BST 210 Lab: Week 2 Linear Regression: Simple and Multiple

Linear Regression

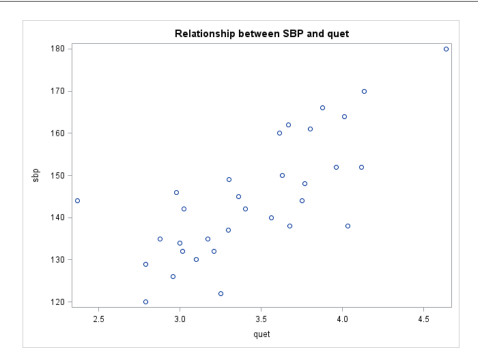
Linear regression is an incredibly useful tool that comes up often in statistical analyses. We use regression for two main reasons:

- To model or predict the values of an outcome of interest.
- To explore and quantify the relationship between an exposure(s) and an outcome.

For this lab, we'll be using the data found in lab2.csv (or in lab2.dat, for those using Stata) to look into simple and multiple regression. The data come from a sample of 32 individuals, and contains information on the systolic blood pressure (SBP), body size as measured by the quetelet index (quet), age (age), and smoking history (smk=0 for non-smoker and 1 for current and previous smoker) of each subject. Our primary outcome of interest will be SBP.

Simple Linear Regression: Example

Let's investigate the relationship between SBP and quet. Before fitting any sort of model, it's always a good idea to take an informal glance at your data. In SAS, we can do this by calling:



Since the relationship appears to be linear, we'll go ahead and fit a simple linear model. What form does this model take? And what assumptions are we making in fitting it?

The model that we fit has the form

$$E[SBP|quet] = \beta_0 + \beta_1 \cdot quet.$$

When fitting a linear regression model, we are assuming (1) a linear relationship between mean SBP and quet, (2) that all responses are independent, (3) that SBP is normally distributed about its mean, and (4) equal variance (homoskedasticity).

```
/* Running simple linear regression for the association between SBP and quet */
title ;
proc reg data=lab;
    model sbp = quet;
run;
```

				De	Mo		MODE		op				
			Nun	nbe	er of (Obser	vatio	ns Re	ad	32			
			Nun	nbe	er of (Obser	vatio	ns Us	ed	32			
					Anal	ysis o	f Vari	ance					
Sou	ırce		ı	DF		Sum o		Me Squ	ean are	F V	alue	Pr >	F
Mod	lel			1	3537.94588		88 35	3537.94588		3	6.75	<.00	01
Erro	r			30	2888.02287		87	96.26743					
Cor	rec	ted Tot	al	31	6425.96875		75						
		Root N	ISE			9.8	31160	R-So	quar	е 0.	5506		
		Depen	den	it M	ean	144.5	3125	Adj	R-S	q 0.	5356		
		Coeff \	/ar			6.7	78856						
					Para	mete	r Estir	nates					
	Variable DF					neter nate		dard Error	t V	alue	Pr >	• t	
	Intercept 1 70.5			7641	12.3	2187		5.73	<.00	001			
	qu	et	1		21.4	9167	3.5	4515		6.06	<.00	001	

What is the estimated slope, $\hat{\beta}_1$? How would you report that result to a collaborator?

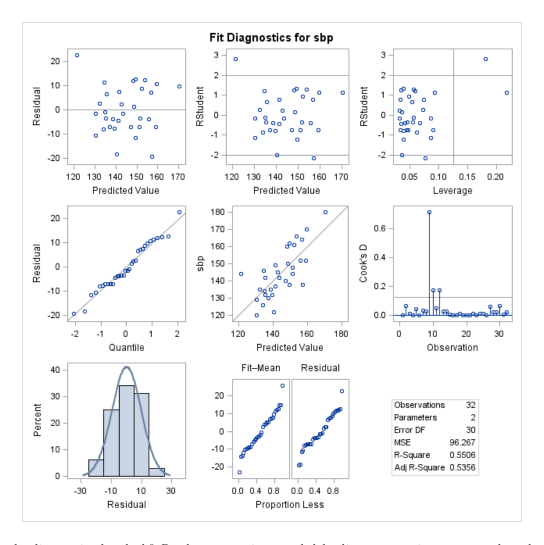
 $\hat{\beta}_1 = 21.492$. A one unit increase in quetelet index is associated with an estimated 21.492 mmHg increase in mean systolic blood pressure.

