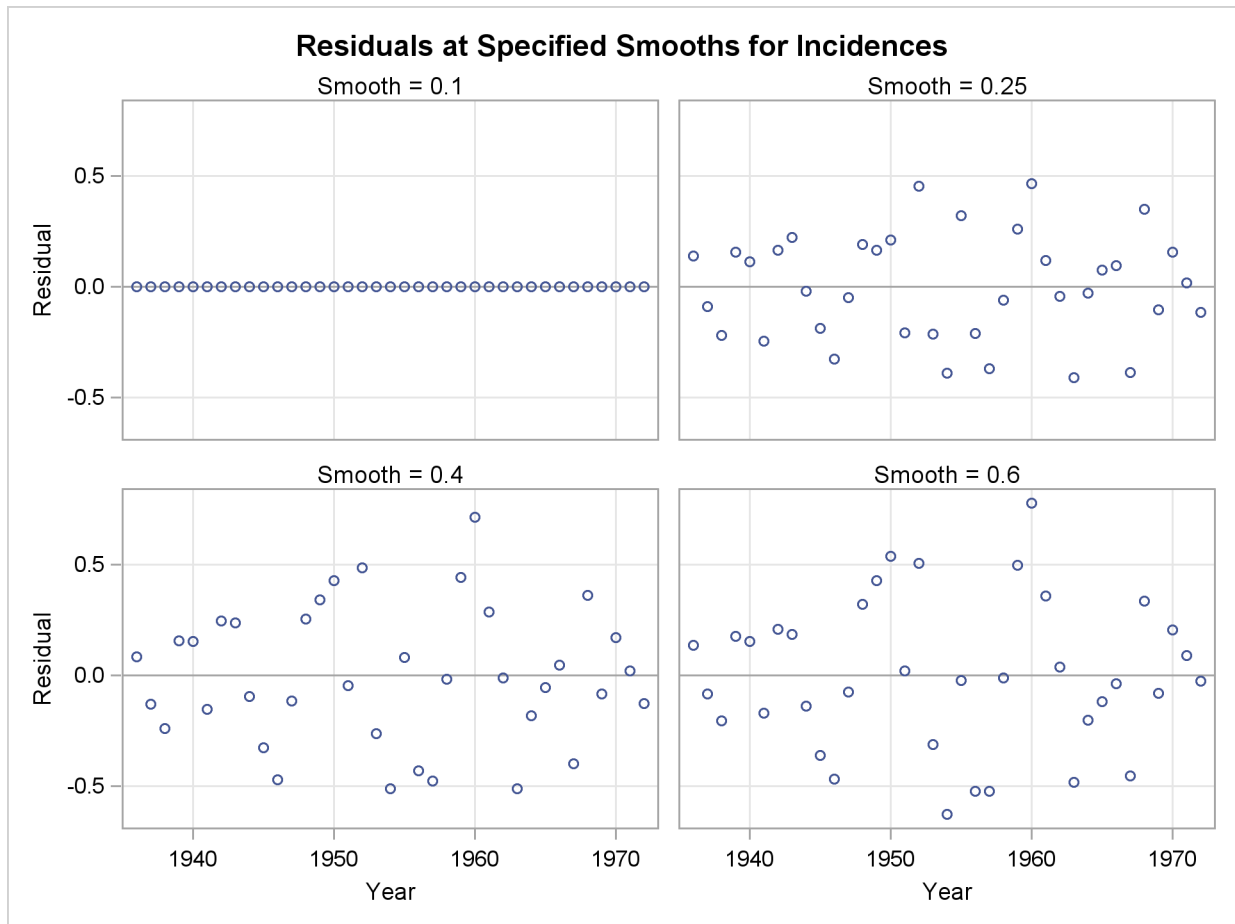
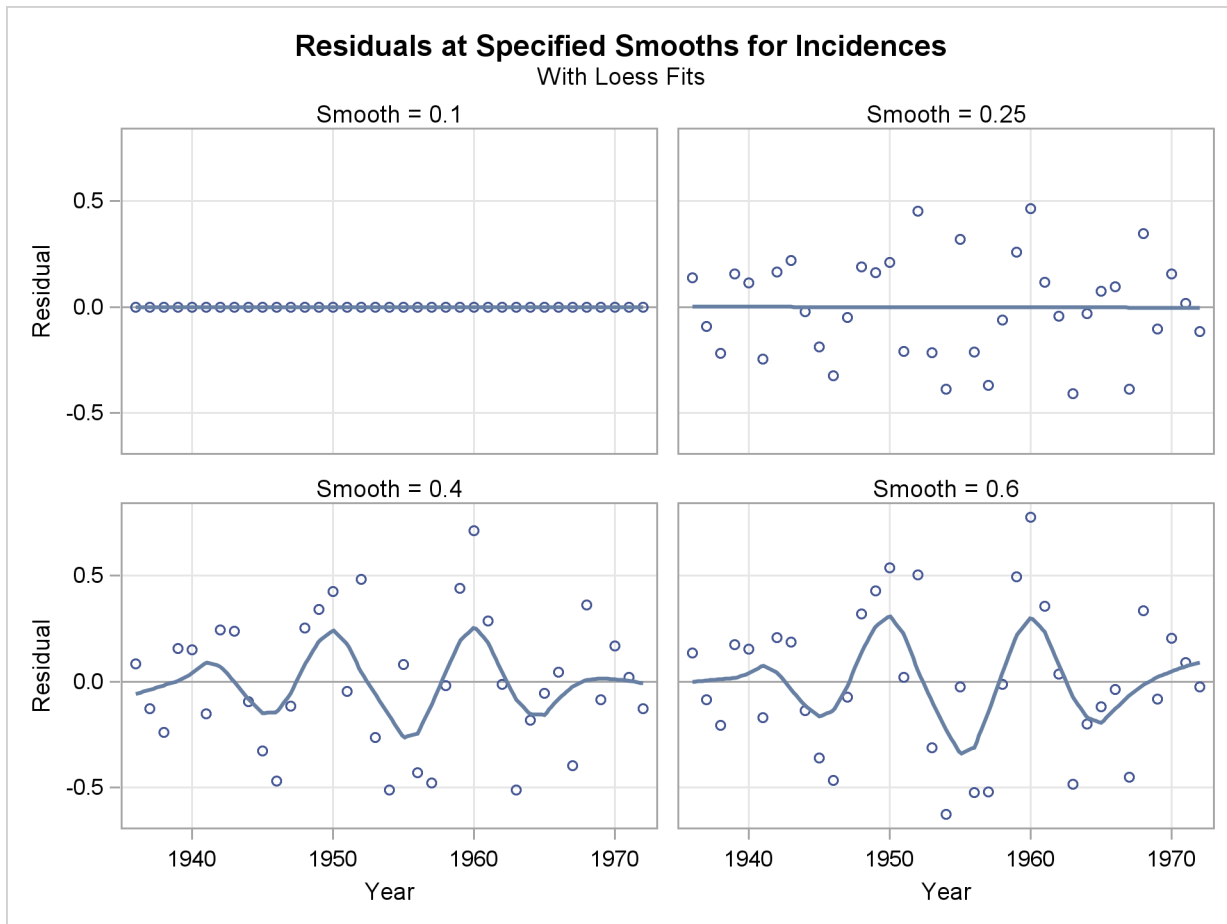
Figure 50.10 Residuals of Loess Fits for a Range of Smoothing Parameters

Figure 50.10 shows a scatter plot of the residuals by year for each smoothing parameter value. One way to discern patterns in these residuals is to superimpose a loess fit on each plot in the panel. You request loess fits on the residual plots in this panel by specifying the `SMOOTH=` suboption of the `PLOTS=RESIDUALSBYSMOOTH` option in the `PROC LOESS` statement. Note that the loess fits that are displayed on each of the residual plots are obtained independently of the loess fit that produces these residuals. The following statements show how you do this for the Melanoma data.

```
proc loess data=Melanoma plots=ResidualsBySmooth(smooth);
  model Incidences=Year/smooth=0.1 0.25 0.4 0.6;
run;
```

Figure 50.11 Residuals with Superimposed Loess Fits

The loess fits shown on the plots in [Figure 50.11](#) help confirm the conclusions obtained when you look at [Figure 50.9](#). Note that residuals for smoothing parameter value 0.25 do not exhibit any pattern, confirming that at this value the loess fit of the melanoma data has successfully modeled the variation in this data. By contrast, the residuals for the fit with smoothing parameter 0.6 retain the variation caused by the sunspot cycle.

The examination of the fits and residuals obtained with a range of smoothing parameter values confirms that the value of 0.257 that PROC LOESS selects automatically is appropriate for these data. The next step in this analysis is to examine fit diagnostics and produce confidence limit for the fit. If you enable ODS Graphics, then a panel of fit diagnostics is produced. Furthermore, you can request prediction confidence limits by adding the CLM option in the [MODEL](#) statement. By default 95% limits are produced, but you can use the ALPHA= option in the [MODEL](#) statement to change the significance level. The following statements request 90% confidence limits.

```
proc loess data=Melanoma;
  model Incidences=Year/clm alpha=0.1;
run;

ods graphics off;
```

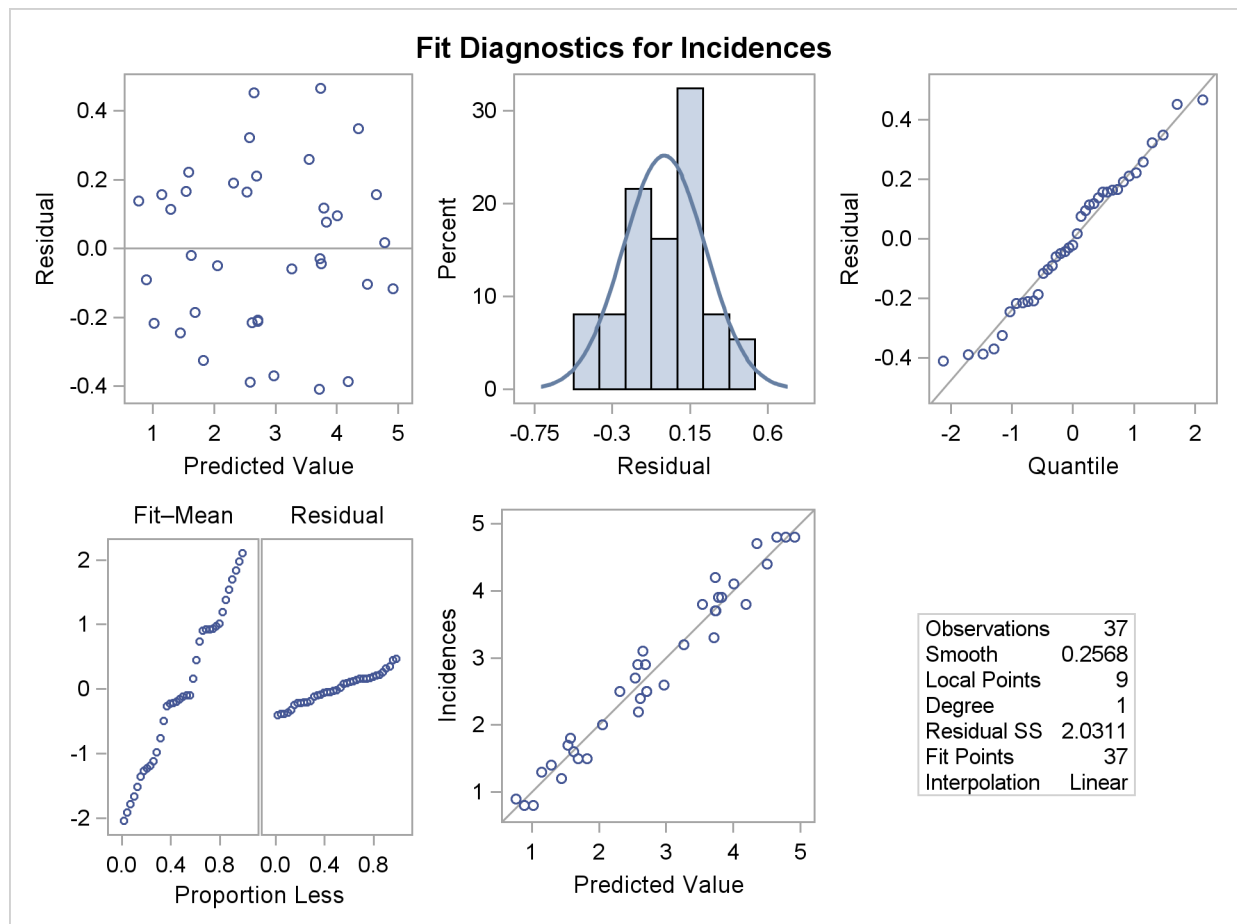
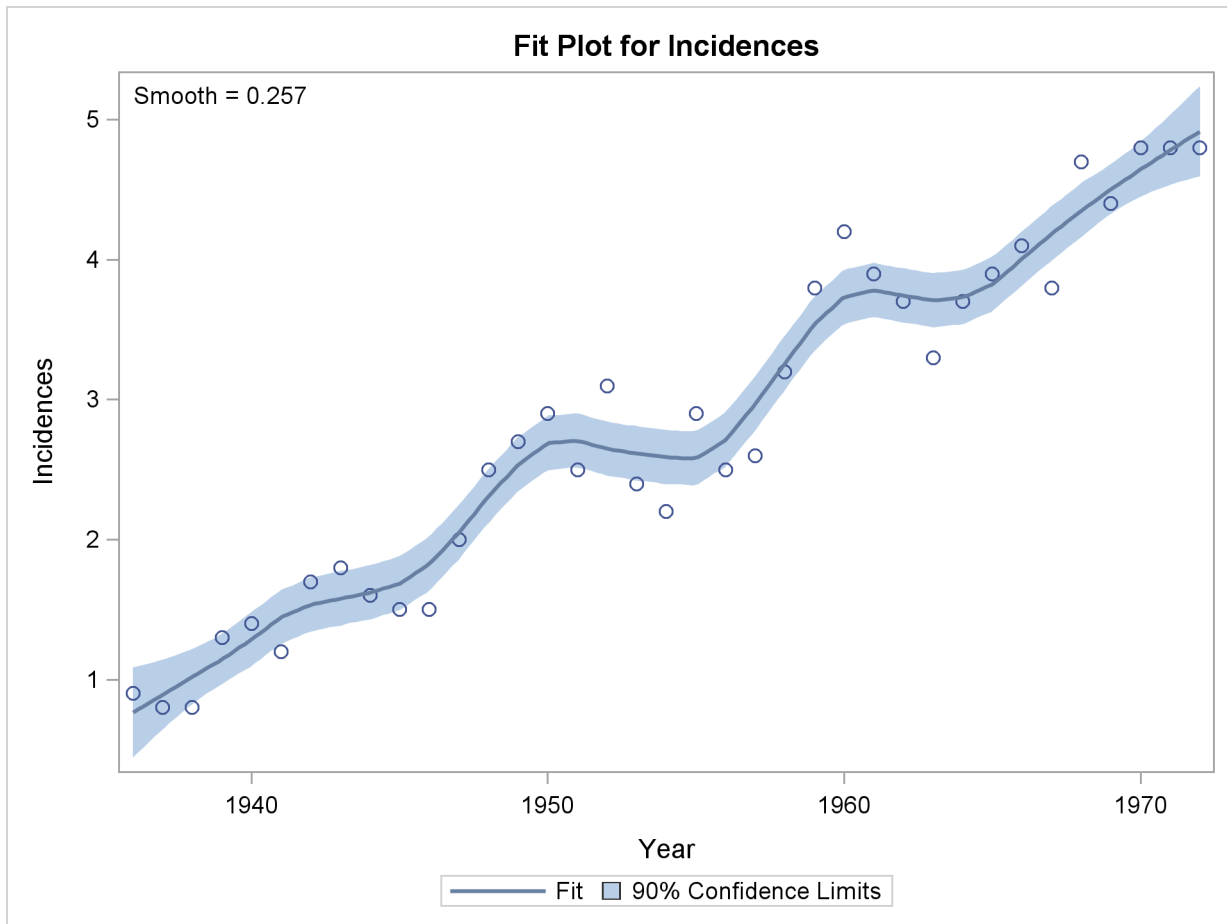
Figure 50.12 Fit Diagnostics

Figure 50.12 shows the fit diagnostics panel. The histogram of the residuals with overlaid normal density estimator and the normal quantile plot show that the residuals do exhibit some small departure from normality. The “Residual-Fit” spread plot shows that the spread in the centered fit is much wider than the spread in the residuals. This indicates that the fit has accounted for most of the variation in the incidences of melanoma in this data. This conclusion is supported by the absence of any clear pattern in the scatter plot of residuals by predicted values and the closeness of the points to the 45-degree reference line in the plot of observed by predicted values.

Figure 50.13 Loess Fit of Melanoma Data with 90% Confidence Limits

Finally, Figure 50.13 shows the selected loess fit with 90% confidence limits.

Syntax: LOESS Procedure

The following statements are available in PROC LOESS:

```
PROC LOESS < DATA=SAS-data-set > ;
  MODEL dependents=regressors < / options > ;
  ID variables ;
  BY variables ;
  WEIGHT variable ;
  SCORE DATA=SAS-data-set < ID=(variable list) > < / options > ;
```

The PROC LOESS and MODEL statements are required. The BY, WEIGHT, and ID statements are optional. The SCORE statement is optional, and more than one SCORE statement can be used.

The statements used with the LOESS procedure, in addition to the PROC LOESS statement, are as follows.

BY	specifies variables to define subgroups for the analysis.
ID	names variables to identify observations in the displayed output.
MODEL	specifies the dependent and independent variables in the loess model, details and parameters for the computational algorithm, and the required output.
SCORE	specifies a data set containing observations to be scored.
WEIGHT	declares a variable to weight observations.

PROC LOESS Statement

PROC LOESS *< options >* ;

The PROC LOESS statement is required. You can specify the following options in the PROC LOESS statement:

DATA=SAS-data-set

names the SAS data set to be used by PROC LOESS. If the DATA= option is not specified, PROC LOESS uses the most recently created SAS data set.

PLOTS *< (global-plot-options) > < = plot-request < (options) > >*

PLOTS *< (global-plot-options) > < = (plot-request < (options) > < ... plot-request < (options) > > >*

controls the plots produced through ODS Graphics. When you specify only one plot request, you can omit the parentheses around the plot request. Here are some examples:

```
plots=none
plots=residuals(smooth)
plots(unpack)=diagnostics
plots(only)=(fit residualHistogram)
```

You must enable ODS Graphics before requesting plots as shown in the following example. For general information about ODS Graphics, see Chapter 21, “[Statistical Graphics Using ODS](#).”

```
ods graphics on;

proc loess;
  model y = x;
run;

proc loess plots=all;
  model y = x;
run;

ods graphics off;
```

The first PROC LOESS step does not specify the PLOTS= option, so the default plots comprising a panel of fit diagnostics, a fit plot, and a plot of residuals by x are produced. The PLOTS=ALL option in the second PROC LOESS step produces the default plots together with individual plots of the plots in the panel of fit diagnostics.

