# BIOS6643 Longitudinal L12 Nonparametric regression

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Department of Biostatistics & Informatics

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## Topics for this lecture:

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- Spline modeling
- ▶ Associated reading: Sections 1 and 2 of the 'Nonparametric and flexible longitudinal regression' notes.

# Parametric, semiparametric and nonparametric regression: introduction and terminology

In modeling a mean function over time (or more generally for predictor x), a researcher may need more flexibility than what standard polynomials or transformations can offer. In this chapter we consider methods to accomplish such flexible fits.

Three classes of regression are parametric, semiparametric and nonparametric. These are discussed in some detail in this chapter (see course notes). Here, we focus on piecewise polynomial regression (parametric) and spline modeling (usually semiparametric or nonparametric).

See the course notes for more detail on nonparametric and semiparametric regression methods.

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# Piecewise polynomial regression and splines

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Summary

Piecewise polynomial regression offers a researcher a more flexible way to model the mean function over time (or more generally over some predictor, x), where the pieces are usually joined together so that the function is continuous but not necessary differentiable.

Spline models further require differentiability so that the entire function is smooth. Cubic terms are commonly used in spline models since they yield a flexible and smooth fit. Quadratic splines can also be used but are less common.

Although smoothness is intuitive in many cases, in certain cases it may be reasonable to allow the function to be continuous but not differentiable at one or more points, such as for a threshold model or when a treatment is applied during an experiment, resulting in a sharp change in the mean function. Such situations are discussed next.

# Piecewise linear regression

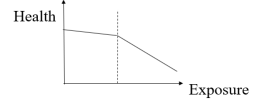
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Consider a health outcome that is modeled as a function of exposure to an environmental risk factor. There may be a negligible or slight dose-response relationship until the level of the risk factor reaches a certain point. Beyond that point, there may be a strong dose-response relationship between this risk factor and the health outcome. Such a model (sometimes called a threshold model) can be fit by joining polynomial functions together into one function. The simplest such function joins two simple linear functions together; the knot is where the two linear pieces join together.

As an introductory example, consider the threshold model described above. Let's assume there is some level of an environmental exposure variable that has the following relationship with a health outcome.



Say that exposure/health data are collected across subjects and the data is 'cross-sectional' in nature.

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Summary

In a GLM regression model, if we know where the knot occurs (say k), we can use the following regression function to fit the linear spline.

$$Y=\beta_0+\beta_1x+\beta_2 max(x-k,\ 0)+\epsilon$$

Note that the extra linear piece only 'kicks in' for  $x \ge k$ .

For x < k:

$$Y = \beta_0 + \beta_1 x + \epsilon$$

For  $x \ge k$ :

$$Y=\beta_0+\beta_1x+\beta_2(x-k)+\epsilon=(\beta_0-\beta_2k)+(\beta_1+\beta_2)x+\epsilon=\beta_0'+(\beta_1+\beta_2)x+\epsilon$$

Thus, the slope of x is  $\beta 1$  for x < k, and  $\beta 1 + \beta 2$  for  $x \ge k$ . Often our data will be longitudinal or clustered in nature, but we can fit splines in a linear mixed model in the same way.

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Illustration: Here is a simplified example of a real data set that I have worked with. Subjects that work in Beryllium metal plants have an increased risk of developing Beryllium sensitization (BeS), which can progress into Chronic Beryllium disease (CBD). We are interested in modeling changes in health over time, and specifically we want to see if there is a pronounced change when they progress from BeS to CBD. The health outcome measure here is y = AADO2R (Alveolar-arterial O2 tension difference at rest); a higher value indicates worse health.