Regression Diagnostics

Once we've fit a linear regression line, we can use the **residuals**—defined as the difference between the true value Y and the fitted value $\hat{Y} = \hat{\beta}_0 + \hat{\beta}_1 \cdot X$ —to assess whether or not the assumptions we made are met.

- 1. **Linearity.** Plot the residuals against the predicted value, \hat{Y} . The residuals should be scattered randomly above and below the zero line, with no discernible pattern.
- 2. **Normality.** Look at a histogram of the residuals. If the normality assumption holds, this histogram should approximate a normal distribution—meaning it should be symmetric and centered around 0. Alternatively, create a quantile-quantile plot (qqplot): if the residuals are normally distributed, the points in the qqplot should fall on the y = x line.
- 3. **Equal Variance.** Again look at the plot of the residuals against the predicted value. The residuals should be just as "spread out" for large values of \hat{Y} as they are for small values of \hat{Y} .

SAS provides diagnostic plots as part of its standard proc reg output. Below are the diagnostic plots for our simple linear regression model:



How do the diagnostic plots look? Do the assumptions needed for linear regression appear to have been met?

The diagnostic plots suggest that the assumptions of linearity, normality, and equal variance have been met! As we can see from the plot of the residuals against the fitted values, our residuals are scattered randomly about the zero line, without any evidence of non-linearity or heteroskedasticity. The histogram and qqplot of the residuals indicate that they are approximately normal.

Assessing Model Fit

Even if our regression line satisfies these four key assumptions, that doesn't automatically guarantee that our model is good, or even useful. To that end, we can use the following statistics to assess how well our model fits the observed data, and how good a job it does at predicting/explaining our outcome of interest:

- R²: the proportion of the overall variability in our outcome that our model is able to explain
- Adjusted \mathbb{R}^2 : a modified version of \mathbb{R}^2 that takes into account the number of predictors in our model
- MSE: the estimated variance of the observed outcome about the fitted regression line—in other words, it estimates the amount variability in our outcome of interest that our model is unable to explain

In general, do we prefer larger R^2 /adjusted R^2 values or smaller R^2 /adjusted R^2 values?

We prefer models with larger R^2 and adjusted R^2 values, as this indicates that our model explains a larger proportion of the variability in our outcome.

What about for the MSE (and root MSE)?

We prefer models with **smaller** mean squared error and root mean squared error.

A quick caveat: these metrics only help us assess how well our linear model predicts our outcome of interest! If we're interested in unbiasedly estimating the association between our predictor and our outcome, there are no statistical tools to tell us whether we have controlled for all confounders and fit the correct model.

SAS reports these model fit statistics as part of its proc reg output:

Root MSE	9.81160	R-Square	0.5506
Dependent Mean	144.53125	Adj R-Sq	0.5356
Coeff Var	6.78856		

How would you explain this \mathbb{R}^2 value to a non-statistical collaborator? How would you explain the root MSE value?

The R^2 for the simple linear regression of systolic blood pressure on body mass (measured by quetelet index) is 0.551, meaning that body mass explains approximately 55% of the observed variability in systolic blood pressure in our sample.

The root MSE tells us to what degree our observed data points differ from the values that our fitted regression line predicts. In this particular instance, the standard deviation of the observed systolic blood pressure values about the fitted regression line is estimated to be 9.82.

Hypothesis Testing

Often times in data analysis, we're interested in determining whether two variables are significantly associated with one other. In the simple linear regression framework, we can formally test for association by looking at the slope:

$$H_0: \beta_1 = 0$$
 vs. $H_1: \beta_1 \neq 0$.

In SAS, the result of this hypothesis test is included in the proc reg output:

Parameter Estimates									
Variable	DF	Parameter Estimate		t Value	Pr > t				
Intercept	1	70.57641	12.32187	5.73	<.0001				
quet	1	21.49167	3.54515	6.06	<.0001				

What conclusion do we reach?