If you have enabled ODS Graphics but do not specify the PLOTS= option, then PROC LOESS produces a default set of plots. The following table lists the default set of plots produced.

Table 50.1 Default ODS Graphics Produced

Plot	Conditional On
ContourFitPanel	SMOOTH= option specified in the MODEL statement
ContourFit	Model with two regressors
CriterionPlot	Smoothing parameter selection performed
DiagnosticsPanel	Unconditional
ResidualsBySmooth	SMOOTH= option specified in the MODEL statement
ResidualPanel	Unconditional
FitPanel	SMOOTH= option specified in the MODEL statement
FitPlot	Model with one regressor
ScorePlot	One or more SCORE statements and a model with one regressor

For models with multiple dependent variables, separate plots are produced for each dependent variable. For models where multiple smoothing parameters are requested with the SMOOTH= option in the [MODEL](#) statement and smoothing parameter value selection is not requested, separate plots are produced for each smoothing parameter. If smoothing parameter value selection is requested with the SELECT= option in the [MODEL](#) statement, then the plots are produced for the selected model only. However, if you specify the STEPS suboption of the SELECT= option, then plots are produced for all smoothing parameters examined in the selection process.

The *global-plot-options* apply to all relevant plots generated by the LOESS procedure, unless they are overridden with a *specific-plot-option*. The *global-plot-options* supported by the LOESS procedure follow.

Global Plot Options

MAXPOINTS=NONE | *number*

specifies that plots with elements that require processing more than *number* points are suppressed. The default is MAXPOINTS=5000. This cutoff is ignored if you specify MAXPOINTS=NONE.

ONLY

suppresses the default plots. Only the plots specifically requested are produced.

UNPACK

suppresses paneling. By default, multiple plots can appear in some output panels. Specify UNPACK to get each plot individually. You can specify PLOTS(UNPACK) to unpack the default plots. You can also specify UNPACK as a suboption with CONTOURFITPANEL, DIAGNOSTICS, FITPANEL, RESIDUALS and RESIDUALSBYSMOOTH.

Specific Plot Options

The following listing describes the specific plots and their options.

ALL

requests that all plots appropriate for the particular analysis be produced. You can specify other options with ALL; for example, to request all plots and unpack only the residuals, specify PLOTS=(ALL RESIDUALS(UNPACK)).

CONTOURFIT <(contour-options)>

produces a contour plot of the fitted surface overlaid with a scatter plot of the data for models with two regressors. Contour plots are not produced if you specify the DIRECT option in the **MODEL** statement. You can use the following *contour-options* to control how the observations are displayed:

OBS=GRADIENT

specifies that observations be displayed as circles colored by the observed response. The same color gradient is used to display the fitted surface and the observations. Observations where the predicted response is close to the observed response have similar colors—the greater the contrast between the color of an observation and the surface, the larger the residual is at that point. OBS=GRADIENT is the default if you do not specify any *contour-options*.

OBS=NONE

suppresses the observations.

OBS=OUTLINE

specifies that observations be displayed as circles with a border but with a completely transparent fill.

OBS=OUTLINEGRADIENT

is the same as OBS=GRADIENT except that a border is shown around each observation. This option is useful to identify the location of observations where the residuals are small, because at these points the color of the observations and the color of the surface are indistinguishable.

CONTOURFITPANEL <(< UNPACK > < contour-options >)>

produces panels of contour plots overlaid with a scatter plot of the data for each smoothing parameter specified in the SMOOTH= option in the **MODEL** statement, for models with two regressors. This plot is not produced if you specify the DIRECT option in the **MODEL** statement. If you do not specify the SMOOTH= option or if the model does not have two regressors, then this plot is not produced. If you specify the SELECT=

option in addition to the SMOOTH= option in the **MODEL** statement, then you need to additionally specify the STEPS suboption of the SELECT= option to obtain this plot. Note that each panel contains at most six plots, and multiple panels are used in the case that there are more than six smoothing parameters in the SMOOTH= option in the **MODEL** statement. See the CONTOURFIT option for a description of the individual plots in this panel. The UNPACK option suppresses paneling, and the *contour-options* are the same as for the CONTOURFIT option.

CRITERIONPLOT | CRITERION

displays a scatter plot of the value of the SELECTION= criterion versus the smoothing parameter value for all smoothing parameter values examined in the selection process. This plot is not produced if smoothing parameter selection is not done.

DIAGNOSTICSPANEL | DIAGNOSTICS <(UNPACK)>

produces a summary panel of fit diagnostics consisting of the following:

- residuals versus the predicted values
- histogram of the residuals
- normal quantile plot of the residuals
- a “Residual-Fit” (or RF) plot consisting of side-by-side quantile plots of the centered fit and the residuals.
- dependent variable values versus the predicted values

You can request the five plots in this panel as individual plots by specifying the UNPACK option. You can also request individual plots in the panel by name without having to unpack the panel. Note that the fit diagnostics panel is produced by default whenever ODS Graphics is enabled.

FITPANEL <(UNPACK)>

produces panels of plots showing the fitted LOESS curve overlaid on a scatter plot of the input data for each smoothing parameter specified in the SMOOTH= option in the **MODEL** statement. If you do not specify the SMOOTH= option or the model has more than one regressor, then this plot is not produced. If you specify the SELECT= option in addition to the SMOOTH= option in the **MODEL** statement, then you need to additionally specify the STEPS suboption of the SELECT= option to obtain this plot. Note that each panel contains at most six plots, and multiple panels are used in the case that there are more than six smoothing parameters in the SMOOTH= option in the **MODEL** statement. If the CLM option is specified in the **MODEL** statement, then a confidence band at the significance level specified in the ALPHA= option is included in each plot in the panels. If you specify the UNPACK option, then all fit panels are unpacked.

FITPLOT | FIT

produces a scatter plot of the input data with the fitted LOESS curve overlaid for models with a single regressor. If the CLM option is specified in the **MODEL** statement, then a confidence band at the significance level specified in the ALPHA= option is included in the plot.

NONE

suppresses all plots.

OBSERVEDBYPREDICTED

produces a scatter plot of the dependent variable values by the predicted values.

QQPLOT | QQ

produces a normal quantile plot of the residuals.

RESIDUALSBYSMOOTH <(< UNPACK > < SMOOTH >)>

produces for each regressor panels of plots showing the residuals of the LOESS fit versus the regressor for each smoothing parameter specified in the SMOOTH= option in the **MODEL** statement. If you do not specify the SMOOTH= option, then this plot is not produced. If you specify the SELECT= option in addition to the SMOOTH= option in the **MODEL** statement, then you need to additionally specify the STEPS suboption of the SELECT= option to obtain this plot. Note that each panel contains at most six plots, and multiple panels are used in the case that there are more than six smoothing parameters in the SMOOTH= option in the **MODEL** statement. If you specify the UNPACK option, then all RESIDUALSBYSMOOTH panels are unpacked.

The SMOOTH option requests that a nonparametric fit line be shown in each plot in the panel. The type of nonparametric fit and the options used are controlled by the template that underlies this plot. In the standard template that is provided, the nonparametric smooth is specified to be a loess fit corresponding to the default options of PROC LOESS, except that the PRESEARCH suboption is always used. It is important to note that the loess fit that is shown in each of the residual plots is computed independently of the loess fit that is used to obtain the residuals.

RESIDUALBYPREDICTED

produces a scatter plot of the residuals by the predicted values.

RESIDUALHISTOGRAM

produces a histogram of the residuals.

RESIDUALPANEL | RESIDUALS <(residual-options) >

produces panels of the residuals versus the regressors in the model. Note that each panel contains at most six plots, and multiple panels are used when there are more than six regressors in the model.

The following *residual-options* are available:

SMOOTH

requests that a nonparametric fit line be shown in each plot in the panel. The type of nonparametric fit and the options used are controlled by the template that underlies this plot. In the standard template that is provided, the nonparametric smooth is specified to be a loess fit corresponding to the default options of PROC LOESS, except that the PRESEARCH suboption is always used. It is important to note that the loess fit that is shown in each of the residual plots is computed independently of the loess fit that is used to obtain the residuals.

UNPACK

suppresses paneling.

RFPLOT | RF

produces a “Residual-Fit” (or RF) plot consisting of side-by-side quantile plots of the centered fit and the residuals. This plot “shows how much variation in the data is explained by the fit and how much remains in the residuals” (Cleveland 1993).

SCOREPLOT | SCORE

produces a scatter plot of the scored values at the score points for each **SCORE** statement. SCORE plots are not produced for models with more than one regressor. If the CLM option is specified in the **MODEL** statement, then confidence bars at the significance level specified in the ALPHA= option are shown at score data points.

BY Statement

BY *variables* ;

You can specify a BY statement with PROC LOESS to obtain separate analyses on observations in groups that are defined by the BY variables. When a BY statement appears, the procedure expects the input data set to be sorted in order of the BY variables. If you specify more than one BY statement, only the last one specified is used.

If your input data set is not sorted in ascending order, use one of the following alternatives:

- Sort the data by using the SORT procedure with a similar BY statement.
- Specify the NOTSORTED or DESCENDING option in the BY statement for the LOESS procedure. The NOTSORTED option does not mean that the data are unsorted but rather that the data are arranged in groups (according to values of the BY variables) and that these groups are not necessarily in alphabetical or increasing numeric order.
- Create an index on the BY variables by using the DATASETS procedure (in Base SAS software).

For more information about BY-group processing, see the discussion in *SAS Language Reference: Concepts*. For more information about the DATASETS procedure, see the discussion in the *Base SAS Procedures Guide*.

ID Statement

ID *variables* ;

The ID statement is optional, and more than one ID statement can be used. The variables listed in any of the ID statements are displayed in the “Output Statistics” table beside each observation. Any

variables specified as a regressor or dependent variable in the **MODEL** statement already appear in the “Output Statistics” table and are not treated as ID variables, even if they appear in the variable list of an ID statement.

MODEL Statement

The **MODEL** statement names the dependent variables and the independent variables. Variables specified in the **MODEL** statement must be numeric variables in the data set being analyzed.

Table 50.2 lists the options available in the **MODEL** statement.

Table 50.2 Summary of MODEL Statement Options

Option	Description
Fit Options	
BUCKET=	specifies the number of points in kd tree buckets
DEGREE=	specifies the degree of local polynomials (1 or 2)
DFMETHOD=	specifies the method of computing lookup degrees of freedom
DIRECT	specifies direct fitting at every data point
DROPSQUARE=	specifies the variables whose squares are to be dropped from local quadratic polynomials
INTERP=	specifies the interpolating polynomials (linear or cubic)
ITERATIONS=	specifies the number of reweighting iterations
SCALE=	specifies the method used to scale the regressor variables
SELECT=	specifies that automatic smoothing parameter selection be done
SMOOTH=	specifies the list of smoothing values
Output Statistics Table Options	
ALL	requests CLM , RESIDUAL , SCALEDINDEP , STD , and T options
CLM	displays confidence limits for mean predictions
RESIDUAL	displays residuals
SCALEDINDEP	displays scaled independent variable coordinates
STD	displays standard errors of the mean predicted values
T	displays <i>t</i> statistics
Other options	
ALPHA=	sets significance level for confidence intervals
DETAILS=	specifies which tables are to be displayed
TRACEL	displays the trace of the smoothing matrix

The following options are available in the **MODEL** statement after a slash (/).

ALL

requests all these options: **CLM**, **RESIDUAL**, **SCALEDINDEP**, **STD**, and **T**.

ALPHA=number

sets the significance level used for the construction of confidence intervals for the current MODEL statement. The value must be between 0 and 1; the default value of 0.05 results in 95% intervals.

BUCKET=number

specifies the maximum number of points in the leaf nodes of the kd tree. The default value used is $s * n / 5$, where s is a smoothing parameter value specified using the SMOOTH= option and n is the number of observations being used in the current BY group. The BUCKET= option is ignored if the DIRECT option is specified.

CLM

requests that $100(1 - \alpha)\%$ confidence limits on the mean predicted value be added to the “Output Statistics” table. By default, 95% limits are computed; the ALPHA= option in the MODEL statement can be used to change the significance level. The use of this option implicitly selects the model option DFMETHOD=EXACT if the DFMETHOD= option has not been explicitly used.

DEGREE=1 | 2

sets the degree of the local polynomials to use for each local regression. The valid values are 1 for local linear fitting and 2 for local quadratic fitting, with 1 being the default.

DETAILS < (tables) >

selects which tables to display, where *tables* is one or more of the specifications KDTREE, MODELSUMMARY, OUTPUTSTATISTICS, and PREDATVERTICES:

- KDTREE displays the kd tree structure.
- MODELSUMMARY displays the fit criteria for all smoothing parameter values that are specified in the SMOOTH= option in the MODEL statement, or that are fit with automatic smoothing parameter selection.
- OUTPUTSTATISTICS displays the predicted values and other requested statistics at the points in the input data set.
- PREDATVERTICES displays fitted values and coordinates of the kd tree vertices where the local least squares fitting is done.

The KDTREE and PREDATVERTICES specifications are ignored if the DIRECT option is specified in the MODEL statement. Specifying the option DETAILS with no qualifying list outputs all tables.

DFMETHOD=NONE | EXACT | APPROX < (approx-options) >

specifies the method used to calculate the lookup degrees of freedom used in performing statistical inference. The default is DFMETHOD=NONE, unless you specify any of the MODEL statement options ALL, CLM, STD, and T, or any SCORE statement CLM option, in which case the default is DFMETHOD=EXACT.

You can specify the following *approx-options* in parentheses after the DFMETHOD=APPROX option:

QUANTILE=number

specifies that the smallest 100(*number*)% of the nonzero coefficients in the smoothing matrix be set to zero in computing the approximate lookup degrees of freedom. The default value is QUANTILE=0.9.

CUTOFF=number

specifies that coefficients in the smoothing matrix whose magnitude is less than the specified value be set to zero in computing the approximate lookup degrees of freedom. Using the CUTOFF= option overrides the QUANTILE= option.

See the section “[Sparse and Approximate Degrees of Freedom Computation](#)” on page 3830 for a description of the method used when the DFMETHOD=APPROX option is specified.

DIRECT

specifies that local least squares fits are to be done at every point in the input data set. When the direct option is not specified, a computationally faster method is used. This faster method performs local fitting at vertices of a kd tree decomposition of the predictor space followed by blending of the local polynomials to obtain a regression surface.

DROPSQUARE=(variables)

specifies the quadratic monomials to exclude from the local quadratic fits. This option is ignored unless the DEGREE=2 option has been specified.

For example,

```
model z=x y / degree=2 dropsquare=(y)
```

uses the monomials 1, x, y, x^2 , and xy in performing the local fitting.

INTERP=LINEAR | CUBIC

specifies the degree of the interpolating polynomials used for blending local polynomial fits at the kd tree vertices. This option is ignored if the DIRECT option is specified in the model statement. INTERP=CUBIC is not supported for models with more than two regressors. The default is INTERP=LINEAR.

ITERATIONS=number

specifies the total number of iterations to be done. The first iteration performs an initial LOESS fit. Subsequent iterations perform iterative reweighting. Such iterations are appropriate when there are outliers in the data or when the error distribution is a symmetric long-tailed distribution. The default number of iterations is 1.

RESIDUAL | R

specifies that residuals be included in the “Output Statistics” table.

SCALE=NONE | SD < (number) >

specifies the scaling method to be applied to scale the regressors. The default is NONE, in which case no scaling is applied. A specification of SD(*number*) indicates that a trimmed standard deviation is to be used as a measure of scale, where *number* is the trimming fraction. A specification of SD with no qualification defaults to 10% trimmed standard deviation.

SCALEDINDEP

specifies that scaled regressor coordinates be included in the output tables. This option is ignored if the SCALE= model option is not used or if SCALE=NONE is specified.

SELECT=criterion < (< **GLOBAL** > < **PRESEARCH** > < **STEPS** > < **RANGE**(lower,upper) >) >

SELECT=DFCriterion < (target < **GLOBAL** > < **PRESEARCH** > < **STEPS** > < **RANGE**(lower,upper) >) >

specifies that automatic smoothing parameter selection be done using the named *criterion* or *DFCriterion*. Valid values for the *criterion* are as follows:

AICC specifies the AIC_C criterion (Hurvich, Simonoff, and Tsai 1998).

AICC1 specifies the AIC_{C_1} criterion (Hurvich, Simonoff, and Tsai 1998).

GCV specifies the generalized cross validation criterion (Craven and Wahba 1979).

The *DFCriterion* specifies the measure used to estimate the model degrees of freedom. The measures implemented in PROC LOESS all depend on prediction matrix L relating the observed and predicted values of the dependent variable. Valid values for the *DFCriterion* are as follows:

DF1 specifies $\text{Trace}(L)$.

DF2 specifies $\text{Trace}(L^T L)$.

DF3 specifies $2\text{Trace}(L) - \text{Trace}(L^T L)$.

For both types of selection, the smoothing parameter value is selected to yield a minimum of an optimization criterion. If you specify *criterion* as one of AICC, AICC1, or GCV, the optimization criterion is the specified *criterion*. If you specify *DFCriterion* as one of DF1, DF2, or DF3, the optimization criterion is $|DFCriterion - target|$, where *target* is a specified target degree of freedom value. Note that if you specify a *DFCriterion*, then you must also specify a target value. See the section “[Automatic Smoothing Parameter Selection](#)” on page 3827 for definitions and properties of the selection criteria.

The selection is done as follows:

- If you specify the SMOOTH=*value-list* option, then PROC LOESS selects the largest value in this list that yields the global minimum of the specified optimization criterion.
- If you do not specify the SMOOTH= option, then PROC LOESS finds a local minimum of the specified optimization criterion by using a golden section search of values less than or equal to one.

You can specify the following suboptions in parentheses after the specified criterion to alter the behavior of the SELECT= option:

GLOBAL

specifies that a global minimum be found within the range of smoothing parameter values examined. This suboption has no effect if you also specify the SMOOTH= option in the MODEL statement.