## Description of variables:

- time can be thought of with units of years
- ▶ CBDX is the time when subjects progressed from BeS to CBD
- ▶ prog\_group = 0/1/2 for those that progress before/during/after the observation period
- stage = stage of illness, 0 for BeS, 1 for CBD
- ➤ Y=AADO2R, as described above

Here is one approach to modeling the data using linear splines:

$$Y_{ij} = \beta_0 + \beta_1 x_{ij} + \beta_2 max(x_{ij} - cbdx_i, \ 0) + pg_h + b_{0i} + \epsilon_{ij}$$

- $ightharpoonup \epsilon_{ij} \sim \mathcal{N}(0, \ \sigma_{\epsilon}^2),$
- $b_{i0} \sim \mathcal{N}(0, \ \sigma_{b_0}^2),$
- h = 1, 2, 3(progression group);
- i = 1, ..., n;
- $j = 1, ..., r_i$

Here,  $r_i = r = 4$  for all i; j = 0, 1, 2, 3, 4.

Above, I'm using x for TIME, pg for PROG\_GROUP. Note: since  $cbdx_i$  depends on i, subjects can have knots at different times. In the code below,  $time_star$  denotes the 'max' term.

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### SAS code

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### Outputs

#### Output: Model Information Type 3 Tests of Fixed Effects Dependent Variable Num Den Covariance Structure Variance Components Effect DE DE EValue Pr≥E Subject Effect time 1 18 4.37 0.0511 Estimation Method REML time\_star 1 18 9.60 0.0062 Residual Variance Method Profile prog group 2 18 2.07 0.1546 Fixed Effects SE Method Model-Based Degrees of Freedom Method Containment Covariance Parameter Estimates Class Level Information Cov Parm Subject Estimate Intercept id 0.7586 Class Levels Values Residual 2.5811 prog group 3 012 Fit Statistics Dimensions -2 Res Log Likelihood 90.0 AIC (smaller is better) 94.0 Covariance Parameters 2 AICC (smaller is better) 94.7 Columns in X BIC (smaller is better) 93.2 Columns in Z Per Subject 1 Subjects Max Obs Per Subject 5

#### Solution for Fixed Effects

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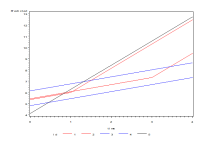
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The test for time\_star indicates that the progression from BeS to CBD causes significant changes to the health-time relationship (p=0.0062). With more data, we can try adding a few more parameters to the model to see if they help describe other patterns in the data.



In the graph, BeS subjects are forced to have the same linear trend and those with CBD are forced to have the same linear trend, but subjects can progress from one stage to the next at different times. Subject 5 progressed before the observation period, so they have the CBD trend; subjects 3 and 4 have the BeS trend since they progress after the observation period; subjects 1 and 2 progress during the observation period, one at time 1 and the other at time 3.

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Quadratic and cubic piecewise polynomial functions can also be fit to data

# Example 1: Potassium data.

These data were obtained via Ed Hess (a former graduate student) based on a consulting project he performed here at the university:

- Units of blood were sampled daily over the course of several weeks and assayed for Potassium level (exterior to the cells).
- ▶ The units were divided into four groups (see following page for plots of each group, given in the order listed as follows:
  - 1. control units that were not irradiated;
  - 2. units irradiated at study initiation;
  - 3. units irradiated at 7 days;
  - 4. units irradiated at 14 days.
- The motivation for this study was the idea that irradiation of bags can cause a release of free potassium which could result in cardiac arrest (such events had been observed during transfusions).
- ▶ The investigators wanted to characterize the rate of change in potassium level after irradiation for units of blood of different ages (i.e. that had been stored after donation for different lengths of time) to see if this had an impact on potassium release after irradiation.

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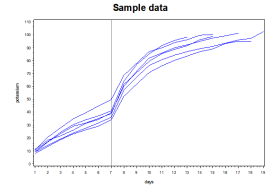
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Here, we consider units irradiated at 7 days. The data illustrate that there was an immediate effect of treatment on potassium levels. In the graph, potassium levels for 6 blood samples were each measured daily for up to 19 days. Responses within samples were joined to yield a spaghetti plot. In terms of spline modeling, it is clear that we want a knot at 7 days. Although the pattern appears to be that of two joined quadratic functions, we actually get a better model fit (lower AIC) including cubic terms in the spline model.