We reject the null hypothesis, and conclude that there is a significant association between quetelet index score and systolic blood pressure (p < 0.001).

Confidence Intervals

An alternative inferential method to hypothesis testing is the construction of a confidence interval. In SAS, we can create a confidence interval for the slope by adding a clb option to proc reg:

```
/* Running proc reg with confidence interval option (95\% is default) */
proc reg data=lab2;
    model sbp = quet / clb;
run;
```

	Parameter Estimates											
Variable	DF	Parameter Estimate		t Value Pr > t		95% Confidence Limits						
Intercept	1	70.57641	12.32187	5.73	<.0001	45.41180	95.74102					
quet	1	21.49167	3.54515	6.06	<.0001	14.25151	28.73182					

 $How\ would\ we\ interpret\ this\ confidence\ interval?$

With 95% confidence, a one unit increase in quetelet index is associated with an estimated increase in mean systolic blood pressure of between 14.25 mmHg and 28.73 mmHg.

Introduction to Multiple Regression

As we saw in class, a simple linear regression model often isn't sufficient to address our question of interest:

- We may believe that just one covariate (X) isn't enough to explain the variability in our outcome (Y)
- The presence of **confounders** or **effect modifiers** may obscure or change the true relationship between our covariate of interest (X) and the outcome (Y)

This is where multiple linear regression comes in handy! It allows us to extend simple linear regression to include other covariates of interest. As we saw in class, the model we fit when performing multiple linear regression has the form

$$E[Y|X_1,\ldots,X_p] = \beta_0 + \beta_1 X_1 + \ldots + \beta_p X_p.$$

How would you interpret the intercept term, β_0 ?

 β_0 is the mean value of Y when every covariate in the model (X_1, \ldots, X_p) is set to 0. Note that this won't always be a meaningful or sensible quantity!

How would you interpret the slope for X_1 , β_1 ?

 β_1 is the estimated change in the mean value of Y corresponding to a one unit increase in X_1 , holding all other covariates constant.

Note that we are still making the same assumptions of **linearity**, **independence**, **normality**, and **equal variances** as we did in the case of simple linear regression!

Multiple Linear Regression & Confounding

A **confounder** is a variable that is associated with both our outcome of interest and the exposure, but that is not a consequence of the exposure. To put that in more concrete terms, let's consider the relationship between smoking status (smoker/non-smoker) and lung cancer:

Would having a genetic predisposition for cancer be a confounder of this relationship? Why or why not?

No, as having a genetic predisposition for lung cancer would not be associated with the exposure—smoking status.

Would number of packs smoked per day be a confounder of this relationship? Why or why not?

No, as the number of packs smoked each day is a consequence of smoking status.

How about age? Why or why not?

Yes, age is a potential confounder of the relationship, as older individuals may be more likely to smoke and also more likely to develop lung cancer.

The presence of confounding causes **bias** in our estimate of the association between the exposure and the outcome—this is why we need to adjust for confounders in linear regression! However, not all confounders actually lead to a large or meaningful amount of bias. So as a rule of thumb, we'll consider a variable to be

a meaningful confounder of the relationship between X and Y if adjusting for it changes the estimated slope by 10% or more.

Let's return to the same dataset we used for simple linear regression: lab2.csv. Suppose we're still interested in quantifying the relationship between quet and SBP. What are some factors that might confound this relationship?

Some potential confounders might include whether or not an individual has a high sodium diet, number of hours of exercise per week, gender, and age.

We'll use multiple regression to adjust for one potential confounder: age! What will our new model look like?

$$E[SBP|age, quet] = \beta_0 + \beta_1 \cdot quet + \beta_2 \cdot age$$

```
/* Running multiple linear regression with both quet and age */
proc reg data=lab;
    model sbp = quet age;
run;
```

	Parameter Estimates											
Variable	DF	Parameter Estimate		t Value	Pr > t							
Intercept	1	55.32344	12.53475	4.41	0.0001							
quet	1	9.75073	5.40246	1.80	0.0815							
age	1	1.04516	0.38606	2.71	0.0113							

What is the estimate for the association between quet and SBP? How would you report this to a collaborater?