PRESEARCH

requests an initial grid search to find a smoothing parameter range within which the subsequent golden section search is done. The initial point in this grid is the smoothing parameter value corresponding to the smallest number of points, n , in the local neighborhoods that yields a fit that does not interpolate all the data points. Subsequent fits with number of local points $n + 1$, $n + 2$, $n + 4$, $n + 8$, ... are evaluated until either the number of local points exceeds the number of fitting points or the **SELECT=criterion** starts increasing. This suboption is ignored if you additionally specify the **GLOBAL** suboption of the **SELECT=** option or if you specify the **SMOOTH=** option in the **MODEL** statement. If you additionally specify the **RANGE=** suboption, then the golden section search is done on the intersection of the range found by this grid search and the range that you specify in the **RANGE=** suboption. This option is useful for data exhibiting features at multiple scales, because in such cases the **SELECT=** criterion often has multiple local minima. Using the **PRESEARCH** option increases the likelihood that the golden section search will find the global minimum of the **SELECT=** criterion. See [Example 50.4](#) for such an example.

RANGE(*lower,upper*)

specifies that only smoothing parameter values greater than or equal to *lower* and less than or equal to *upper* be examined.

STEPS

specifies that all models evaluated in the selection process be displayed.

For models with one dependent variable, if you specify neither the **SELECT=** nor the **SMOOTH=** options in the **MODEL** statement, then PROC LOESS uses **SELECT=AICC**.

The following table summarizes how the smoothing parameter values are chosen for various combinations of the **SMOOTH=** option, the **SELECT=** option, and the **SELECT=** option modifiers.

Table 50.3 Smoothing Parameter Value(s) Used for Combinations of **SMOOTH=** and **SELECT=** OPTIONS for Models with One Dependent Variable

Syntax	Search Method	Search Domain
<i>default</i>	golden section using AICC	(0, 1]
SMOOTH=list	no selection	values in <i>list</i>
SMOOTH=list SELECT=criterion	global	values in <i>list</i>
SMOOTH=list SELECT=criterion (RANGE(<i>l, u</i>))	global	values in <i>list</i> within [<i>l, u</i>]
SELECT=criterion	golden section	(0, 1]
SELECT=criterion (RANGE(<i>l,u</i>))	golden section	[<i>l, u</i>]
SELECT=criterion (GLOBAL)	global	(0, 1]
SELECT=criterion (GLOBAL RANGE(<i>l, u</i>))	global	[<i>l, u</i>]

Some examples of using the SELECT= option follow:

SELECT=GCV	specifies selection that uses the GCV <i>criterion</i> .
SELECT=DF1(6.3)	specifies selection that uses the DF1 <i>DFCriterion</i> with target value 6.3.
SELECT=AICC(STEPS)	specifies selection that uses the AICC <i>criterion</i> , showing all step details.
SELECT=DF2(7 GLOBAL)	specifies selection that uses a global search algorithm to find the smoothing parameter that yields the DF2 <i>DFCriterion</i> closest to the target value 7.

NOTE: The SELECT= option cannot be used for models with more than one dependent variable.

SMOOTH=*value-list*

specifies a list of positive smoothing parameter values. If you do not specify the SELECT= option in the MODEL statement, then a separate fit is obtained for each SMOOTH= value specified. If you do specify the SELECT= option, then models with all values specified in the SMOOTH= list are examined, and PROC LOESS selects the value that minimizes the criterion specified in the SELECT= option.

For models with two or more dependent variables, if the SMOOTH= option is not specified in the MODEL statement, then SMOOTH=0.5 is used as a default.

STD

specifies that standard errors of the mean predicted values be included in the “Output Statistics” table. The use of this option implicitly selects the model option DFMETHOD=EXACT if the DFMETHOD= option has not been explicitly used.

T

specifies that *t* statistics are to be included in the “Output Statistics” table. The use of this option implicitly selects the model option DFMETHOD=EXACT if the DFMETHOD= option has not been explicitly used.

TRACEL

specifies that the trace of the prediction matrix as well as the GCV and AICC statistics be included in the “Fit Summary” table. The use of any of the MODEL statement options ALL, CLM, DFMETHOD=EXACT, DIRECT, SELECT=, STD, and T implicitly selects the TRACEL option.

SCORE Statement

SCORE < DATA=SAS-data-set > < ID=(variable list) > < / options > ;

The fitted loess model is used to score the data in the specified SAS data set. This data set must contain all the regressor variables specified in the **MODEL** statement. Furthermore, when a **BY**

statement is used, the score data set must also contain all the BY variables sorted in the order of the BY variables. A SCORE statement is optional, and more than one SCORE statement can be used. SCORE statements cannot be used if the DIRECT option is specified in the MODEL statement. The optional ID= (*variable list*) specifies ID variables to be included in the “Score Results” table.

You find the results of the SCORE statement in the “Score Results” table. This table contains all the data in the data set named in the SCORE statement, including observations with missing values. However, only those observations with nonmissing regressor variables are scored. If no data set is named in the SCORE statement, the data set named in the PROC LOESS statement is scored. You use the PRINT option in the SCORE statement to request that the “Score Results” table be displayed. You can place the “Score Results” table in an output data set by using an ODS OUTPUT statement even if this table is not displayed.

The following options are available in the SCORE statement after a slash (/).

CLM

requests that $100(1-\alpha)\%$ confidence limits on the mean predicted value be added to the “Score Results” table. By default the 95% limits are computed; the ALPHA= option in the MODEL statement can be used to change the significance level. The use of this option implicitly selects the model option DFMETHOD=EXACT if the DFMETHOD= option has not been explicitly used.

PRINT < (VAR=*variables*) >

specifies that the “Score Results” table be displayed. By default only the variables named in the MODEL statement, the variables listed in the ID list in the SCORE statement, and the scored dependent variables are displayed. You can use the VAR= option to specify additional variables in the score data set that are to be included in the displayed output. Note, however, that all columns in the SCORE data set are placed in the SCORE results table, even if you do not request that they be included in the displayed output.

RESIDUAL | R

requests that residuals be added to the “Score Results” table. If the data set you specify in DATA= option in the SCORE statement does not contain one or more of the model dependent variables, then the corresponding residual values in the “Score Results” table are set to missing.

SCALEDINDEP

specifies that scaled regressor coordinates be included in the “Score Results” table. This option is ignored if the SCALE= option is not specified in the MODEL statement.

STEPS

requests that all models evaluated during smoothing parameter value selection be scored, provided that the SELECT= option together with the STEPS modifier is specified in the MODEL statement. By default only the selected model is scored.

WEIGHT Statement

WEIGHT *variable* ;

The WEIGHT statement specifies a variable in the input data set that contains values to be used as a priori weights for a loess fit.

The values of the weight variable must be nonnegative. If an observation's weight is zero, negative, or missing, the observation is deleted from the analysis.

Details: LOESS Procedure

Missing Values

PROC LOESS deletes any observation with missing values for any variable specified in the [MODEL](#) statement. This enables the procedure to reuse the kd tree for all the dependent variables that appear in the [MODEL](#) statement. If you have multiple dependent variables with different missing value structures for the same set of independent variables, you might want to use separate PROC LOESS steps for each dependent variable.

Output Data Sets

PROC LOESS assigns a name to each table it creates. You can use the ODS OUTPUT statement to place one or more of these tables in output data sets. See the section “[ODS Table Names](#)” on page 3831 for a list of the table names created by PROC LOESS. For detailed information about ODS, see Chapter 20, “[Using the Output Delivery System](#).”

For example, the following statements create an output data set named MyOutStats containing the “Output Statistics” table and an output data set named MySummary containing the “Fit Summary” table.

```
proc loess data=Melanoma;
  model Incidences=Year;
  ods output OutputStatistics = MyOutStats
             FitSummary       = MySummary;
run;
```

Often, a single MODEL statement describes more than one model. For example, the following statements fit eight different models (four smoothing parameter values for each dependent variable).

```
proc loess;
  model y1 y2 = x1 x2 x3/smooth =0.1 to 0.7 by 0.2;
  ods output OutputStatistics = MyOutStats;
run;
```

The eight “Output Statistics” tables for these models are stacked in a single data set called MyOutStats. The data set contains a column named DepVarName and a column named SmoothingParameter that

distinguish each model (see [Figure 50.8](#) for an example). If you want the “Output Statistics” table for each model to be in its own data set, you can use the `MATCH_ALL` option in the `ODS OUTPUT` statement. The following statements create eight data sets named `MyOutStats`, `MyOutStats1`, ..., `MyOutStats7`.

```
proc loess;
  model y1 y2 = x1 x2 x3/smooth =0.1 to 0.7 by 0.2;
  ods output OutputStatistics(match_all) = MyOutStats;
run;
```

For further options available in the `ODS OUTPUT` statement, see Chapter 20, “[Using the Output Delivery System](#).”

Only the “Scale Details” and “Fit Summary” tables are displayed by default. The other tables are optionally displayed by using the `DETAILS` option in the `MODEL` statement and the `PRINT` option in the `SCORE` statement. Note that it is not necessary to display a table in order for that table to be used in an `ODS OUTPUT` statement. For example, the following statements display the “Output Statistics” and “kd Tree” tables but place the “Output Statistics” and “Prediction at Vertices” tables in output data sets.

```
proc loess data=Melanoma;
  model Incidences=Year/details(OutputStatistics kdTree);
  ods output OutputStatistics = MyOutStats
             PredAtVertices   = MyVerticesOut;
run;
```

Using the `DETAILS` option alone causes all tables to be displayed.

The `MODEL` statement options `CLM`, `RESIDUAL`, `STD`, `SCALEDINDEP`, and `T` control which optional columns are added to the `OutputStatistics` table. For example, to obtain an `OutputStatistics` output data set containing residuals and confidence limits in addition to the model variables and predicted value, you need to specify the `RESIDUAL` and `CLM` options in the `MODEL` statement as in the following example:

```
proc loess data=Melanoma;
  model Incidences=Year/residual clm;
  ods output OutputStatistics = MyOutStats;
run;
```

Finally, note that using the `ALL` option in the `MODEL` statement causes all optional columns to be included in the output. Also, `ID` columns can be added to the `OutputStatistics` table by using the `ID` statement.

Data Scaling

The loess algorithm to obtain a predicted value at a given point in the predictor space proceeds by doing a least squares fit that uses all data points close to the given point. Thus the algorithm depends critically on the metric used to define closeness. This has the consequence that if you have more than one predictor variable and these predictor variables have significantly different scales, then closeness

depends almost entirely on the variable with the largest scaling. It also means that merely changing the units of one of your predictors can significantly change the loess model fit.

To circumvent this problem, it is necessary to standardize the scale of the independent variables in the loess model. The `SCALE=` option in the `MODEL` statement is provided for this purpose. PROC LOESS uses a symmetrically trimmed standard deviation as the scale estimate for each independent variable of the loess model. This is a robust scale estimator in that extreme values of a variable are discarded before estimating the data scaling. For example, to compute a 10% trimmed standard deviation of a sample, you discard the smallest and largest 5% of the data and compute the standard deviation of the remaining 90% of the data points. In this case, the trimming fraction is 0.1.

For example, the following statement specifies that the variables `Temperature` and `Catalyst` are scaled before performing the loess fitting. In this case, because the trimming fraction is 0.1, the scale estimate used for each of these variables is a 10% trimmed standard deviation.

```
model Yield=Temperature Catalyst / scale = SD(0.1);
```

The default trimming fraction used by PROC LOESS is 0.1 and need not be specified by the `SCALE=` option. Thus the following `MODEL` statement is equivalent to the previous `MODEL` statement.

```
model Yield=Temperature Catalyst / scale = SD;
```

If the `SCALE=` option is not specified, no scaling of the independent variables is done. This is appropriate when there is only a single independent variable or when all the independent variables are a priori scaled similarly.

When the `SCALE=` option is specified, the scaling details for each independent variable are added to the `ScaleDetails` table (see [Output 50.3.2](#) for an example). By default, this table contains only the minimum and maximum values of each independent variable in the model. Finally, note that when the `SCALE=` option is used, specifying the `SCALEDINDEP` option in the `MODEL` statement adds the scaled values of the independent variables to the `OutputStatistics` and `PredAtVertices` tables. If the `SCALEDINDEP` option is specified in the `SCORE` statement, then scaled values of the independent variables are included in the `ScoreResults` table. By default, only the unscaled values are placed in these tables.

Direct versus Interpolated Fitting

Local regression to obtain a predicted value at a given point in the predictor space is done by doing a least squares fit that uses all data points in a local neighborhood of the given point. This method is computationally expensive because a local neighborhood must be determined and a least squares problem must be solved for each point at which a fitted value is required. A faster method is to obtain such fits at a representative sample of points in the predictor space and to obtain fitted values at all other points by interpolation.

PROC LOESS can fit models by using either of these two methods. By default, PROC LOESS uses fitting at a sample of points and interpolation. The method fitting a local model at every data point is selected by specifying the `DIRECT` option in the `MODEL` statement.

kd Trees and Blending

PROC LOESS uses a kd tree to divide the box (also called the *initial cell* or *bucket*) enclosing all the predictor data points into rectangular cells. The vertices of these cells are the points at which local least squares fitting is done.

Starting from the initial cell, the direction of the longest cell edge is selected as the split direction. The median of this coordinate of the data in the cell is the split value. The data in the starting cell are partitioned into two child cells. The left child consists of all data from the parent cell whose coordinate in the split direction is less than the split value. This procedure is repeated for each child cell that has more than a prespecified number of points, called the *bucket size* of the kd tree.

You can specify the bucket size with the BUCKET= option in the **MODEL** statement. If you do not specify the BUCKET= option, the default value used is the largest integer less than or equal to $ns/5$, where n is the number of observations and s is the value of the smoothing parameter. Note that if fitting is being done for a range of smoothing parameter values, the bucket size can change for each value.

The set of vertices of all the cells of the kd tree are the points at which PROC LOESS performs its local fitting. The fitted value at an original data point (or at any other point within the original data cell) is obtained by blending the fitted values at the vertices of the kd tree cell that contains that data point.

The univariate blending methods available in PROC LOESS are linear and cubic polynomial interpolation, with linear interpolation being the default. You can request cubic interpolation by specifying the INTERP=CUBIC option in the **MODEL** statement. In this case, PROC LOESS uses the unique cubic polynomial whose values and first derivatives match those of the fitted local polynomials evaluated at the two endpoints of the kd tree cell edge.

In the multivariate case, such univariate interpolating polynomials are computed on each edge of the kd tree cells and are combined using blending functions (Gordon 1971). In the case of two regressors, if you specify INTERP=CUBIC in the **MODEL** statement, PROC LOESS uses Hermite cubic polynomials as blending functions. If you do not specify INTERP=CUBIC, or if you specify a model with more than two regressors, then PROC LOESS uses linear polynomials as blending functions. In these cases, the blending method reduces to tensor product interpolation from the 2^p vertices of each kd tree cell, where p is the number of regressors.

While the details of the kd tree and the fitted values at the vertices of the kd tree are implementation details that seldom need to be examined, PROC LOESS does provide options for their display. Each kd tree subdivision of the data used by PROC LOESS is placed in the “kdTree” table. The predicted values at the vertices of each kd tree are placed in the “PredAtVertices” table. You can request these tables by using the DETAILS option in the **MODEL** statement.

Local Weighting

The size of the local neighborhoods that PROC LOESS uses in performing local fitting is determined by the smoothing parameter value s . When $s < 1$, the local neighborhood used at a point x contains the s fraction of the data points closest to the point x . When $s \geq 1$, all data points are used.

Suppose q denotes the number of points in the local neighborhoods and d_1, d_2, \dots, d_q denote the distances in increasing order of the q points closest to x . The point at distance d_i from x is given a weight w_i in the local regression that decreases as the distance from x increases. PROC LOESS uses a tricube weight function to define

$$w_i = \frac{32}{5} \left(1 - \left(\frac{d_i}{d_q} \right)^3 \right)^3$$

If $s > 1$, then d_q is replaced by $d_q s^{1/p}$ in the previous formula, where p is the number of predictors in the model.

Finally, note that if a weight variable has been specified using a **WEIGHT** statement, then w_i is multiplied by the corresponding value of the specified weight variable.

Iterative Reweighting

PROC LOESS can do iterative reweighting to improve the robustness of the fit in the presence of outliers in the data. Iterative reweighting is also appropriate when statistical inference is requested and the error distribution is symmetric but not Gaussian.

The number of iterations is specified by the **ITERATIONS=** option in the **MODEL** statement. The default is **ITERATIONS=1**, which corresponds to no reweighting.

At iterations beyond the first iteration, the local weights w_i of the previous section are replaced by $r_i w_i$, where r_i is a weight that decreases as the residual of the fitted value at the previous iteration at the point corresponding to d_i increases. Refer to Cleveland and Grosse (1991) and Cleveland, Grosse, and Shyu (1992) for details.

Specifying the Local Polynomials

PROC LOESS uses linear or quadratic polynomials in doing the local least squares fitting. The option **DEGREE =** in the **MODEL** statement is used to specify the degree of the local polynomials used by PROC LOESS, with **DEGREE = 1** being the default. In addition, when **DEGREE = 2** is specified, the **MODEL** statement **DROPSQUARE=** option can be used to exclude specific monomials during the least squares fitting.

For example, the following statements use the monomials 1, x_1 , x_2 , $x_1 \times x_2$, and $x_2 \times x_2$ for the local least squares fitting.

```
proc loess;
  model y= x1 x2/ degree=2 dropsquare=(x1);
run;
```

Smoothing Matrix

When no iterative reweighting is done, the “Smoothing Matrix” denoted by L defines the linear relationship between the fitted and observed dependent variable values of a loess model. You can obtain the predicted values of a loess fit from the observed values via

$$\hat{y} = Ly$$

where y is the vector of observed values and \hat{y} is the corresponding vector of predicted values of the dependent variable. Note that L is an n by n matrix, where n is the number of observations in the analysis. PROC LOESS does not explicitly form L if the DFMETHOD=EXACT option is not explicitly or implicitly selected.

Model Degrees of Freedom

The approximate model degrees of freedom in a nonparametric fit is a number that is analogous to the number of free parameters in a parametric model. There are three commonly used measures of model degrees of freedom in nonparametric models. These criteria are as follows:

$$\begin{aligned} \text{DF1} &\equiv \text{Trace}(L) \\ \text{DF2} &\equiv \text{Trace}(L^T L) \\ \text{DF3} &\equiv 2\text{Trace}L - \text{Trace}(L^T L) \end{aligned}$$

A discussion of their properties can be found in Hastie and Tibshirani (1990). DF2 is also referred to as the “Equivalent Number of Parameters,” and this is the name that PROC LOESS uses for DF2 when it appears in the “Fit Summary” table.