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### Model

Here is a possible model for the data:

$$Y_{ij} = \beta_0 + \beta_1 x_{ij} + \beta_2 x_{ij}^2 + \beta_3 x_{ij}^3 + \beta_4 s_{ij1}^1 + \beta_5 s_{ij2}^2 + \beta_6 s_{ij3}^3 + b_{1i} x_{ij} + \epsilon_{ij}$$

- indexes subject, j indexes observation,  $i=1,\;...,\;n;\;j=1,\;...,\;r_i$
- $Y_{ij} = j$ th weight observation for mouse i.
- $x_{ij} =$ day that jth observation was taken on mouse i.

```
data k;
set long.potassium;
if day416;
sl = (max(0,day-7))**1;
s2 = (max(0,day-7))**2;
s3 = (max(0,day-7))**3;
run;
run;
proc mixed data=k;
class sample;
model potassium= day day*day day*day*day
sl = 2 sample;
sl = 2 sample;
sl = 2 sample;
run;
proc mixed data=k;
class sample;
model potassium= day day*day day*day*day
sl = 2 sample;
run;
sl = (max(0,day-7))**3;
run;
proc mixed data=k;
class sample;
model potassium= day day*day day*day*day
sl = 2 sample;
run;
sl = 2 sample
```

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### Abbreviated output:

#### The Mixed Procedure

#### Covariance Parameter Estimates

#### Dimensions

Covariance Parameters 3
Columns in X 7
Columns in Z Per Subject 1
Subjects 6
Max Obs Per Subject 15
Number of Obs. Used 88

Cov Parm Subject Estimate day sample 0.1368 AR(1) sample 0.8678 Residual 8.5462

#### Fit Statistics

-2 Res Log Likelihood 344.8 AIC (smaller is better) 350.8 AICC (smaller is better) 351.1 BIC (smaller is better) 350.2

#### Solution for Fixed Effects

#### Standard

 Effect
 Estimate
 Error
 DF
 t Value
 Pr> |t|

 Intercept
 0.623
 1.7519
 76
 0.35
 0.7512

 day
 10.2237
 1.4932
 5
 6.85
 0.0010

 day\*day\* - 1.1590
 0.4193
 5
 2.76
 0.038

 day\*day\*day\*
 1.07170
 0.08440
 5
 2.08
 0.09

 1
 1.73723
 1.0657
 76
 1.60
 0.00
 6

 5
 0.167
 0.0387
 6
 -9.80
 <0.001</td>
 6

 5
 0.117
 0.0381
 76
 5
 0.00
 0.005

#### Solution for Random Effects

#### Std Err

Effect	san	nple	Estima	ate	Pre	d	DF	t Va	lue	Pr>	Įŧ
day	1	0.	09050	0.21	L55	76		0.42	0.	6756	
day	2	-0	.4602	0.21	55	76	-	2.14	0.	0359	
day	3	0	.3793	0.22	83	76	1	1.66	0.3	1007	
day	4	-0	.3115	0.21	55	76	-	1.45	0.	1523	
day	5	0.	02884	0.21	L55	76		0.13	0.	8939	
day	6	0	.2731	0.21	55	76	1	L.27	0.3	2088	

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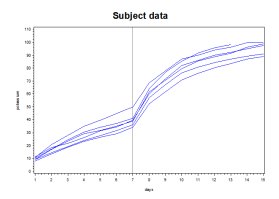
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The raw data (up through day 15 only) is shown below, followed by a graph of predicted values. For both, responses are joined by lines within subjects.



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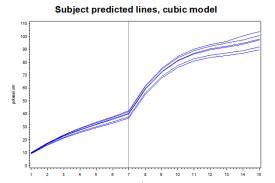
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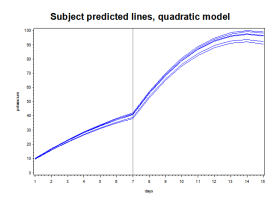
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If we drop the cubic terms (day and s3), we yield a much higher AIC of 416.6 for the quadratic model (shown below). Note that the predicted values start to bend back down at higher days, a pattern not evident in the data. Thus, the cubic model is superior both visually and quantitatively.