After adjusting for age, the new estimate is $\hat{\beta}_1 = 9.75$.

Among individuals of the same age, a one unit increase in quet score is associated with an estimated increase of 9.75 mmHg in mean SBP.

Does age appear to be a meaningful confounder? Why or why not?

Yes! The estimated slope for quet changed from 21.492 in the unadjusted model to 9.7507 in the adjusted model. This is a change of well over 10%—in fact the decrease is over 50%—so we can conclude that age is a meaningful confounder.

Note: the fact that the p-value associated with age is significant is not evidence that age is a meaningful confounder—this just tells us that there is a significant association between age and SBP, even after adjusting for quetelet index.

Suppose that age was a meaningful confounder of the relationship between SBP and quet score, but that its p-value was not significant. Would we still want to include it in the model?

This depends in part on what the purpose of our model is. If we're just looking to build the model that best predicts SBP—without actually caring about the relationship between any one predictor and SBP—then we might want to exclude age. BUT if we're interested in assessing the relationship between quet and SBP, then we need to include in the model any meaningful confounders of that relationship, no matter whether or not the confounders themselves have significant slopes.

Multiple Linear Regression & Effect Modification

We consider a random variable to be an **effect modifier** if the magnitude of the association between our predictor and our outcome varies across its different levels.

Let's (once again!) go back to the blood pressure data set and consider the relationship between quet score and SBP. SBP will still be our response variable, but this time let's also look at the dichotomous variable, smk. There are three different models that we can fit:

1. The **Coincident Model**, which assumes that smokers and nonsmokers have the same intercept and slope:

$$E[\mathtt{SBP}|\mathtt{quet}] = \beta_0 + \beta_1 \cdot \mathtt{quet}$$

2. The **Parallel Model**, which assumes that smokers and nonsmokers have different intercepts, but the same slope:

$$E[SBP|quet, smk] = \beta_0 + \beta_1 \cdot quet + \beta_2 \cdot smk$$

3. The **Full Model**, which assumes that smokers and nonsmokers have different intercepts and different slopes:

$$E[\mathtt{SBP}|\mathtt{quet},\,\mathtt{smk}] = \beta_0 + \beta_1 \cdot \mathtt{quet} + \beta_2 \cdot \mathtt{smk} + \beta_3 \cdot \mathtt{quet} \cdot \mathtt{smk}$$

This third model, which looks at the interaction between smk and quet, is the one that will help us assess whether effect modification is occurring.

Let's assume we fit model (3). What is the regression model for individuals with smk = 0?

$$E[SBP|quet, smk] = \beta_0 + \beta_1 \cdot quet$$

What is the regression model for individuals with smk = 1?

$$E[SBP|quet, smk] = (\beta_0 + \beta_2) + (\beta_1 + \beta_3) \cdot quet$$

```
/* In SAS, we first need to create a new variable for the interaction (quet)*(smk) */
data lab2;
    set lab;
    quet_smk = quet*smk;
run;

proc reg data=lab2;
    model sbp = quet smk quet_smk;
run;
```

	Parameter Estimates										
Variable	DF	Parameter Estimate		t Value	Pr > t						
Intercept	1	49.31177	19.97234	2.47	0.0199						
quet	1	26.30282	5.70349	4.61	<.0001						
smk	1	29.94356	24.16355	1.24	0.2256						
quet_smk	1	-6.18478	6.93171	-0.89	0.3799						

Based on the output above, does smk appear to be an effect modifier of the relationship between quet and SBP?

The interaction term between smk and quet has a non-significant slope (p = 0.38). Given that we fail to reject the null hypothesis that $\beta_3 = 0$, it seems that smoking status does not modify the relationship between quet and SBP. So smk does not appear to be an effect modifier.

Based on the model fit above, what is the estimated association between quet and SBP among smk = 0? How would you report this to a collaborator?