Statistical Inference and Lookup Degrees of Freedom

If you denote the i th measurement of the response by y_i and the corresponding measurement of predictors by x_i , then

$$y_i = g(x_i) + \epsilon_i$$

where g is the regression function and ϵ_i are independent random errors with mean zero. If the errors are normally distributed with constant variance, then you can obtain confidence intervals for the predictions from PROC LOESS. You can also obtain confidence limits in the case where ϵ_i is heteroscedastic but $a_i\epsilon_i$ has constant variance and a_i are a priori weights that are specified using the **WEIGHT** statement of PROC LOESS. You can do inference in the case in which the error

distribution is symmetric by using iterative reweighting. Formulas for doing statistical inference under the preceding conditions can be found in Cleveland and Grosse (1991) and Cleveland, Grosse, and Shyu (1992). Cleveland and Grosse (1991) show that standardized residuals for a loess model follow a t distribution with ρ degrees of freedom where

$$\begin{aligned}\delta_1 &\equiv \text{Trace}(I - L)^T(I - L) \\ \delta_2 &\equiv \text{Trace}\left((I - L)^T(I - L)\right)^2 \\ \rho &\equiv \text{Lookup Degrees of Freedom} \\ &\equiv \delta_1^2/\delta_2\end{aligned}$$

The residual standard error that you find in the “Fit Summary” table is defined by

$$\text{Residual Standard Error} \equiv \sqrt{\text{Residual SS}/\delta_1}$$

The determination of ρ is computationally expensive and is not done by default. It is computed if you specify the `DFMETHOD=EXACT` or `DFMETHOD=APPROX` option in the **MODEL** statement. It is also computed if you specify any of the options `CLM`, `STD`, and `T` in the **MODEL** statement. Note that the values of δ_1 , δ_2 , and ρ are reported in the “Fit Summary” table.

If you specify the `CLM` option in the **MODEL** statement, confidence limits are added to the `OutputStatistics` table. By default, 95% limits are computed, but you can change this by using the `ALPHA=` option in the **MODEL** statement.

Automatic Smoothing Parameter Selection

There are several methodologies for automatic smoothing parameter selection. One class of methods chooses the smoothing parameter value to minimize a criterion that incorporates both the tightness of the fit and model complexity. Such a criterion can usually be written as a function of the error mean square, $\hat{\sigma}^2$, and a penalty function designed to decrease with increasing smoothness of the fit. This penalty function is usually defined in terms of the smoothing matrix L (see the section “[Smoothing Matrix](#)” on page 3826).

Examples of specific criteria are generalized cross validation (Craven and Wahba 1979) and the Akaike information criterion (Akaike 1973). These classical selectors have two undesirable properties when used with local polynomial and kernel estimators: they tend to undersmooth small data sets and tend to be nonrobust in the sense that small variations of the input data can change the choice of smoothing parameter value significantly. Hurvich, Simonoff, and Tsai (1998) obtained several corrected AIC criteria that address the small-sample bias and perform comparably with the *plug-in selectors* (Ruppert, Sheather, and Wand 1995). PROC LOESS provides automatic smoothing parameter selection that uses two of these corrected AIC criteria, named AIC_{C_1} and AIC_C in Hurvich, Simonoff, and Tsai (1998), and generalized cross validation, denoted by GCV.

The relevant formulas are

$$\begin{aligned}
 AICC_1 &= n \log(\hat{\sigma}^2) + n \frac{\delta_1/\delta_2(n + \nu_1)}{\delta_1^2/\delta_2 - 2} \\
 AICC &= \log(\hat{\sigma}^2) + 1 + \frac{2(\text{Trace}(L) + 1)}{n - \text{Trace}(L) - 2} \\
 GCV &= \frac{n\hat{\sigma}^2}{(n - \text{Trace}(L))^2}
 \end{aligned}$$

where n is the number of observations and

$$\begin{aligned}
 \delta_1 &\equiv \text{Trace}(I - L)^T(I - L) \\
 \delta_2 &\equiv \text{Trace}\left((I - L)^T(I - L)\right)^2 \\
 \nu_1 &\equiv \text{Equivalent Number of Parameters} \\
 &\equiv \text{Trace}(L^T L)
 \end{aligned}$$

You invoke these methods for automatic smoothing parameter selection by specifying the `SELECT=criterion` option in the **MODEL** statement, where *criterion* is AICC1, AICC, or GCV. The LOESS procedure evaluates the specified criterion for a sequence of smoothing parameter values and selects the value in this sequence that minimizes the specified criterion. If multiple values yield the optimum, then the largest of these values is selected.

A second class of methods seeks to set an approximate measure of model degrees of freedom to a specified target value. These methods are useful for making meaningful comparisons between loess fits and other nonparametric and parametric fits. Three approximate model degrees of freedom for a loess model are defined in the section “[Model Degrees of Freedom](#)” on page 3826. You invoke these methods by specifying the `SELECT=DFCriterion(target)` option in the **MODEL** statement, where *DFCriterion* is DF1, DF2, or DF3. The criterion that is minimized is given in the following table.

Table 50.4 Minimization Criteria

Syntax	Minimization Criterion
SELECT=DF1(<i>target</i>)	$\text{Trace}(L) - \text{target}$
SELECT=DF2(<i>target</i>)	$\text{Trace}(L^T L) - \text{target}$
SELECT=DF3(<i>target</i>)	$2\text{Trace}(L) - \text{Trace}(L^T L) - \text{target}$

The results are summarized in the “Smoothing Criterion” table. This table is displayed whenever automatic smoothing parameter selection is performed. You can obtain details of the sequence of models examined by specifying the `DETAILS(MODELSUMMARY)` option in the **MODEL** statement to display the “Model Summary” table.

There are several ways in which you can control the sequence of models examined by PROC LOESS. If you specify the `SMOOTH=value-list` option in the `MODEL` statement, then only the values in this list are examined in performing the selection. For example, the following statements select the model that minimizes the AICC1 criterion among the three models with smoothing parameter values 0.1, 0.3, and 0.4:

```
proc loess;
  model y= x1/ smooth=0.1 0.3 0.4 select=AICC1;
run;
```

If you do not specify the `SMOOTH=` option in the `MODEL` statement, then by default PROC LOESS uses a golden section search method to find a local minimum of the specified criterion in the range (0, 1]. You can use the `RANGE(lower,upper)` modifier in the `SELECT=` option to change the interval in which the golden section search is performed. For example, the following statements request a golden section search to find a local minimizer of the GCV criterion for smoothing parameter values in the interval [0.1,0.5]:

```
proc loess;
  model y= x1/select=GCV( range(0.1,0.5) );
run;
```

If you want to be sure of obtaining a global minimum in the range of smoothing parameter values examined, you can specify the `GLOBAL` modifier in the `SELECT=` option. For example, the following statements request that a global minimizer of the AICC criterion be obtained for smoothing parameter values in the interval [0.2, 0.8]:

```
proc loess;
  model y= x1/select=AICC( global range(0.2,0.8) );
run;
```

Note that even though the smoothing parameter is a continuous variable, a given range of smoothing parameter values corresponds to a finite set of local models. For example, for a data set with 100 observations, the range [0.2, 0.4] corresponds to models with 20, 21, 22, . . . , 40 points in the local neighborhoods. If the `GLOBAL` modifier is specified, all possible models in the range are evaluated sequentially.

Note that by default PROC LOESS displays a “Fit Summary” and other optionally requested tables only for the selected model. You can request that these tables be displayed for all models in the selection process by adding the `STEPS` modifier in the `SELECT=` option. Also note that by default scoring requested with `SCORE` statements is done only for the selected model. However, if you specify the `STEPS` in both the `MODEL` and `SCORE` statements, then all models evaluated in the selection process are scored.

In terms of computation, $AICC$, GCV , and $DF1$ depend on the smoothing matrix L only through its trace. In the direct method, this trace can be computed efficiently. In the interpolated method that uses kd trees, there is some additional computational cost but the overall work is not significant compared to the rest of the computation. In contrast, the quantities δ_1 , δ_2 , and ν_1 that appear in the $AICC_1$ criterion, and the $DF2$ and $DF3$ criteria, depend on the entire L matrix and for this reason, the time needed to compute these quantities dominates the time required for the model fitting. Hence `SELECT=AICC1`, `SELECT=DF2`, and `SELECT=DF3` are much more computationally expensive than `SELECT=AICC`, `SELECT=GCV`, and `SELECT=DF1`, especially when combined with the `GLOBAL` modifier. Hurvich, Simonoff, and Tsai (1998) note that $AICC$ can be regarded as an approximation of $AICC_1$ and that “the $AICC$ selector generally performs well in all circumstances.”

For models with one dependent variable, PROC LOESS uses `SELECT=AICC` as its default, if you specify neither the `SMOOTH=` nor the `SELECT=` option in the **MODEL** statement. With two or more dependent variables, automatic smoothing parameter selection needs to be done separately for each dependent variable. For this reason automatic smoothing parameter selection is not available for models with multiple dependent variables. In such cases you should use a separate PROC LOESS step for each dependent variable, if you want to use automatic smoothing parameter selection.

Sparse and Approximate Degrees of Freedom Computation

As noted in the section “Statistical Inference and Lookup Degrees of Freedom” on page 3826, obtaining confidence limits in loess models requires the computation of the lookup degrees of freedom. This in turn requires the computation of

$$\delta_2 \equiv \text{Trace} \left((I - L)^T (I - L) \right)^2$$

where L is the loess smoothing matrix (see the section “Smoothing Matrix” on page 3826).

The work in a direct implementation of this formula grows as n^3 , where n is the number of observations in analysis. For large n , this work dominates the time needed to fit the loess model itself. To alleviate this computational bottleneck, Cleveland and Grosse (1991) and Cleveland, Grosse, and Shyu (1992) developed approximate methods for estimating this quantity in terms of more readily computable statistics. A different approach to obtaining a computationally cheap estimate of δ_2 has been implemented in PROC LOESS.

For large data sets with significant local structure, the loess model is often used with small values of the smoothing parameter. Recalling that the smoothing parameter defines the fraction of the data used in each local regression, this means that the loess fit at any point in regressor space depends on only a small fraction of the data. This is reflected in the smoothing matrix L whose (i, j) th entry is nonzero only if the i th and j th observations lie in at least one common local neighborhood. Hence the smoothing matrix is a sparse matrix (has mostly zero entries) in such cases. By exploiting this sparsity, PROC LOESS now computes δ_2 orders of magnitude faster than in previous implementations.

When each local neighborhood contains a large subset of the data—i.e., when the smoothing parameter is large—then it is no longer true that the smoothing matrix is sparse. However, since a point in a local neighborhood is given a local weight that decreases with its distance from the center of the neighborhood, many of the coefficients in the smoothing matrix turn out to be nonzero but with orders of magnitude smaller than that of the larger coefficients in the matrix. The approximate method for computing δ_2 that has been implemented in PROC LOESS exploits these disparities in magnitudes of the elements in the smoothing matrix by setting the small elements to zero. This creates a sparse approximation of the smoothing matrix to which the fast sparse methods can be applied.

In order to decide the threshold at which elements in the smoothing matrix are set to zero, PROC LOESS samples the elements in the smoothing matrix to obtain the value of the element in a specified lower quantile in this sample. The magnitude of the element at this quantile is used as a cutoff value, and all elements in the smoothing matrix whose magnitude is less than this cutoff are set to zero.

for the approximate computation. By default all elements in the lower 90th percentile are set to zero. You can use the `DFMETHOD=APPROX(QUANTILE=)` option in the **MODEL** statement to change this value. As you increase the value for the quantile to be zeroed, you speed up the degrees of freedom computation at the expense of increasing approximation errors. You can also use the `DFMETHOD=APPROX(CUTOFF=)` option in the **MODEL** statement to specify the cutoff value directly.

For small data sets, the approximate computation is not needed and would be rougher than for larger data sets. Hence PROC LOESS performs the exact computation for analyses with fewer than 500 points, even if `DFMETHOD=APPROX` is specified in the model statement. Also, for small values of the smoothing parameter, elements in the lower specified quantile might already all be zero. In such cases the approximate method is the same as the exact method. PROC LOESS labels as approximate any statistics that depend on the approximate computation of δ_2 only in the cases where the approximate computation was used and is different from the exact computation.

Scoring Data Sets

One or more **SCORE** statements can be used with PROC LOESS. A data set that includes all the variables specified in the **MODEL** and **BY** statements must be specified in each **SCORE** statement. Score results are placed in the ScoreResults table. This table is not displayed by default, but specifying the **PRINT** option in the **SCORE** statement produces the table. If you specify the **CLM** option in the **SCORE** statement, confidence intervals are included in the ScoreResults table.

Note that scoring is not supported when the **DIRECT** option is specified in the **MODEL** statement. Scoring at a point specified in a score data set is done by first finding the cell in the kd tree containing this point and then interpolating the scored value from the predicted values at the vertices of this cell. This methodology precludes scoring any points that are not contained in the box that surrounds the data used in fitting the loess model.

ODS Table Names

PROC LOESS assigns a name to each table it creates. You can use these names to reference the table when using the Output Delivery System (ODS) to select tables and create output data sets. These names are listed in the following table. For more information about ODS, see Chapter 20, “Using the Output Delivery System.”

Table 50.5 ODS Tables Produced by PROC LOESS

ODS Table Name	Description	Statement	Option
FitSummary	Specified fit parameters and fit summary		default
kdTree	Structure of kd tree used	MODEL	DETAILS(kdTree)

Table 50.5 *continued*

ODS Table Name	Description	Statement	Option
ModelSummary	Summary of all models evaluated	MODEL	DETAILS(ModelSummary)
OutputStatistics	Coordinates and fit results at input data points	MODEL	DETAILS(OutputStatistics)
PredAtVertices	Coordinates and fitted values at kd tree vertices	MODEL	DETAILS(PredAtVertices)
ScaleDetails	Extent and scaling of the independent variables		default
ScoreResults	Coordinates and fit results at scoring points	SCORE	PRINT
SmoothingCriterion	Criterion value and selected smoothing parameter	MODEL	SELECT

ODS Graphics

To request graphics with PROC LOESS, you must first enable ODS Graphics by specifying the **ODS GRAPHICS ON** statement. See Chapter 21, “[Statistical Graphics Using ODS](#),” for more information. You can reference every graph produced through ODS Graphics with a name. The names of the graphs that PROC LOESS generates are listed in [Table 50.6](#), along with the relevant PLOTS= options.

Table 50.6 ODS Graphics Produced by PROC LOESS

ODS Graph Name	Plot Description	PLOTS Option
ContourFitPanel	Panel of loess contour surfaces overlaid on scatter plots of data	CONTOURFITPANEL
ContourFit	Loess contour surface overlaid on scatter plot of data	CONTOURFITPANEL
DiagnosticsPanel	Panel of fit diagnostics	DIAGNOSTICS
FitPanel	Panel of loess curves overlaid on scatter plots of data	FITPANEL
FitPlot	Loess curve overlaid on scatter plot of data	FIT
ObservedByPredicted	Dependent variable versus loess fit	OBSERVEDBYPREDICTED
QQPlot	Normal quantile plot of residuals	QQPLOT
ResidualsBySmooth	Panel of residuals versus regressor by smoothing parameter values	RESIDUALSBYSMOOTH
ResidualByPredicted	Residuals versus loess fit	RESIDUALBYPREDICTED
ResidualHistogram	Histogram of fit residuals	RESIDUALHISTOGRAM
ResidualPanel	Panel of residuals versus regressors for fixed smoothing parameter value	RESIDUALS

Table 50.6 *continued*

ODS Graph Name	Plot Description	PLOTS Option
ResidualPlot	Plot of residuals versus regressor	RESIDUALS
RFPlot	Side-by-side plots of quantiles of centered fit and residuals	RFPLOT
ScorePlot	Loess fit evaluated at scoring points	SCOREPLOT
CriterionPlot	Selection criterion versus smoothing parameter	CRITERION

Examples: LOESS Procedure

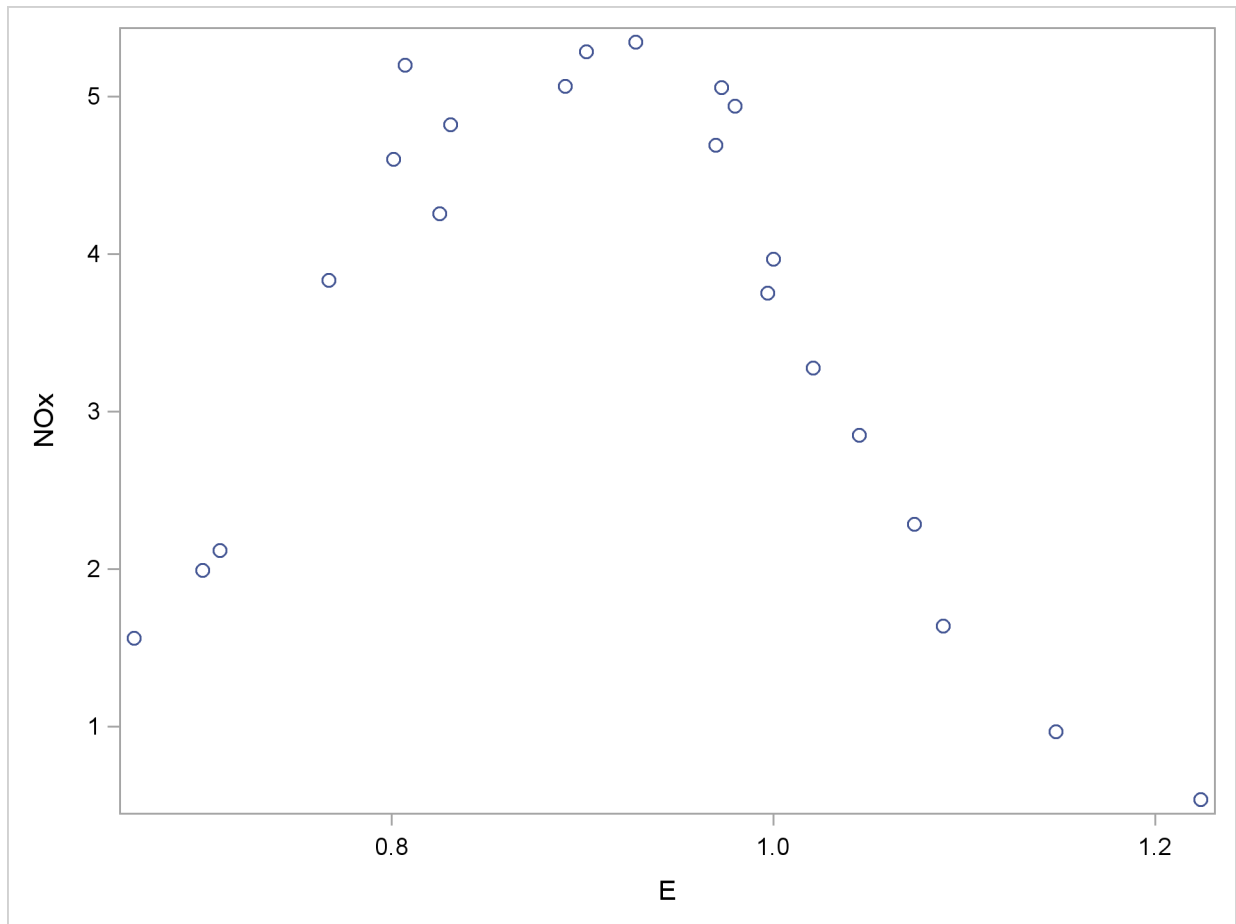
Example 50.1: Engine Exhaust Emissions

Investigators studied the exhaust emissions of a one-cylinder engine (Brinkman 1981). The SAS data set Gas contains the results data. The dependent variable, NO_x, measures the concentration, in micrograms per joule, of nitric oxide and nitrogen dioxide normalized by the amount of work of the engine. The independent variable, E, is a measure of the richness of the air and fuel mixture.

```
data Gas;
  input NOx E @@;
  format NOx f3.1;
  format E f3.1;
datalines;
4.818 0.831 2.849 1.045
3.275 1.021 4.691 0.97
4.255 0.825 5.064 0.891
2.118 0.71 4.602 0.801
2.286 1.074 0.97 1.148
3.965 1 5.344 0.928
3.834 0.767 1.99 0.701
5.199 0.807 5.283 0.902
3.752 0.997 0.537 1.224
1.64 1.089 5.055 0.973
4.937 0.98 1.561 0.665
;
```