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For the cubic model, we can test for significance of at least one of the spline terms at day 7,  $H_0: \beta_4=\beta_5=\beta_6=0$ , using an F-test. This is accomplished by adding the following contrast statement in the PROC MIXED code:

It is not surprising that the test is very significant, given the previous output. This test is just confirming what we have already observed, that irradiation gives a strong boost to potassium levels.

We can compare the slope just before vs. just after irradiation by taking the derivatives of the fitted function at fixed days. Specifically, let f(x) = E[Y|x] for the mixed model, where x = days; let f'(x) denote the derivative of f(x). Note that

$$\begin{split} f'(x) &= \beta_1 + 2\beta_2 x + 3\beta_3 x_{ij}^2 &\quad for \; x < 7 \\ f'(x) &= \beta_1 + 2\beta_2 x + 3\beta_3 x_{ij}^2 + \beta_4 + 2\beta_5 (x - 7) + 3\beta_6 (x - 7)^2 &\quad for \; x \geq 7 \end{split}$$
 Using the fitted equations, we find that:

\$f'(6) = \$ \_\_\_\_\_ \$f'(8) = \$

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Cubic splines have a natural appeal due to their flexible fit, and although they are considered in the class of nonparametric regression modeling, the model can still often be expressed easily in parametric form. So far we have considered piecewise polynomial functions that may have a hard change point (i.e., continuous but not differentiable), but now we consider piecewise polynomial functions that are smooth. To obtain smoothness, lower-order terms are not included at the change points. Specifically, a piecewise polynomial cubic spline model has the form

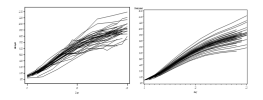
$$f(x) = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3 + \sum_{k=1}^p \beta_{k+3} s_k^3$$

where  $s_k=max(0,\ x-c_k)$  and  $c_k$  is the location of knot k with respect to the x-axis,  $k=1,\ ...,\ p.$ 

Unlike the previous examples, we only include the cubic terms  $(s_k^3)$ , but not the lower-order terms  $(s_k,\ s_k^2)$ , which forces differentiability across the entire function.

In some cases, we may want to include multiple knots in the spline model, and it may not be so clear where the knots should be. These are true particularly when we are more concerned about getting a flexible fit for the data - in the direction of nonparametric regression. To illustrate, consider the mouse growth data graphed below.

These data were obtained from Rob Weiss's (Dept. of Biostatistics, UCLA) web site. In the graph to the lower left, the weights of mice are measured over their first days of life; to the right are the predicted values based on the mixed model fit of the model described below.



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You may notice with the data that the quickest growth occurs around days 3 to 8, while the growth is not so steep shortly after birth, and then after day 10 or so. This suggests some type of cubic function may work for these data. Also, we may try modeling a random slope for time across subjects in order to account for the expanding variability between mice over time.

Using knots at days 3, 8 and 13 (where change points seem to be occurring), here is a possible model for the data:

$$Y_{ij} = \beta_0 + \beta_1 x_{ij} + \beta_2 x_{ij}^2 + \beta_3 x_{ij}^3 + \beta_4 s_{ij1}^3 + \beta_5 s_{ij2}^3 + \beta_6 s_{ij3}^3 + b_{1i} x_{ij} + \epsilon_{ij}$$

- lack i indexes subject, j indexes observation,  $i=1,\;...,\;n;\;j=1,\;...,\;r_i$
- ightharpoonup Yij = jth weight observation for mouse i
- $lacktriangledown x_{ij} = {\sf day} \ {\sf that} \ j{\sf th} \ {\sf observation} \ {\sf was} \ {\sf taken} \ {\sf on} \ {\sf mouse} \ i$
- $\begin{array}{l} \bullet \quad s_{ijk} = max(X_{ij} v_k, \ 0) \text{ where } k \text{ denotes knot, knots were fixed} \\ \text{at } v_1 = 3.3, \ v_2 = 8.3, \ v_3 = 13.3 \text{ days.} \\ \epsilon_{ij} \sim \mathcal{N}(0, \ \sigma_{\epsilon}^2); \ b_{i1} \sim \mathcal{N}(0, \ \sigma_{b_1}^2) \end{array}$

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In this case, the lower order spline terms were not included in the model, which is often not done in spline modeling with multiple knots.