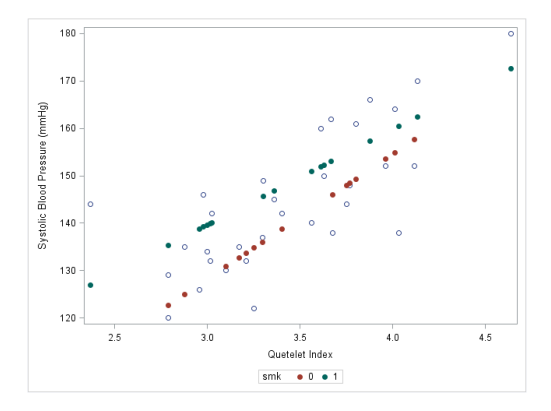
The estimated effect is $\hat{\beta}_1 = 26.303$. Among non-smokers, a one unit change in quetelet score is associated with an estimated 26.303 mmHg increase in mean systolic blood pressure.

What is the estimated association between quet and SBP among smk = 1? How would you report this to a collaborator?

The estimated effect among smokers is $\hat{\beta}_1 + \hat{\beta}_3 = 26.303 - 6.185 = 20.118$. Among current or previous smokers, a one unit change in quetelet score is associated with an estimated 20.118 mmHg increase in mean systolic blood pressure.

If we want to plot the fitted lines for smokers and non-smokers—and overlay these plots on top of a scatterplot of the observed data—we can do so using the following commands in SAS:

```
/* Plotting Fitted Regression Lines Over a Scatterplot */
/* We first need to output a dataset that includes our fitted/predicted values */
proc reg data=lab2;
       model sbp = quet smk quet_smk;
        output out=pred
                p=yhat;
run;
/* We can then overlay these fitted values on top of our scatter plot */
proc sgplot data=pred;
        xaxis label = "Quetelet Index";
        yaxis label = "Systolic Blood Pressure (mmHg)";
        scatter x=quet y=sbp / ;
        scatter x=quet y=yhat / GROUP=smk markerattrs = (symbol = circlefilled);
        * GROUP specified color-coding by group, and markerattrs changes the specific
        * attributes of the points, here making them filled dots;
run;
```



Additional Topic: Confidence Intervals for Predicted Values

In this course, we will mostly be concerned with estimating—and constructing confidence intervals for—regression coefficients. But in many clinical settings, we aren't just interested in building regression models to estimate associations—we also want to use our models to *predict* health outcomes for particular patient subpopulations, or even for particular patients.

To make our predictions more meaningful, we would like to have some way of quantifying our certainty (or uncertainty) in these predictions. This is the motivation for constructing intervals for predicted subpopulation means and predicted individual patient outcomes!

We'll once again use the blood pressure data set, and will return to the regression model we fit in the first part of lab:

$$E[\text{sbp}|\text{quet}] = 70.576 + 21.493 \cdot \text{quet}.$$

Confidence Intervals for a Predicted Subpopulation Mean

Suppose that a quetelet index of 3.5 has some sort of clinical significance, and that we want to estimate the average systolic blood pressure among all patients with that value. Based on the estimated regression coefficients, the subpopulation individuals with a quetelet index of 3.5 has an average systolic blood pressure of

$$70.576 + 21.493 \cdot 3.5 \approx 145.8$$
 mmHg.

We can construct a confidence interval for this subpopulation mean in SAS using the clm option in proc reg:

```
/* To generate a prediction for an unobserved value , we first need to add it as an
   additional observation in our dataset */
data lab3;
        set lab2 end=eof;
        output;
        if eof then do;
                person=33;
                quet=3.5;
                sbp=.;
                output;
        end:
run;
/* Running proc reg with confidence interval for mean value option */
proc reg data=lab3;
        model sbp = quet / clm;
run;
```

The REG Procedure
Model: MODEL1
Dependent Variable: sbp

	Output Statistics													
Obs	Dependent Variable	Predicted Value	Std Error Mean Predict	95% CL Mean		Residual								
1	135	132.3864	2.6499	126.9747	137.7982	2.6136								
2	122	140.4458	1.8608	136.6456	144.2460	-18.4458								
3	130	137.2006	2.1144	132.8824	141.5187	-7.2006								
4	148	151.5570	2.0860	147.2968	155.8172	-3.5570								
5	146	134.6001	2.3858	129.7276	139.4725	11.3999								

. . .