The following PROC SGPLOT statements produce the simple scatter plot of these data displayed in [Output 50.1.1](#).

```
proc sgplot data=Gas;
  scatter x=E y=NOx;
run;
```

Output 50.1.1 Scatter Plot of the Gas Data

The following statements fit two loess models for these data. Because this is a small data set, it is reasonable to do direct fitting at every data point. As there is substantial curvature in the data, quadratic local polynomials are used. An ODS OUTPUT statement creates two output data sets containing the “Output Statistics” and “Fit Summary” tables.

```
ods graphics on;

proc loess data=Gas;
  ods output OutputStatistics = GasFit
             FitSummary=Summary;
  model NOx = E / degree=2 select=AICC(steps) smooth = 0.6 1.0
                direct alpha=.01 all details;
run;

ods graphics off;
```

Output 50.1.2 Fit Summary Table

The LOESS Procedure	
Selected Smoothing Parameter: 0.6	
Dependent Variable: NOx	
Fit Summary	
Fit Method	Direct
Number of Observations	22
Degree of Local Polynomials	2
Smoothing Parameter	0.60000
Points in Local Neighborhood	13
Residual Sum of Squares	1.71852
Trace[L]	6.42184
GCV	0.00708
AICC	-0.45637
AICC1	-9.39715
Delta1	15.12582
Delta2	14.73089
Equivalent Number of Parameters	5.96950
Lookup Degrees of Freedom	15.53133
Residual Standard Error	0.33707

The “Fit Summary” table for smoothing parameter value 0.6, shown in [Output 50.1.2](#), records the fitting parameters specified and some overall fit statistics. See the section “[Smoothing Matrix](#)” on page 3826 for a definition of the smoothing matrix L , and the sections “[Model Degrees of Freedom](#)” on page 3826 and “[Statistical Inference and Lookup Degrees of Freedom](#)” on page 3826 for definitions of the statistics that appear in this table.

The “Output Statistics” table for smoothing parameter value 0.6 is shown in [Output 50.1.3](#). Note that, because the ALL option is specified in the **MODEL** statement, this table includes all the relevant optional columns. Furthermore, because the ALPHA=0.01 option is specified in the **MODEL** statement, the confidence limits in this table are 99% limits.

Output 50.1.3 Output Statistics Table

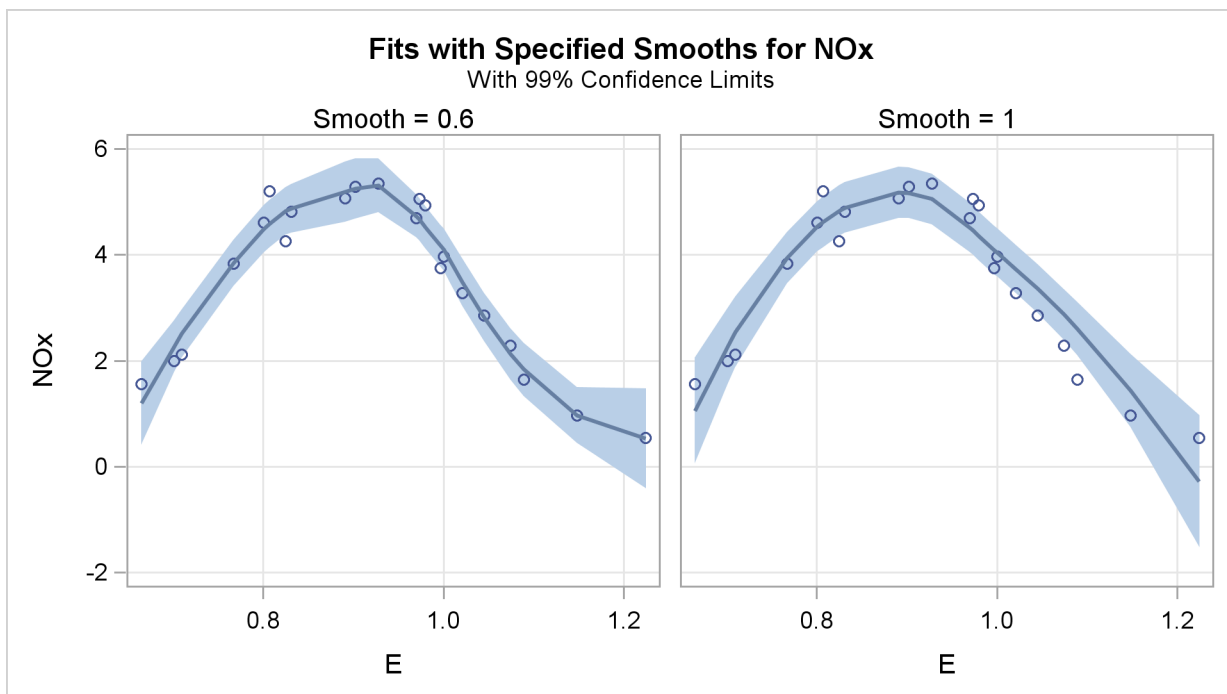
The LOESS Procedure						
Selected Smoothing Parameter: 0.6						
Dependent Variable: NOx						
Output Statistics						
Obs	E	NOx	Predicted NOx	Estimated Prediction Std Deviation	Residual	t Value
1	0.8	4.8	4.87377	0.15528	-0.05577	-0.36
2	1.0	2.8	2.81984	0.15380	0.02916	0.19
3	1.0	3.3	3.48153	0.15187	-0.20653	-1.36
4	1.0	4.7	4.73249	0.13923	-0.04149	-0.30
5	0.8	4.3	4.82305	0.15278	-0.56805	-3.72
6	0.9	5.1	5.18561	0.19337	-0.12161	-0.63
7	0.7	2.1	2.51120	0.15528	-0.39320	-2.53
8	0.8	4.6	4.48267	0.15285	0.11933	0.78
9	1.1	2.3	2.12619	0.16683	0.15981	0.96
10	1.1	1.0	0.97120	0.18134	-0.00120	-0.01
11	1.0	4.0	4.09987	0.13477	-0.13487	-1.00
12	0.9	5.3	5.31258	0.17283	0.03142	0.18
13	0.8	3.8	3.84572	0.14929	-0.01172	-0.08
14	0.7	2.0	2.26578	0.16712	-0.27578	-1.65
15	0.8	5.2	4.58394	0.15363	0.61506	4.00
16	0.9	5.3	5.24741	0.19319	0.03559	0.18
17	1.0	3.8	4.16979	0.13478	-0.41779	-3.10
18	1.2	0.5	0.53059	0.32170	0.00641	0.02
19	1.1	1.6	1.83157	0.17127	-0.19157	-1.12
20	1.0	5.1	4.66733	0.13735	0.38767	2.82
21	1.0	4.9	4.52385	0.13556	0.41315	3.05
22	0.7	1.6	1.19888	0.26774	0.36212	1.35
Output Statistics						
Obs	99% Confidence Limits					
1	4.41841	5.32912				
2	2.36883	3.27085				
3	3.03617	3.92689				
4	4.32419	5.14079				
5	4.37503	5.27107				
6	4.61855	5.75266				
7	2.05585	2.96655				
8	4.03444	4.93089				
9	1.63697	2.61541				
10	0.43942	1.50298				
11	3.70467	4.49507				
12	4.80576	5.81940				
13	3.40794	4.28350				
14	1.77571	2.75584				
15	4.13342	5.03445				
16	4.68089	5.81393				
17	3.77457	4.56502				
18	-0.41278	1.47397				
19	1.32933	2.33380				
20	4.26456	5.07010				
21	4.12632	4.92139				
22	0.41375	1.98401				

Output 50.1.4 Output Statistics Table

Optimal Smoothing Criterion	
AICC	Smoothing Parameter
-0.45637	0.60000

The combination of the options `SELECT=AICC` and `SMOOTH=0.6 1` in the **MODEL** statement specifies that PROC LOESS fit models with smoothing parameters of 0.6 and 1 and select the model that yields the smaller value of the AICC statistic. The “Smoothing Criterion” shown in **Output 50.1.4** shows that PROC LOESS selects the model with smoothing parameter value 0.6 as it yields the smaller value of the AICC statistic.

With ODS Graphics enabled, PROC LOESS produces a panel of fit plots whenever you specify the `SMOOTH=` option in the **MODEL** statement. These fit plots include confidence limits if you additionally specify the `CLM` option in the **MODEL** statement.

Output 50.1.5 Loess Fits with 99% Confidence Limits for the Gas Data

Output 50.1.5 shows the “Fit Panel” that displays the fitted models with 99% confidence limits overlaid on scatter plots of the data.

Based on the AICC criterion, the model with smoothing parameter 0.6 is preferred. You can address the question of whether the differences between these models are significant using analysis of variance. You do this by using the model with smoothing parameter value 1 as the null model.

The statistic

$$F = \frac{(\text{rss}^{(n)} - \text{rss}) / (\delta_1^{(n)} - \delta_1)}{\text{rss} / \delta_1}$$

has a distribution that is well approximated by an F distribution with

$$\nu = \frac{(\delta_1^{(n)} - \delta_1)^2}{\delta_2^{(n)} - \delta_2}$$

numerator degrees of freedom and ρ denominator degrees of freedom (Cleveland and Grosse 1991). Here quantities with superscript n refer to the null model, rss is the residual sum of squares, and δ_1 , δ_2 , and ρ are defined in the section “Statistical Inference and Lookup Degrees of Freedom” on page 3826.

The “Fit Summary” tables contain the information needed to carry out such an analysis. These tables have been captured in the output data set named `Summary` by using an `ODS OUTPUT` statement. The following statements extract the relevant information from this data set and carry out the analysis of variance:

```
data h0 h1;
  set Summary(keep=SmoothingParameter Label1 nValue1
               where=(Label1 in ('Residual Sum of Squares', 'Delta1',
                                'Delta2', 'Lookup Degrees of Freedom')));
  if SmoothingParameter = 1 then output h0;
  else output h1;
run;

proc transpose data=h0(drop=SmoothingParameter Label1) out=h0;

data h0(drop=_NAME_); set h0;
  rename Col1 = RSSNull
         Col2 = delta1Null
         Col3 = delta2Null;

proc transpose data=h1(drop=SmoothingParameter Label1) out=h1;

data h1(drop=_NAME_); set h1;
  rename Col1 = RSS      Col2 = delta1
         Col3 = delta2   Col4 = rho;

data ftest; merge h0 h1;
  nu = (delta1Null - delta1)**2 / (delta2Null - delta2);
  Numerator = (RSSNull - RSS) / (delta1Null - delta1);
  Denominator = RSS / delta1;
  FValue = Numerator / Denominator;
  PValue = 1 - ProbF(FValue, nu, rho);
  label nu      = 'Num DF'   rho      = 'Den DF'
        FValue = 'F Value'  PValue = 'Pr > F';

proc print data=ftest label;
  var nu rho Numerator Denominator FValue PValue;
  format nu rho FValue 7.2 PValue 6.4;
run;
```

The results are shown in [Output 50.1.6](#).

Output 50.1.6 Test ANOVA for Loess Models of Gas Data

Obs	Num DF	Den DF	Numerator	Denominator	F Value	Pr > F
1	2.67	15.53	1.05946	0.11362	9.32	0.0012

The small p -value confirms that the fit with smoothing parameter value 0.6 is significantly different from the loess model with smoothing parameter value 1.

Example 50.2: Sulfate Deposits in the U.S. for 1990

The following data set contains measurements in grams per square meter of sulfate (SO₄) deposits during 1990 at 179 sites throughout the 48 contiguous states.

```
data SO4;
  input Latitude Longitude SO4 @@;
  format Latitude f4.0;
  format Longitude f4.0;
  format SO4 f4.1;
datalines;
32.45833 87.24222 1.403 34.28778 85.96889 2.103
33.07139 109.86472 0.299 36.07167 112.15500 0.304
31.95056 112.80000 0.263 33.60500 92.09722 1.950
... more lines ...
43.87333 104.19222 0.306 44.91722 110.42028 0.210
45.07611 72.67556 2.646
;
```

As longitudes decrease from west to east in the western hemisphere, the roles of east and west get interchanged if you use these longitudes on the horizontal axis of a plot. You can address this by using negative values to represent longitudes in the western hemisphere. The following statements change the sign of longitude in the SO4 data set and define a format to display these negative values with a suffix of “W”.

```
proc format;
  picture latitude -90 - 0 = '000S'
                0 - 90 = '000N';
  picture longitude -180 - 0 = '000W'
                0 - 180 = '000E';
run;
data SO4;
  set SO4;
  format longitude longitude. latitude latitude.;
  longitude = -longitude;
run;
```

The following statements use ODS Graphics to plot the locations of the sulfate measurements. The circles indicating the locations are colored using a gradient that denotes the value of SO4.

```
proc template;
  define statgraph gradientScatter;
    beginGraph;
      layout overlay;
        scatterPlot x=longitude y=latitude /
          markercolorgradient = SO4
          markerattrs         = (symbol=circleFilled)
          colormodel           = ThreeColorLowHigh
          name                 = "Scatter";
        scatterPlot x=longitude y=latitude /
          markerattrs         = (symbol=circle);
        continuousLegend "Scatter"/title= "SO4";
      endlayout;
    endgraph;
  end;
run;

proc sgrender data=SO4 template=gradientScatter;run;
```

Output 50.2.1 Sulfate Measurements

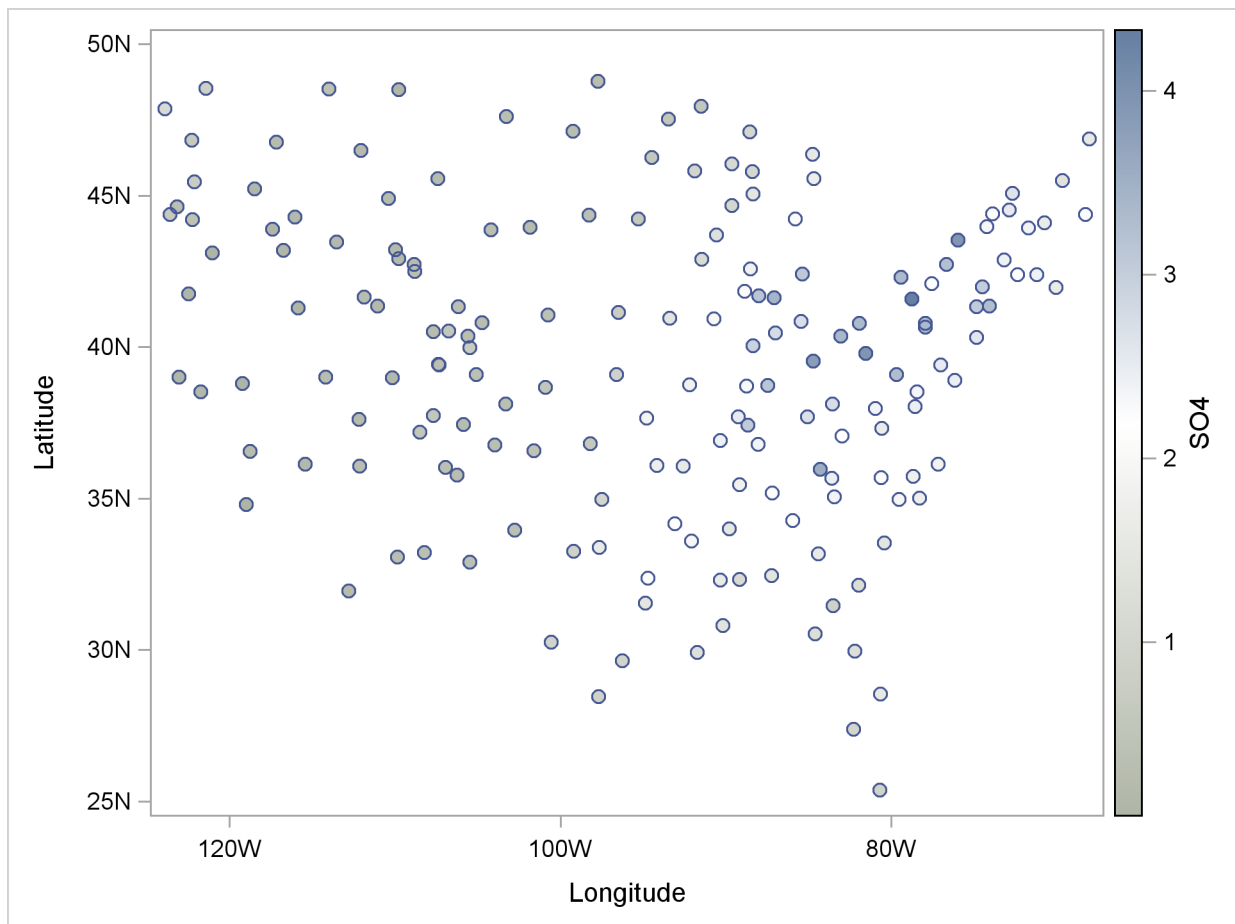


Figure 50.2.1 shows that the largest concentrations of sulfate deposits occur in the northeastern United States.