For both of the examples in this subsection, note that we included a random term for time, but no random intercept. This worked since all experimental units had the same value, 0, at the start time.

But generally, I would caution against such an approach unless it makes sense.

Generally, I would warn against excluding the random intercept simply based on p-value, just as I would warn against dropping the fixed intercept term based on p-value.

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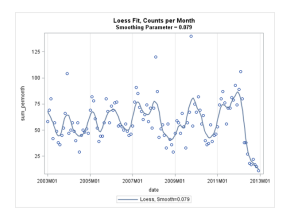
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The study took place in the San Luis Valley; hospital admission counts (for a medical facility in Alamosa) was compared with daily PM10 data (i.e., coarse particulate matter in the air) between 2003 and 2013.

Here, we consider larger number of knots to be able to get a 'nonparametric' fit to the data. I use nonparametric in quotes since really the spline data can still technically be expressed parametrically. However, most consider it a class of nonparametric regression.

When such spline variables are combined in models with predictors that are used in the standard way, then we typically call this a semi-parametric regression model. In the models discussed below, we use splines for time (so treat it 'nonparametrically'), and use standard variables for the pollutant, meteorological variables, month and day of week, and thus have a semi-parametric model.

# San Luis hospital admission counts and PM10.



Circles show monthly hospital counts, and a LOESS (kernel-type) nonparametric regression was used to get the fitted function. This was used for descriptive purposes only. LOESS regression is discussed in more detail in the next section of notes.

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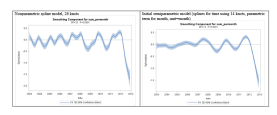
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Models below are only initial models that examine hospital counts as a function of time (left) and time and month (right). Here, canned procedures were used to obtain fits.



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Summary

The final model needs to include a flexible fit for time, account for serial correlation, and allow for testing for effects of interest (primarily the pollutant variable).

Once we define the variables associated with the splines, we actually have a parametric representation of the spline data and can include the variables in a standard parametric longitudinal model, like an LMM or GzLM with GEE. Since we have count data, we will use the latter to do all of this

Note that with these data, there is only one 'subject', the hospital at which we're measuring the daily admission counts. We will be able to fit the model as we have ample longitudinal data, although inference is limited to the population that uses this facility.

With a piecewise smooth cubic spline function, we include the (initial) intercept, linear, quadratic and cubic terms, and then k knots, where each knot has a related cubic 'spline' variable that kicks for  $\times$  greater than the knot. By including only the cubic terms associated with the knots, we keep the function smooth. (Also see the mouse data described previously.)

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Summary

The initial analyses suggested placing knots at roughly yearly intervals if we consider the more smoothed function. This would relate to 2-year cycles, which may be sufficient since the model will already have 'month' included, which should take care of trends within a year (e.g., a yearly cycle).

With about 10 years of data, we can place 9 equally spaced knots in the interior. This means there are 13 degrees of freedom including the initial intercept, linear, quadratic and cubic terms, and the spline terms associated with the 9 knots.

A 'b-spline' approach is essentially a transformation of the X matrix (for the spline variables) so that rows add up to 1. In this case, x variables act more like weights, and variables will have 0's for some elements, indicating that certain spline parameters are not used in predicting values if they are far away from point of interest. (See the SAS Appendix for a comparison of piecewise splines (or 'psplines') that we're familiar with, and basis-splines (or 'bsplines').

One advantage of b-splines is that the covariance between spline terms can be reduced, compared with p-splines. Another spline approach is to use natural b splines, which force the 2nd derivative of the function to be 0 at the beginning and ending knots.

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For practical purposes, I do not see much difference in models that use the pspline, bspline and nbspline approaches. While estimates and SE's of the spline terms may differ (including the intercept), those for the other terms in the model are either exactly the same or close to the same (they are exactly the same for pspline and bpline approaches, and close to the same for the natural bspline approach).