31	152	155.7264	2.5335	150.5523	160.9005	-3.7264
32	164	156.7580	2.6601	151.3254	162.1906	7.2420
33	-	145.7972	1.7470	142.2294	149.3651	

So with 95% confidence, the mean systolic blood pressure for the population of individuals with a quetelet index of 3.5 is between 142.23 and 149.37 mmHg.

Prediction Intervals for a Specific Individual

Sometimes, we may be interested in predicting the systolic blood pressure of a specific patient with a quetelet index of 3.5. In this case, our best estimate for that particular individual's systolic blood pressure value is still the population average SBP for all patients with a quetelet index of 3.5,

$$70.576 + 21.493 \cdot 3.5 \approx 145.8 \text{ mmHg}$$

since we have no reason to believe that this particular patient differs systematically from the population.

But we're a lot less certain of this prediction, since we know there can be a lot of biological variability from person to person. So our confidence interval (also called a prediction interval) for this individual prediction will be different from those above.

SAS, R, and Stata all allow us to construct a prediction interval for the predicted systolic blood pressure of an individual with quet=3.5. In SAS, that code looks like:

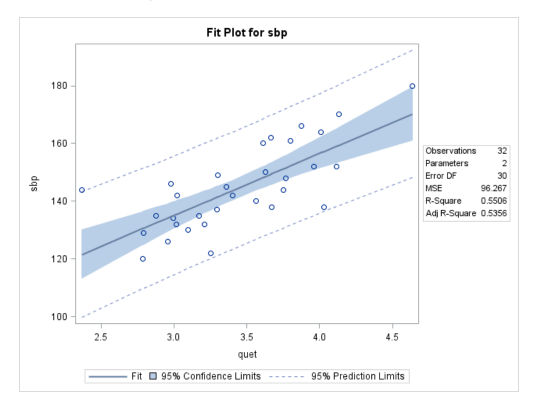
	The REG Procedure Model: MODEL1 Dependent Variable: sbp											
		Ou	tput Stati	stics								
Obs	Std Error Dependent Predicted Mean Obs Variable Value Predict 95% CL Predict Residual											
1	135	132.3864	2.6499	111.6306	153.1423	2.6136						
2	122	140.4458	1.8608	120.0507	160.8409	-18.4458						
3	130	137.2006	2.1144	116.7026	157.6985	-7.2006						
4	148	151.5570	2.0860	131.0712	172.0428	-3.5570						
5	146	134.6001	2.3858	113.9782	155.2219	11.3999						

. . .

31	152	155.7264	2.5335	135.0312	176.4216	-3.7264
32	164	156.7580	2.6601	135.9967	177.5193	7.2420
33		145.7972	1.7470	125.4441	166.1504	

We are 95% confident that the systolic blood pressure for an individual with a quetelet index of 3.5 is between 125.44 and 166.15 mmHg.

proc reg in SAS also automatically returns the following plot, which overlays both the confidence intervals and prediction intervals for each predicted SBP value:



Note that in the above figure, the confidence bands for the predicted subpopulation means are always narrower than the prediction bands for the predicted individual means. This makes intuitive sense: there is often much greater variability in the characteristics of a particular individual than there is variability in the average characteristics of a subpopulation. For example, we can be fairly certain that the mean height of women in this class is between 5 ft 2 in and 5 ft 6 in, but we're not as sure that the next woman who walks through the door will be between 5 ft 2 in and 5 ft 6 in tall.

It makes mathematical sense as well: our uncertainty about an individual prediction is made up of uncertainty about the mean PLUS the variability of individual observations about that mean. When we estimate the subpopulation mean, we're estimating

$$E[Y|X] = \beta_0 + \beta_1 \cdot X,$$

and so our uncertainty in the subpopulation means comes from our uncertainty in the estimation of β_0 and β_1 . But when we predict outcome values for an individual, we are instead concerned with

$$Y = \beta_0 + \beta_1 \cdot X + \epsilon$$
$$= E[Y|X] + \epsilon,$$

where the error term ϵ captures the additional variability of the individual observations about the subpopulation mean.