The following statements fit a loess model.

```
ods graphics on;

proc loess data=SO4;
  model SO4=Longitude Latitude / degree=2 interp=cubic;
run;

ods graphics off;
```

Output 50.2.2 Fit Plot for the SO4 Data

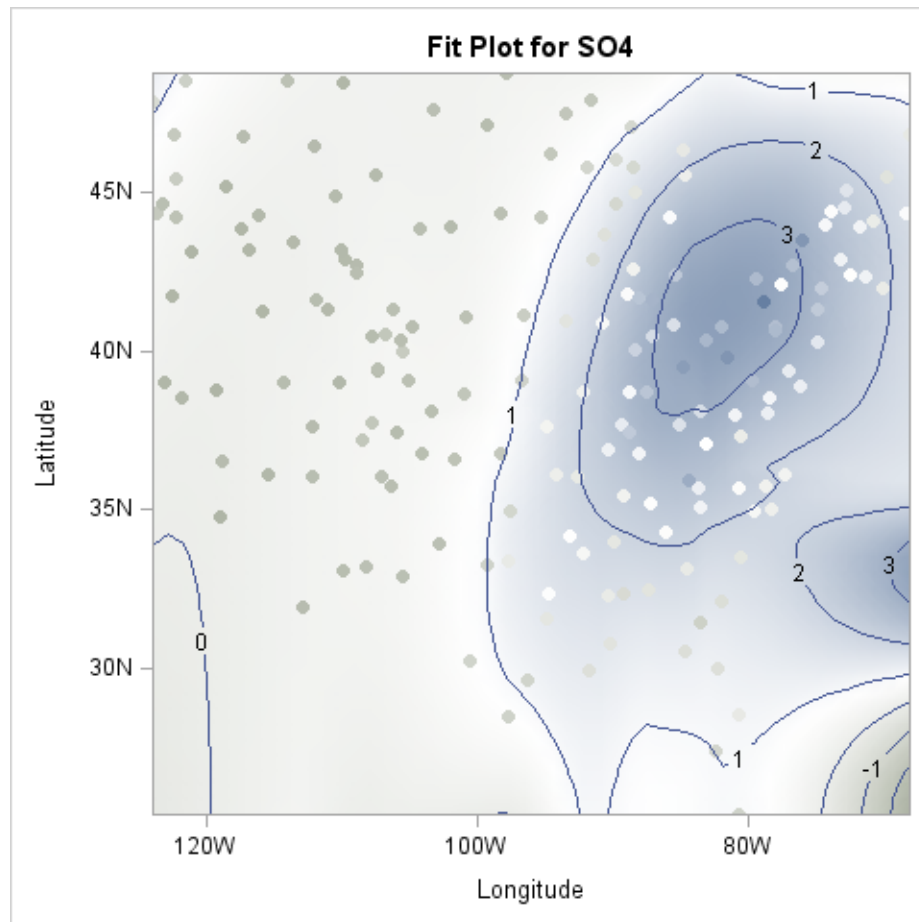


Figure 50.2.2 shows a contour plot of the fitted loess surface overlaid with a scatter plot of the data. The data are colored by the observed sulfate concentrations, using the same color gradient as the gradient-filled contour plot of the fitted surface. Note that for observations where the residual is small, the observations blend in with the contour plot. The greater the size of the residual, the greater the contrast between the observation color and the surface color.

The sulfate measurements are irregularly spaced. To facilitate producing a plot of the fitted loess surface, you can create a data set containing a regular grid of longitudes and latitudes and then use

the **SCORE** statement to evaluate the loess surface at these points. The following statements show how you do this:

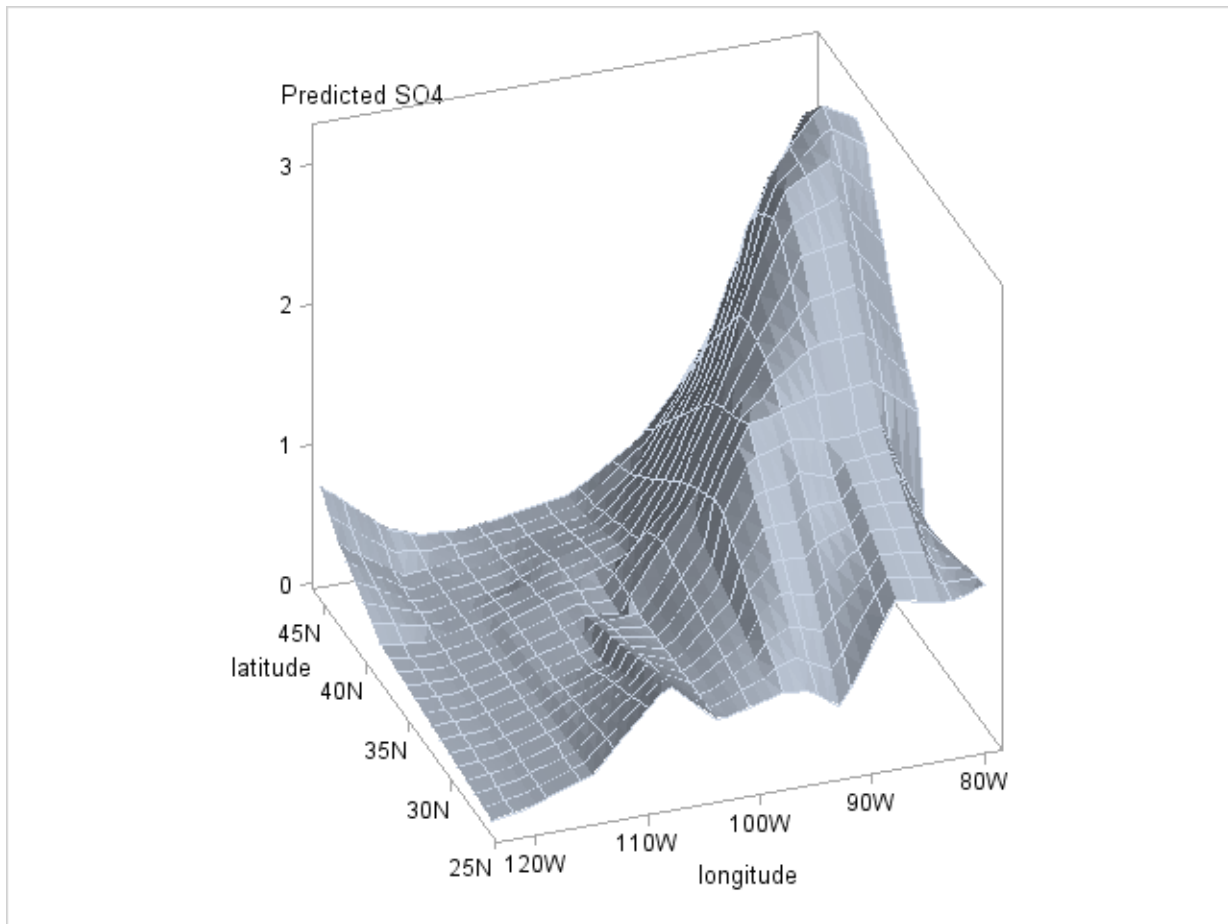
```
data PredPoints;
  format longitude longitude.
         latitude latitude.;
  do Latitude = 26 to 46 by 1;
    do Longitude = -79 to -123 by -1;
      output;
    end;
  end;
run;

proc loess data=S04;
  model S04=Longitude Latitude;
  score data=PredPoints / print;
  ods Output ScoreResults=ScoreOut;
run;
```

The PRINT option in the **SCORE** statement requests that the “Score Results” table be displayed as part of the PROC LOESS output. The ODS OUTPUT statement outputs this table to a data set named ScoreOut. If you do not want to display the score results but you do want the score results in an output data set, then you can omit the PRINT option from the **SCORE** statement. To plot the surface shown in [Figure 50.2.3](#) by using ODS Graphics, use the following statements:

```
proc template;
  define statgraph surface;
    begingraph;
      layout overlay3d / rotate=340 tilt=30 cube=false;
      surfaceplotparm x=Longitude y=Latitude z=p_S04;
    endlayout;
  endgraph;
end;
run;

proc sgrender data=ScoreOut template=surface;
run;
```

Output 50.2.3 Loess Fit of SO₄ Surface**Example 50.3: Catalyst Experiment**

The following data set records the results of an experiment to determine how the yield of a chemical reaction varies with temperature and amount of a catalyst used.

```
data Experiment;
  input Temperature Catalyst MeasuredYield;
  if ranuni(1) < 0.1
    then CorruptedYield = MeasuredYield + 10 * ranuni(1);
    else CorruptedYield = MeasuredYield;
datalines;
80    0.000    6.85601
80    0.002    7.26355
80    0.004    7.41448

... more lines ...

140   0.078    5.20562
140   0.080    5.49371
;
```

The aim of this example is to show how you can use PROC LOESS for robust fitting in the presence of outliers. To simulate an intermittent equipment malfunction, the variable `CorruptedYield` is the same as the variable `MeasuredYield` except for about 10% of the observations where an offset has been added. This example shows how you can use PROC LOESS obtain a fit for `CorruptedYield` that is close to the fit you obtain for `MeasuredYield`.

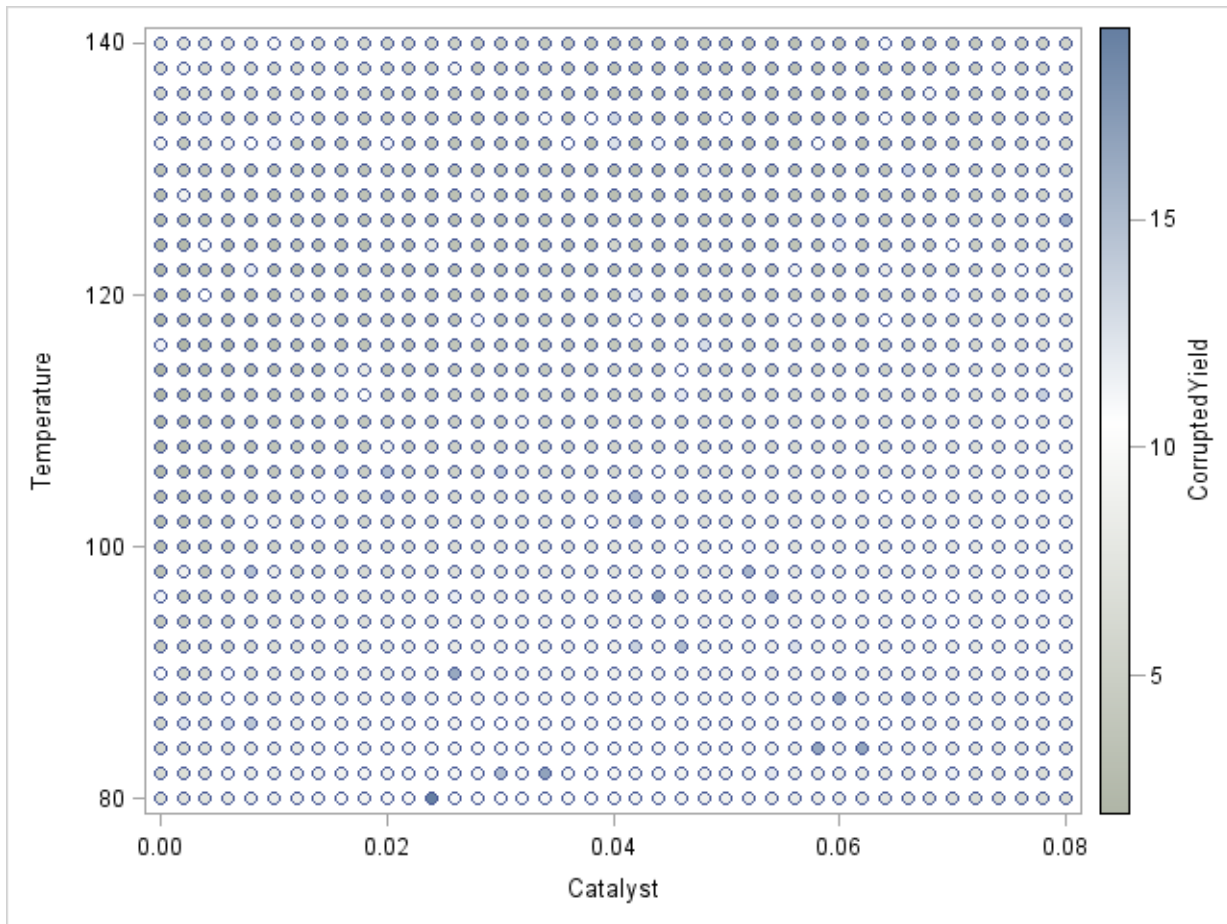
The following statements produce a scatter plot of Temperature by Catalyst where the observations are colored by `CorruptedYield`:

```
proc template;
  define statgraph gradientScatter;
    beginGraph;
      layout overlay;
        scatterPlot x=Catalyst y=Temperature /
          markercolorgradient = CorruptedYield
          markerattrs         = (symbol=circleFilled)
          colormodel           = ThreeColorLowHigh
          name                 = "Yield";

        scatterPlot x=Catalyst y=Temperature /
          markerattrs         = (symbol=circle);

        continuousLegend "Yield" / title= "CorruptedYield";
      endlayout;
    endgraph;
  end;
run;

proc sgrender data=Experiment template=gradientScatter;run;
```

Output 50.3.1 Scatter Plot of Experiment Data Colored by CorruptedYield

Output 50.3.1 shows a scatter plot of the data where the observations are shaded by the value of CorruptedYield. The darkly shaded points that are surrounded by lightly shaded points are points where the simulated incorrect measurements occur.

The following code fits a loess model to the measured data:

```
ods graphics on;

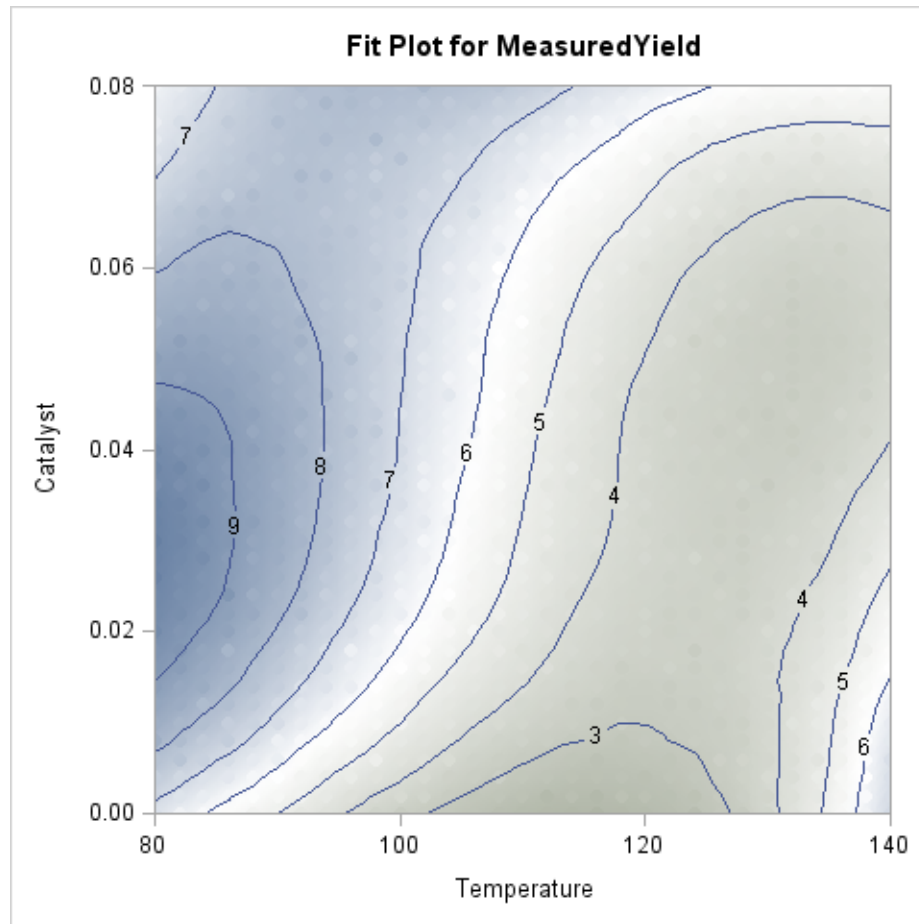
proc loess data=Experiment;
  model MeasuredYield = Temperature Catalyst / scale=sd(0.1);
run;
```

Output 50.3.2 Scale Details for the Experiment Data

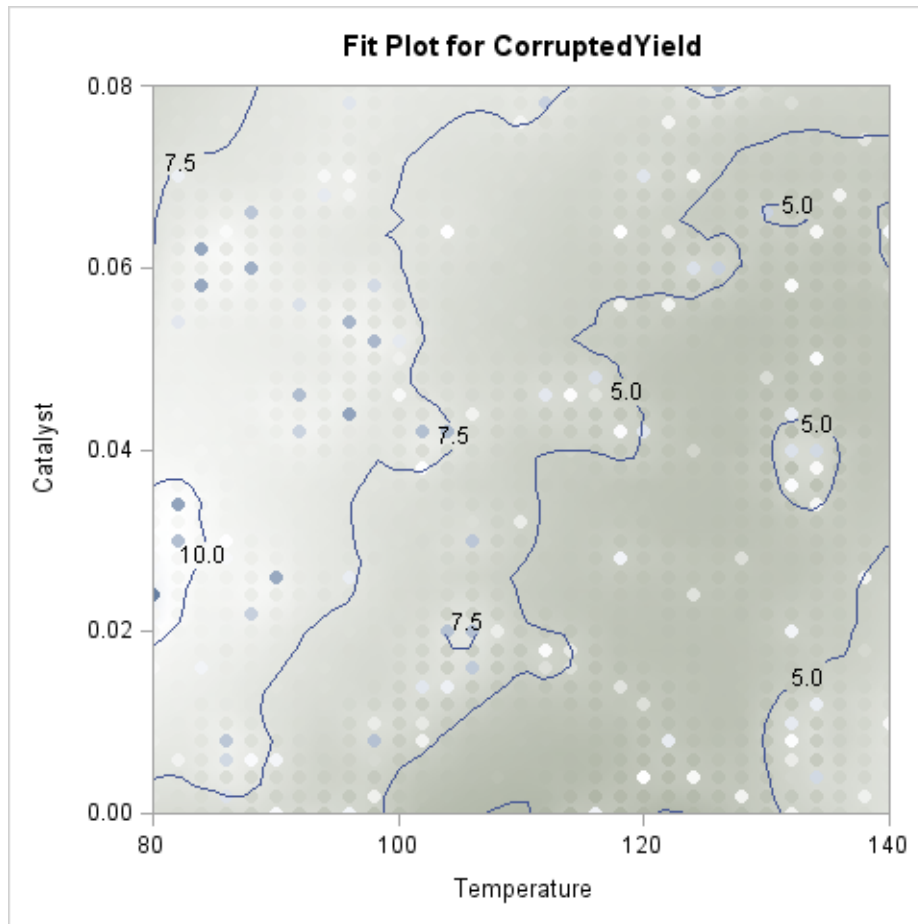
The LOESS Procedure		
Independent Variable Scaling		
Scaling applied: 10% trimmed standard deviation		
Statistic	Temperature	Catalyst
Minimum Value	80.00000	0
Maximum Value	140.00000	0.08000
Trimmed Mean	110.00000	0.04000
Trimmed Standard Deviation	14.32149	0.01894

The SCALE=SD(0.1) option in the **MODEL** statement specifies that the independent variables in the model are to be divided by their respective 10% trimmed standard deviations before the fitted model is computed. This is appropriate because the independent variables Temperature and Catalyst are not similarly scaled. The “Scale Details” table in **Output 50.3.2** displays the details of ranges of the regressors and the scale factors applied to each regressor.