# Spline matrices can be obtained with software and code as follows:

- SAS: PROC TRANSREG
  - PSPLINE for piecewise spline
  - ► BSPLINE for basis spline
- R: SPLINE package
  - bs for basis splines
  - ns for natural splines

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There are many different types of spline approaches, only some of which are discussed here. Also, be careful with the terminology, it is not always consistent.

The SAS code demonstrates the program for total hospital count , using 3-day moving average for the pollutant, total hospital count. One run for each of bspline and pspline approaches is shown.

```
Smacro june (var.dist.splinetype.svar begin.svar end.polvarl.polvar2.
polvar3);
*use SAS to get variables;
proc transreg data=alldata;
 model identity(&var) = &aplinetype(cday
   / knots=0.350 to 3.150 by 0.350):
 output outwess splines improximations predicted; run;
data alldata2; merge alldata nbepline bepline sas splines; run;
proc genmod data=alldata2 /*descending*/; *where muni02<300;
class dayofweek month subject;
model &var = &polvar1 &polvar2 &polvar3
/* if pspline is used, vars will be cday 1-cday 12*/
/* if bepline is used, vars will be cday 0-cday 12*/
/* if R version of b-spline is used, vars are bs0-bs12*/
/* if R version of natural b-spline is used, vars are nbs0-nbs12*/
&svar begin - &svar end
dayofweek month temp pressure precip
/ dist=&dist corrb; output out=modfit predicted=p;
ods output GeeEmpPest=est1;
repeated subject=subject / type=/*ar(1)*/mdep(4) modelse;
*estimate 'line after knot' &polvar1 1 &polvar2 1; run;
%mend june:
% jume (n tot, poisson, bapline, cday 0, cday 12, logmuni02,,);
% june (n tot, poisson, pspline, cday 1, cday 12, logmuni02,,);
```

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## **BSPLINE** approach

```
BSPLINE approach
                                                                                      1.1469 0.3355 3.42 0.0006
                                                                        cday 1
                                                                                      1.2261 0.3300 3.72
                                                                        cday 2
The GENMOD Procedure
                                                                        cday 3
                                                                                      0.9632 0.3109 3.10
Model Information
                                                                        cday_4
                                                                                      1.5810 0.3069 5.15
          WORK.ALLDATA2
Data Set
                                                                        cday 5
                                                                                      0.8336 0.3101 2.69
                                                                                                          0.0072
Distribution Poisson
                                                                        cday 6
                                                                                      2.0013 0.3038 6.59
                                                                                                           <.0001
Link Function Law
                                                                                      0.3656 0.3201 1.14
                                                                                                          0.2533
                                                                        cday 7
Dependent Variable n tot
                                                                                      1.9698 0.2952 6.67
                                                                        cday 8
                                                                        cday 9
                                                                                      0.6050 0.3434 1.76
Number of Observations Read 3469
                                                                        cday 10
                                                                                      1.7775 0.2710 6.56
Number of Observations Used 3274
                                                                        cday_11
                                                                                      1 7970 0 4189 4 29
Missing Values
                                                                        cday 12
                                                                                      0.0000 0.0000
                                                                        dayofweek 1 0.2668 0.0444 6.01
Class Level Information
                                                                        dayofweek 2 0.1326 0.0468 2.83
Class Levels Values
                                                                        dayofweek 3 0.0487 0.0461 1.06
dayofweek 7 1234567
                                                                        dayofweek 4 0.0435 0.0459 0.95
month
                  123456789101112
                                                                        dayofweek 5 -0.0264 0.0482 -0.55 0.5838
           1
                                                                        dayofweek 6 0.0452 0.0467 0.97 0.3331
subject
                                                                        dayofweek 7 0.0000 0.0000 .
GEE Model Information
                                                                        month
                                                                                   1 0.0922 0.0746
                                                                                                                  0.2166
Correlation Structure 4-Dependent
                                                                        month
                                                                                    2 0.3782 0.0721
                                                                                                          5.24
                                                                                                                  < .0001
Subject Effect subject (1 levels)
                                                                                    3 0.2771 0.0784
                                                                                                          3.54
                                                                                                                  0.0004
Number of Clusters 1
                                                                                    4 0.0604 0.0897
                                                                                                          0.67
                                                                                                                  0.5005
Clusters With Missing Values 1
                                                                        month
                                                                                    5 -0.0048 0.0997 -0.05 0.9614
Correlation Matrix Dimension 3469
                                                                        month
                                                                                    6 -0.1366 0.1130 -1.21 0.2270
Maximum Cluster Size 3274
                                                                        month
                                                                                    7 -0.2933 0.1288 -2.28 0.0228
Minimum Cluster Size 3274
                                                                        month
                                                                                    8 -0.2159 0.1222 -1.77 0.0773
                                                                                    9 0.0376 0.1071
                                                                                                           0.35 0.7255
Algorithm converged.
                                                                        month 10 0.0599 0.0921 0.65 0.5150
GEE Fit Criteria
                                                                        month 11 -0.0044 0.0811 -0.05
OIC 3406 1776
                                                                        month 12 0.0000 0.0000
OlCu 3474 1776
                                                                        temo
                                                                                      0.0014 0.0019 0.75 0.4514
                                                                                      -0.3134 0.0977 -3.21 0.0013
                                                                        pressure
Analysis Of GEE Parameter Estimates using Model-Based SE Estimates
                                                                                      0.0044 0.1818 0.02 0.9808
                                                                        precip
(SE's were all 0 when using Empirical SE estimates)
                                                                        Scale 1.0276
```