Output 50.3.3 displays the loess fit. Because the fitted surface is a good fit of the observed data, the observations on this plot are not clearly distinguishable from the fitted surface. The results are dramatically different when the outliers are included. The following statements fit a loess model to the corrupted response, using the same smoothing parameter that was selected for the measured response.

Output 50.3.3 Fit for MeasuredYield

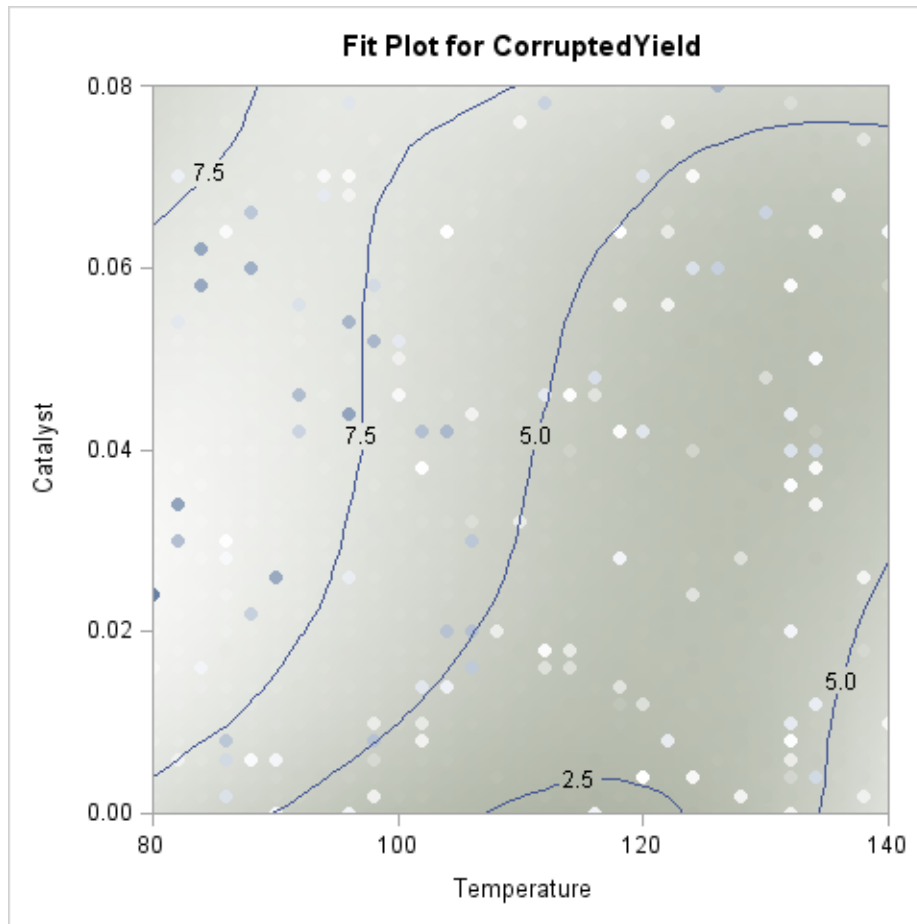
```
proc loess data=Experiment;  
  model CorruptedYield = Temperature Catalyst /  
    scale=sd(0.1) smooth=0.018;  
run;
```

Output 50.3.4 Fit for CorruptedYield

Output 50.3.4 displays the loess fit. The fit is pulled upward in the neighborhoods of these outliers. If you use a larger smoothing parameter value, then these local perturbations in the fit get smoothed out, but at the expense of smoothing away the information in the underlying measured response. In such cases a robust fitting method is indicated. The following statements show how you do this:

```
proc loess data=Experiment;
  model CorruptedYield = Temperature Catalyst /
    scale = sd(0.1)
    smooth = 0.018
    iterations=4;
run;
```

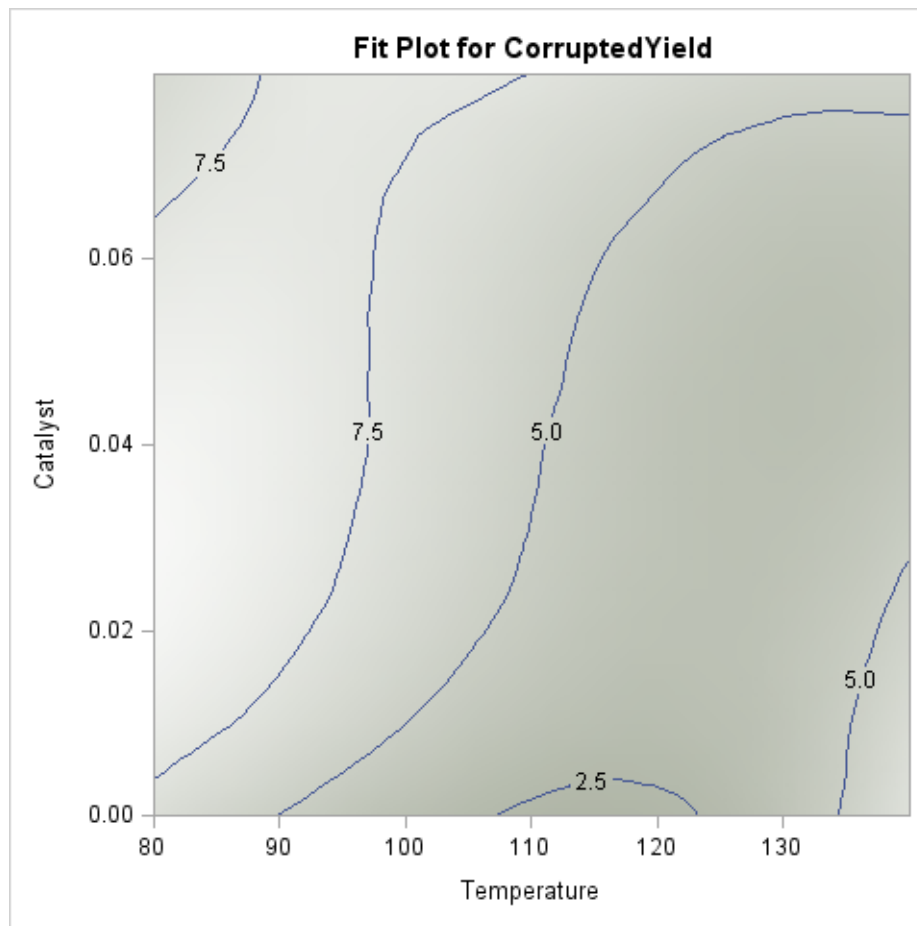
The ITERATIONS=4 option in the **MODEL** statement requests the initial loess fit followed by three iteratively reweighted iterations.

Output 50.3.5 Robust Fit for CorruptedYield

You can see the impact of the robust fitting by comparing the robust fit shown in [Output 50.3.5](#) with the nonrobust fit in [Output 50.3.4](#). In the robust fit you see that the local perturbations caused by the outliers have been eliminated as these the outlying observations get down-weighted during the robustness iterations. By comparing the labeled contours on the fit plot for the uncorrupted response shown in [Output 50.3.3](#) with the labeled contours for the corrupted response shown in [Output 50.3.4](#), you can see that the robust fit has produced a reasonable fit for the underlying measured data. The color gradient in [Output 50.3.5](#) is chosen to accommodate the outliers that are present in the observed data, and so you cannot easily compare the color gradient in this plot with that in [Output 50.3.3](#). The following statements repeat the robust analysis with an option added to suppress the display of the observations on the fit plot:

```
proc loess data=Experiment plots=contourFit(obs=None);
  model CorruptedYield = Temperature Catalyst /
    scale = sd(0.1)
    smooth = 0.018
    iterations=4;
run;

ods graphics off;
```

Output 50.3.6 Robust Fit for CorruptedYield with Observations Suppressed

Output 50.3.6 shows the robust fit with the observations suppressed. The range of the fitted surface values in this plot is similar to the range in **Output 50.3.3**. By comparing this contour plot with the contour plot in **Output 50.3.3**, you clearly see that the robust loess fit has successfully modeled the underlying surface despite the presence of the outliers.

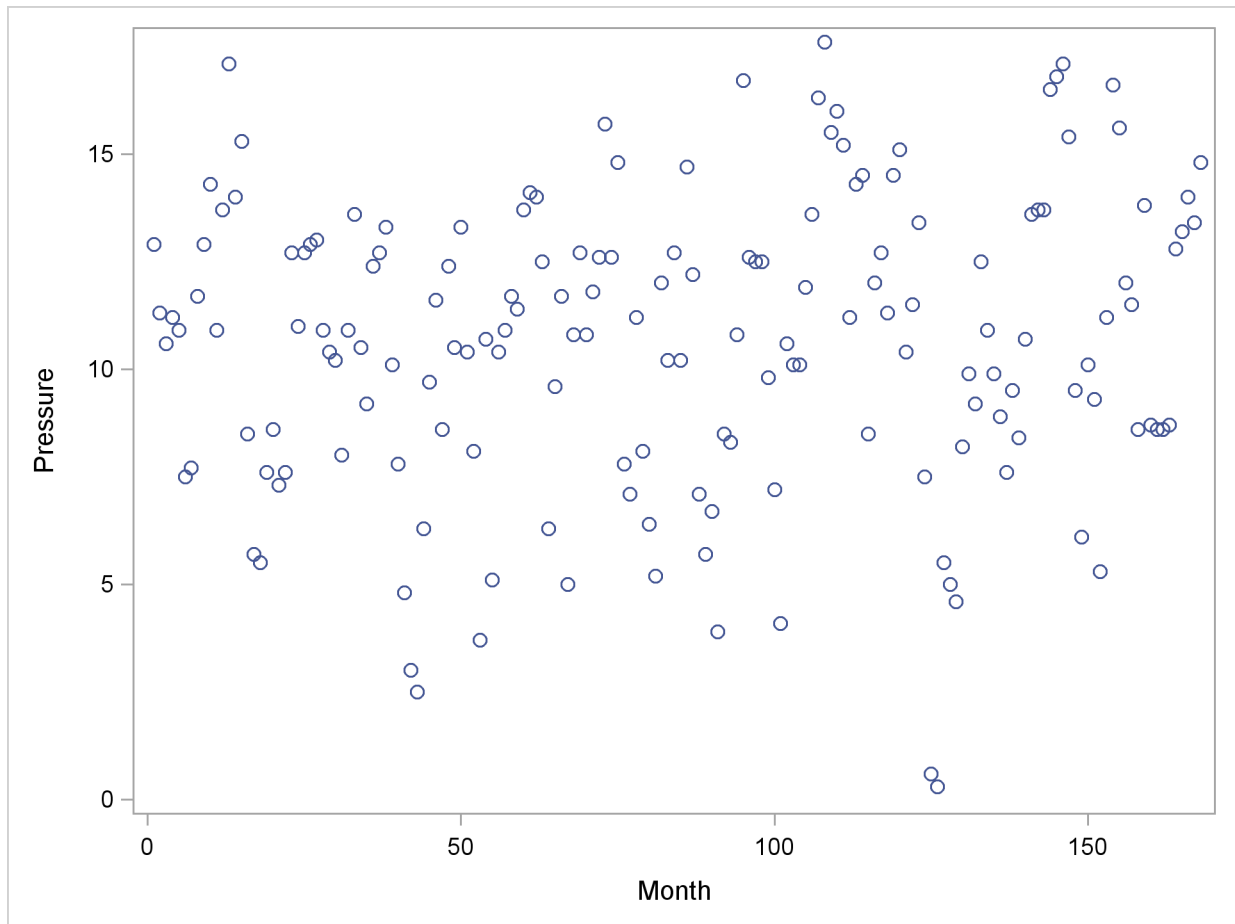
Example 50.4: El Niño Southern Oscillation

The following data set contains measurements of monthly averaged atmospheric pressure differences between Easter Island and Darwin, Australia, for a period of 168 months (National Institute of Standards and Technology 1998):

```
data ENSO;
  input Pressure @@;
  Month=_N_;
  format Pressure 4.1;
  format Month 3.0;
datalines;
12.9 11.3 10.6 11.2 10.9 7.5 7.7 11.7
12.9 14.3 10.9 13.7 17.1 14.0 15.3 8.5
5.7 5.5 7.6 8.6 7.3 7.6 12.7 11.0
12.7 12.9 13.0 10.9 10.4 10.2 8.0 10.9
13.6 10.5 9.2 12.4 12.7 13.3 10.1 7.8
4.8 3.0 2.5 6.3 9.7 11.6 8.6 12.4
10.5 13.3 10.4 8.1 3.7 10.7 5.1 10.4
10.9 11.7 11.4 13.7 14.1 14.0 12.5 6.3
9.6 11.7 5.0 10.8 12.7 10.8 11.8 12.6
15.7 12.6 14.8 7.8 7.1 11.2 8.1 6.4
5.2 12.0 10.2 12.7 10.2 14.7 12.2 7.1
5.7 6.7 3.9 8.5 8.3 10.8 16.7 12.6
12.5 12.5 9.8 7.2 4.1 10.6 10.1 10.1
11.9 13.6 16.3 17.6 15.5 16.0 15.2 11.2
14.3 14.5 8.5 12.0 12.7 11.3 14.5 15.1
10.4 11.5 13.4 7.5 0.6 0.3 5.5 5.0
4.6 8.2 9.9 9.2 12.5 10.9 9.9 8.9
7.6 9.5 8.4 10.7 13.6 13.7 13.7 16.5
16.8 17.1 15.4 9.5 6.1 10.1 9.3 5.3
11.2 16.6 15.6 12.0 11.5 8.6 13.8 8.7
8.6 8.6 8.7 12.8 13.2 14.0 13.4 14.8
;
```

The following PROC SGLOT statements produce the simple scatter plot of these data, displayed in [Output 50.4.1](#).

```
proc sgplot data=ENSO;
  scatter y=Pressure x=Month;
run;
```

Output 50.4.1 Scatter Plot of ENSO Data

You can compute a loess fit and obtain graphical results for these data by using the following statements:

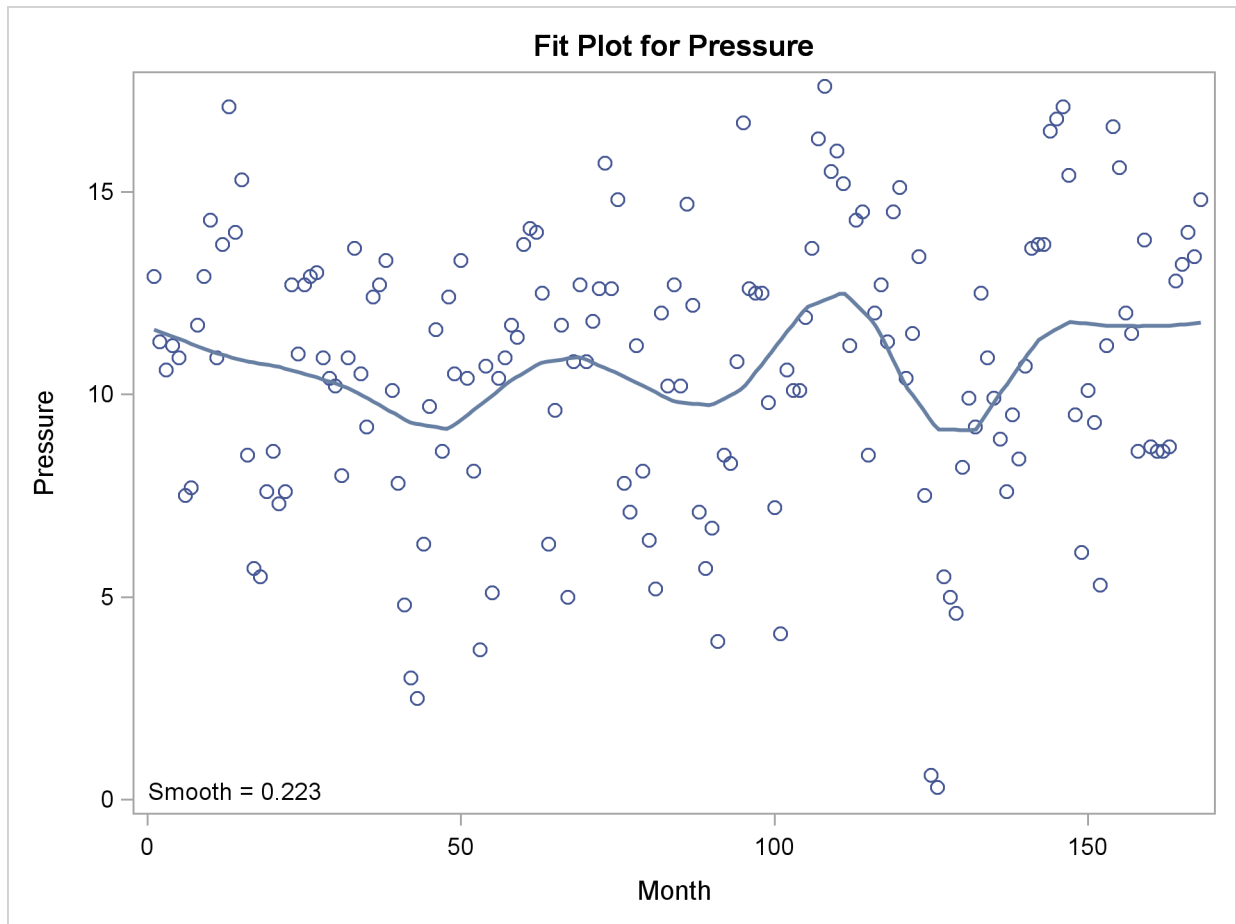
```
ods graphics on;

proc loess data=ENSO plots=residuals(smooth);
  model Pressure=Month;
run;
```

The “Smoothing Criterion” and “Fit Summary” tables are shown in [Output 50.4.2](#), and the fit plot is shown in [Output 50.4.3](#).

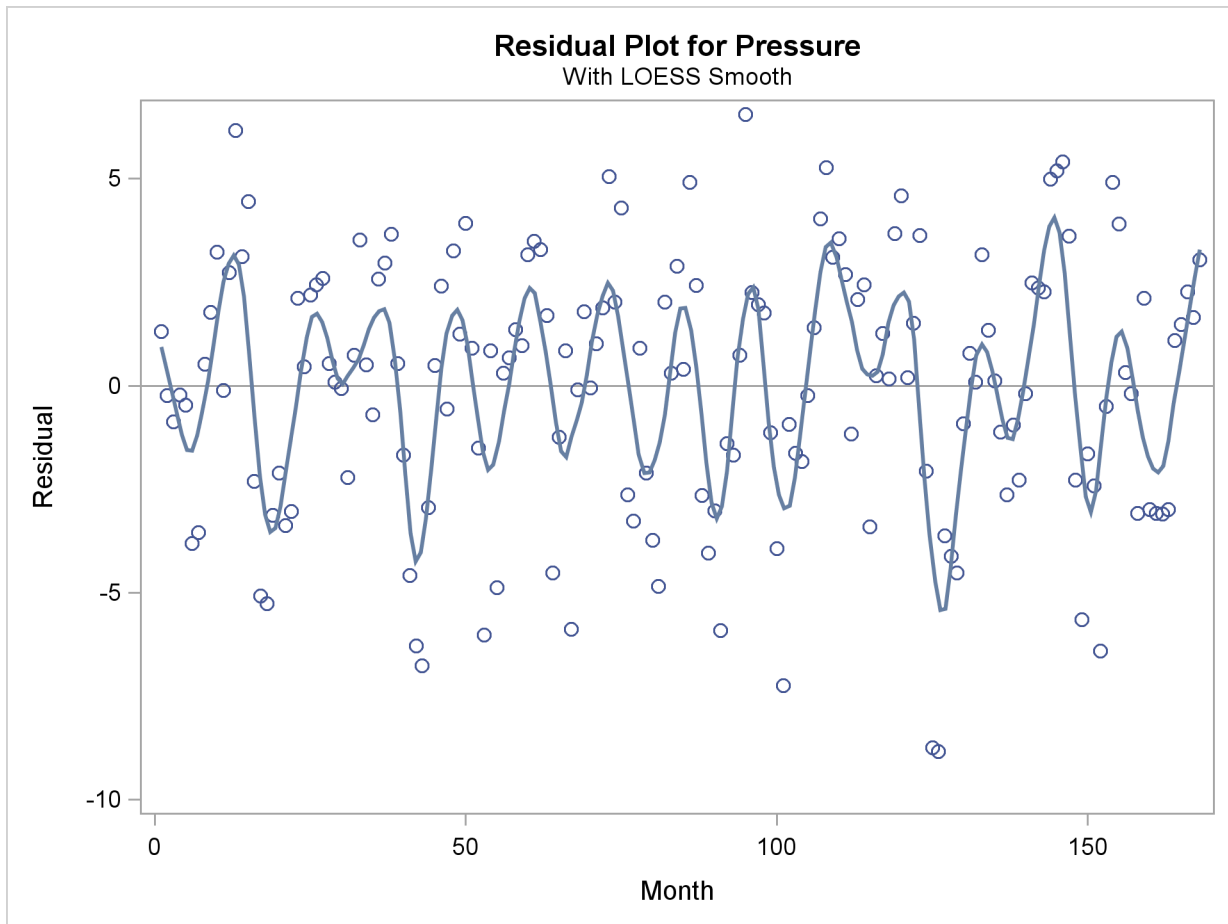
Output 50.4.2 Output from PROC LOESS

The LOESS Procedure	
Dependent Variable: Pressure	
Optimal Smoothing Criterion	
AICC	Smoothing Parameter
3.41105	0.22321
The LOESS Procedure	
Selected Smoothing Parameter: 0.223	
Dependent Variable: Pressure	
Fit Summary	
Fit Method	kd Tree
Blending	Linear
Number of Observations	168
Number of Fitting Points	33
kd Tree Bucket Size	7
Degree of Local Polynomials	1
Smoothing Parameter	0.22321
Points in Local Neighborhood	37
Residual Sum of Squares	1654.27725
Trace[L]	8.74180
GCV	0.06522
AICC	3.41105

Output 50.4.3 Oversmoothed Loess Fit for the ENSO Data

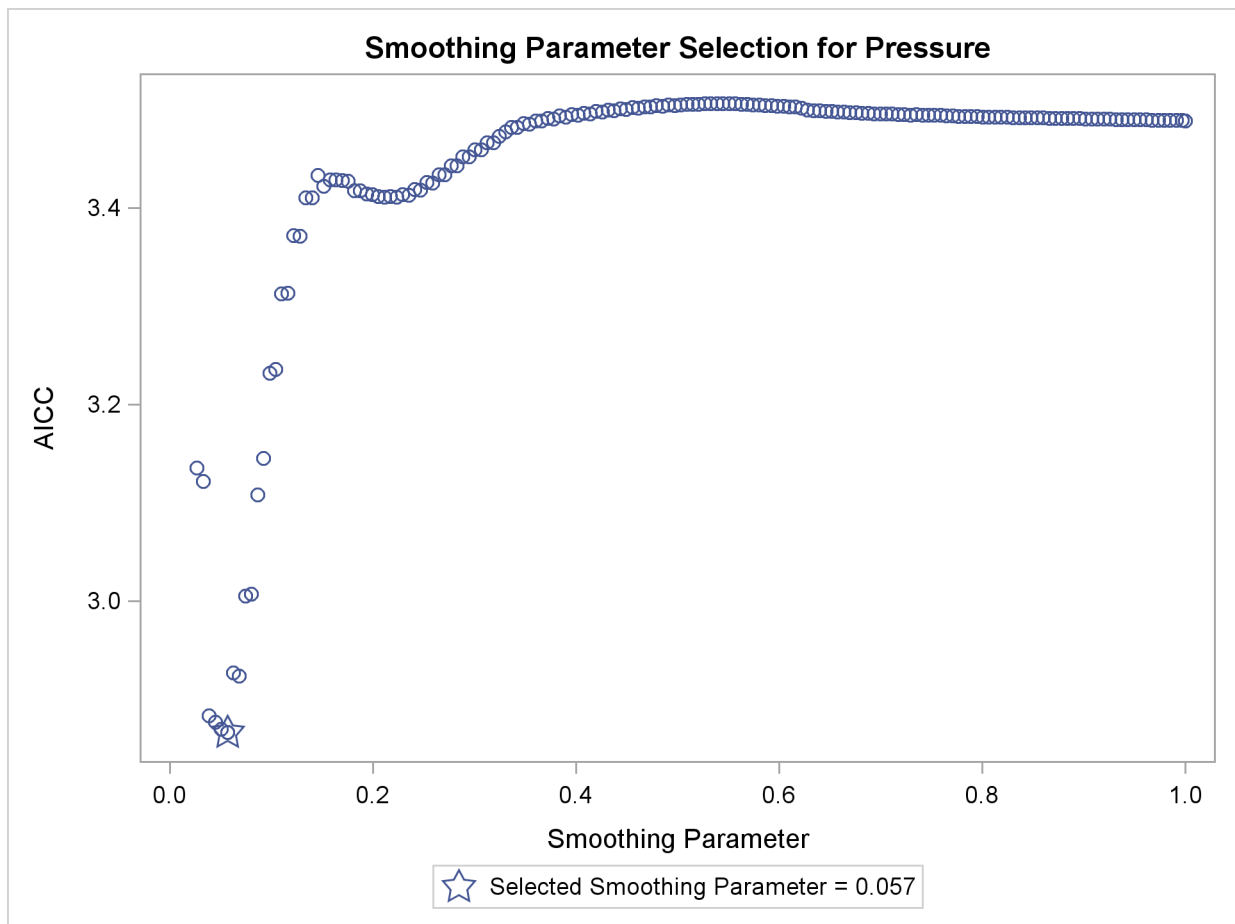
This weather-related data should exhibit an annual cycle. However, the loess fit in [Output 50.4.3](#) indicates a longer cycle but no annual cycle. This suggests that the loess fit is oversmoothed. One way to detect oversmoothing is to look for patterns in the fit residuals. With ODS Graphics enabled, PROC LOESS produces a scatter plot of the residuals versus each regressor in the model. To aid in visually detecting patterns in these scatter plots, it is useful to superimpose a nonparametric fit on these scatter plots. You can request this by specifying the SMOOTH suboption of the PLOTS=RESIDUALS option in the [PROC LOESS](#) statement. The nonparametric fit that is produced is again a loess fit that is produced independently of the loess fit used to obtain these residuals.

With the superimposed loess fit shown in [Output 50.4.4](#), you can clearly identify an annual cycle in the residuals, which confirms that the loess fit for the ENSO is oversmoothed. What accounts for this poor fit?

Output 50.4.4 Residuals for the Loess Fit for the ENSO Data

The smoothing parameter value used for the loess fit shown in [Output 50.4.3](#) was chosen using the default method of PROC LOESS, namely a golden section minimization of the AICC criterion over the interval (0, 1]. One possibility is that the golden section search has found a local rather than a global minimum of the AICC criterion. You can test this by redoing the fit requesting a global minimum. You do this with the following statements:

```
proc loess data=ENSO;
  model Pressure=Month/select=AICC(global);
run;
```

Output 50.4.5 AICC versus Smoothing Parameter Showing Local Minima

The explanation for the oversmoothed fit in [Output 50.4.3](#) is now apparent. [Output 50.4.5](#) shows that the golden section search algorithm found the local minimum that occurs near the value 0.22 of the smoothing parameter rather than the global minimum that occurs near 0.06. Note that if you restrict the range of smoothing parameter values examined to lie below 0.2, then the golden section search finds the global minimum, as the following statements demonstrate:

```
proc loess data=ENSO;
  model Pressure=Month/select=AICC(range(0.03,0.2));
run;
```

Output 50.4.6 Selected Smoothing Parameter Value

The LOESS Procedure	
Dependent Variable: Pressure	
Optimal Smoothing Criterion	
AICC	Smoothing Parameter
2.86660	0.05655

Output 50.4.6 shows that with the restricted range of smoothing parameter values examined, PROC LOESS finds the global minimum of the AICC criterion. Often you might not know an appropriate range of smoothing parameter values to examine. In such cases, you can use the PRESEARCH suboption of the SELECT= option in the MODEL statement. When you specify this option, PROC LOESS does a preliminary search to try to locate a smoothing parameter value range that contains just the first local minimum of the criterion being used for the selection. The following statements provide an example.

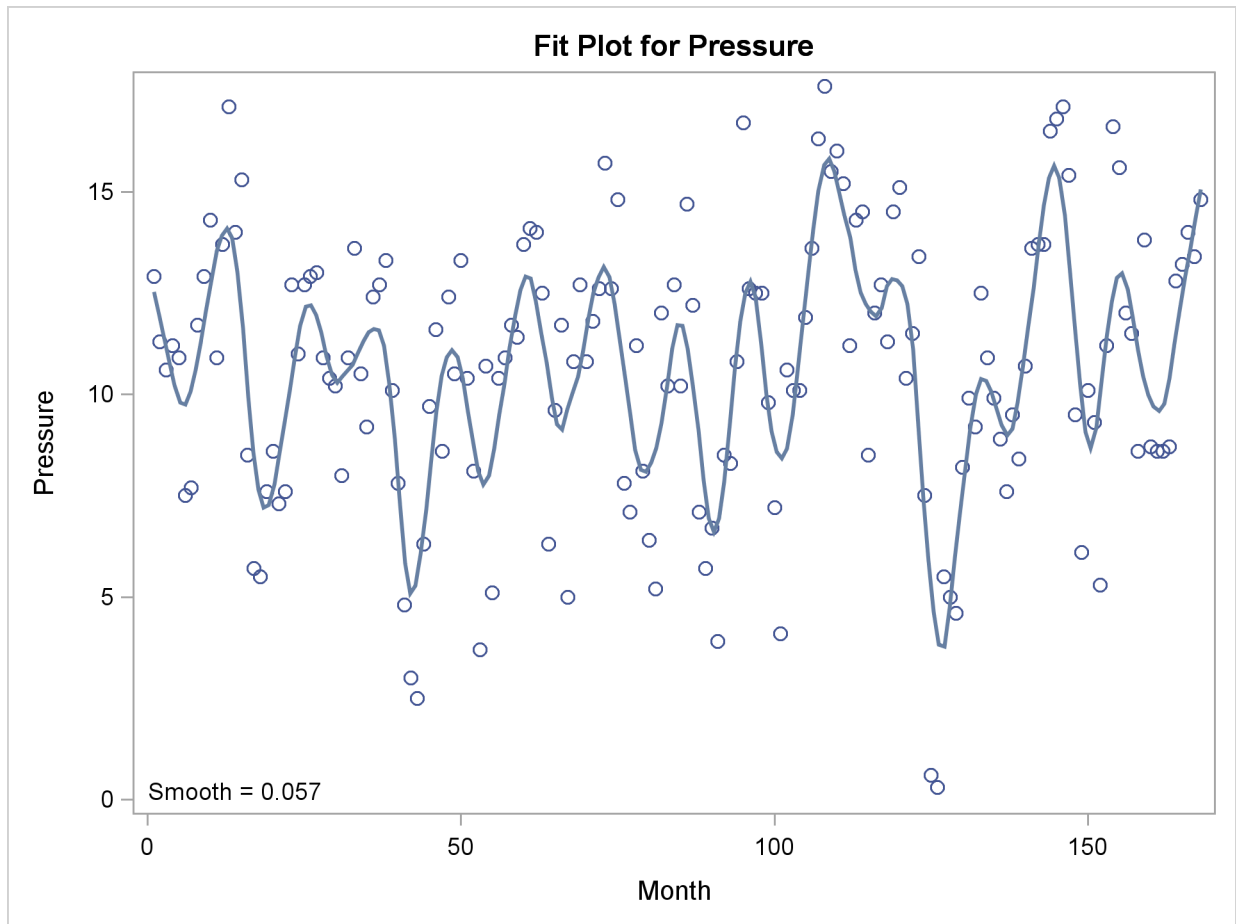
```
proc loess data=ENSO plots=residuals(smooth);
  model Pressure=Month/select=AICC(presearch);
run;

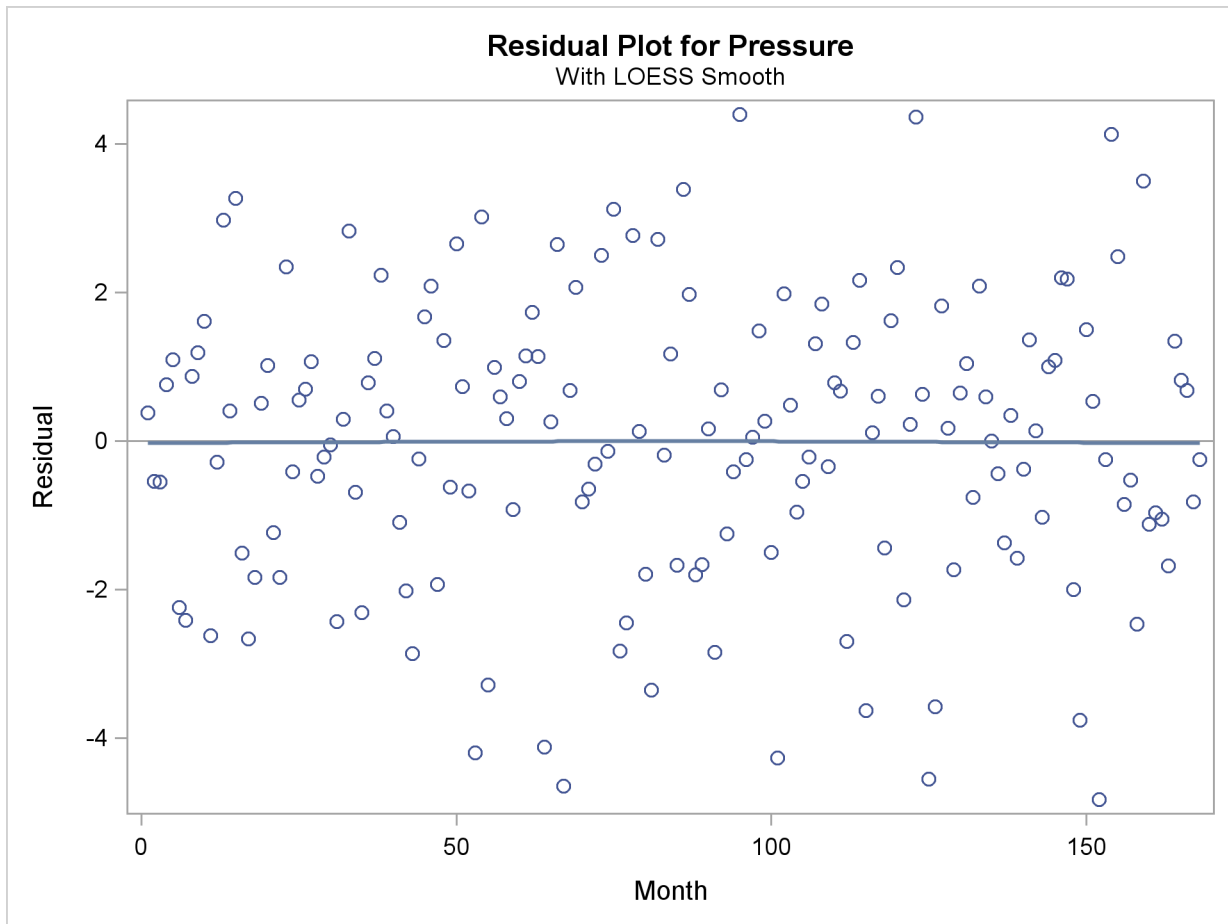
ods graphics off;
```

Output 50.4.7 Selected Smoothing Parameter Value When Presearch Is Specified

The LOESS Procedure	
Dependent Variable: Pressure	
Optimal Smoothing Criterion	
AICC	Smoothing Parameter
2.86660	0.05655

Output 50.4.7 shows that with the PRESEARCH suboption specified, PROC LOESS selects the smoothing parameter value that yields the global minimum of the AICC criterion. The fit obtained is shown in Output 50.4.8, and a plot of the residuals with a superimposed loess fit is shown in Output 50.4.9.

Output 50.4.8 Loess Fit Showing an Annual Cycle

Output 50.4.9 Residuals of the Selected Model

In contrast to the residual plot shown in [Output 50.4.4](#), the residuals plotted in [Output 50.4.9](#) do not exhibit any pattern, indicating that the corresponding loess fit has captured all the systematic variation in the data.

An interesting question is whether there is some phenomenon captured in the data that would explain the presence of the local minimum near 0.22 in the AICC curve. Note that there is some evidence of a cycle of about 42 months in the oversmoothed fit in [Output 50.4.3](#). You can see this cycle because the strong annual cycle in [Output 50.4.8](#) has been smoothed out. The physical phenomenon that accounts for the existence of this cycle has been identified as the periodic warming of the Pacific Ocean known as “El Niño.”

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