The scale parameter for GEE estimation was computed

as the square root of the normalized Pearson's chi-square.

Estimate SE Z Pro ZI

6.3184 2.2385 2.82 0.0048

0.0202 0.0333 0.61 0.5437

1.1343 0.3309 3.43 0.0006

Intercept

cday 0

loemuni02

BIOS6643 Longitudinal

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regressoin

Case study

Bspline and Pspline

#### PSPLINE approach The GENMOD Procedure Model Information Data Set WORK,ALLDATA2 Distribution Poisson Link Function Los Dependent Variable Number of Observations Read 3469 Number of Observations Used 2274 Missing Values 195 Class Level Information Class Levels Values dayofweek7 1234567 month 12 123456789101112 subject 1 GEE Model Information Correlation Structure 4-Dependent Subject Effect subject (1 levels) Number of Clusters 1 Clusters With Missing Values1 Correlation Matrix Dimension Maximum Cluster Size Minimum Cluster Size Algorithm converged. GEE Fit Criteria 3406.1776 3474.1776

Analysis Of GEE Parameter Estimates using Model-Based SE Estimates

(SE's were all 0 when using Empirical SE estimates)

12 0.0000 0.0000 0.0014 0.0019 pressure +0.3134.0.0977+3.21.0.0013 0.0044 0.1818 0.02 0.9808 1.0276 . . .

0.1072 2.7133 0.04

-13.6431 7.2355 -1.89

41.8347 5.9230 7.06

dayofweek 5 -0.0264 0.0482 -0.55 0.5838

5 -0.0048 0.0997 -0.05 0.9614

6 -0.1366 0.1130 -1.21 0.2270

7 -0.2933 0.1288 -2.28 0.0228

8 -0.2159 0.1222 -1.77 0.0773

11 -0.0044 0.0811 -0.05 0.9567

-47.6282 6.1466 -7.75

-31.0938 5.7758 -5.38

cday 6 21,4826 5,9278 3,62

cday\_10 45.6791 6.5321 6.99

cday 11 -36.3350 8.1614 -4.45

dayofweek 1 0.2668 0.0444

dayofweek 2 0.1326 0.0468

devertises 3 0.0487 0.0461

dayofweek 4 0.0435 0.0459

dayofweek 6 0.0452 0.0467

dayofweek 7 0.0000 0.0000

month 1 0.0922 0.0746

3 0.2771 0.0784

9 0.0376 0.1071

10 0.0599 0.0921

cday\_12 -35.5942 23.6427 -1.51

0.6692 11.1875 0.06 0.9523 cday 3

0.0003

<.0001

< 0001

< 0001

<.0001

<.0001

0.1322

2.83 0.0046

1.06 0.2911

0.95 0.3441

0.97 0.3331

1.24 0.2166

5.24 < 0001

2.54 0.0004

0.67 0.5005

0.35 0.7255

0.65 0.5150

0.75 0.4514

< 0001 6.01

-2.1248 13.0953 -0.16 0.8711

7.0367 16.8737 0.42 0.6767

cday 1 cday 2

cday 4

cday 5

cday 7

cday 8

cday\_9

month 2 0.3782 0.0721

month 4 0.0604 0.0897

month

month

month

month

month

month

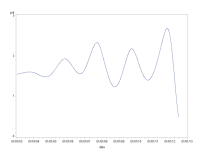
temp

```
Parameter Estimate SE Z Pr> Z
                                                                                  Note: The scale parameter for GEE estimation was computed
Intercept 7.4526 2.2297 3.34 0.0008
                                                                             as the square root of the normalized Pearson's chi-square.
Ioemuni02 0.0202 0.0333 0.61 0.5437
```

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**Bspline and Pspline** 



The graph above shows predicted counts based on the GzLM/GEE model fit. The fit represents month and day of week at reference values (December and Saturday, respectively). Otherwise, other covariates in the model besides those involving date (i.e., the spline terms) were set to their mean values. Predicted values are exactly the same, whether the PSPLINE or BSPLINE approaches are used.

The pollutant effect is not significant, but is going in the expected direction (positive). Some other models yielded p<0.05 for the pollutant variable, e.g., model with a binary pollutant variable based on a particular cut point.

# BIOS6643 Longitudinal

lonparametric egressoin

Piecewise polynomial regressoin

Cubic spline model

Bspline and Pspline

Pspline correlation between spline parameter estimates (intercept not included)

Bspline correlation between spline parameter estimates (intercept not included)

Nbspline correlation between spline parameter estimates (intercept not included)

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Nonparametric egressoin

regressoin

Bspline and Pspline

# Comparing piecewise polynomial and b-splines: bases and properties

Note: this section is taken from SAS Help Documentation, with some minor editing. An algorithm for generating the B-spline basis is given in de Boor (1978, pp. 134-135). B-splines are both a computationally accurate and efficient way of constructing a basis for piecewise polynomials; however, they are not the most natural method of describing splines. Consider an initial scaling vector  $\boldsymbol{x}=(1\ 2\ 3\ 4\ 5\ 6\ 7\ 8\ 9))^{\top}$  and a degree-three spline with interior knots at 3.5 and 6.5. The natural piecewise polynomial spline basis (X matrix for associated variables) is the left matrix, and the B-spline basis for the transformation is the right matrix.

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Bspline and Pspline

Piec	ewis	se Po	lynomi	al Splines	B-Split	S-Spline Basis					
/1	1	1	1	0	0 \	/1.000	0.000	0.000	0.000	0	0 \
/ 1	2	4	8	0	0 \	0.216	0.608	0.167	0.009	0	0 \
1	3	9	27	0	0	0.008	0.458	0.461	0.073	0	0
1	4	16	64	0.125	0	0	0.172	0.585	0.241	0.001	0
1	5	25	125	3.375	0	0	0.037	0.463	0.463	0.037	0
1	6	36	216	15.625	0	0	0.001	0.241	0.585	0.172	0
1	7	49	343	42.875	0.125	0	0	0.073	0.461	0.458	0.0008
\ 1	8	64	512	91.125	3.375	\ 0	0	0.009	0.167	0.608	0.216
\1	9	81	729	166.375	15.625	\ 0	0	0.000	0.000	0.000	1.000

The two matrices span the same column space. The numbers in the B-spline basis do not have a simple interpretation like the numbers in the natural piecewise polynomial basis. The B-spline basis has a diagonally banded structure and the band shifts one column to the right after every knot. The number of entries in each row that can potentially be nonzero is one greater than the degree. The elements within a row always sum to one. The B-spline basis is accurate because of the smallness of the numbers and the lack of extreme collinearity inherent in the piecewise polynomials.

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Piecewise polynomia

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### Bspline and Pspline

# Summary

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Spline and Pspline