POLS/CSSS 503: Advanced Quantitative Political Methodology

Linear Regression: Specification and Fitting

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Specification

\[ Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \ldots + \varepsilon \]

(Note: for the moment, we use \( X_1 \) rather than \( x_1 \) to refer to the first covariate)

Model specification refers to the modeler’s choice of \( X_1, X_2, \) etc.: which \( X \)'s we include in our model, which we exclude, & how we transform them.

We need to get this right for substantive & statistical reasons.

In observational research, a large % of criticism regard specification.

The goal of most experimental or quasi-experimental work, is avoiding specification altogether, but even then not always possible.

First, we’ll talk about what different specifications imply substantively.

Later, we’ll talk about how to choose a specification: fitting the model.
Outline of topics

Omitted variable bias & Specification

Transforming covariates

Transforming response variables

Diagnosing heteroskedasticity & misspecification

Goodness of Fit tests
Omitted variable bias

Let’s imagine we know that the true model for some data $Y$ is

$$Y_i = \beta_{0}^{\text{true}} + \beta_{1}^{\text{true}} X_i + \beta_{2}^{\text{true}} Z_i + \varepsilon_i^{\text{true}}$$

Data generating process (DGP): the model generating the population we sample (usually a fiction)

Which quantitative methods will recover the truth about the model’s parameters?

Unpacking the question:

• “recover the truth”: yield an efficient, unbiased estimate; one that is true on average (and close to the truth) over many samples of $X, Y, Z$

• “methods”: includes the method of estimation; e.g. least squares

• “model”: includes the choice of specification; e.g., which controls, transformations, and interactions to include on the RHS
Omitted variable bias

Because $Y_i$ was constructed by adding together

$$\beta_0^{\text{true}} + \beta_1^{\text{true}} X_i + \beta_2^{\text{true}} Z_i + \text{a Normal disturbance},$$

estimation by least squares will be not only be unbiased, but also the best unbiased method (most efficient)

But it will only be unbiased if we choose the right specification

If we estimate

$$Y_i = \beta_0^* + \beta_1^* X_i + \varepsilon_i^*$$

We won’t get any estimate for $\beta_2$ (because we assumed it was zero by omitting it)

Moreover, it will often be the case that $\hat{\beta}_1^*$ is a biased estimate of $\beta_1$
Omitted variable bias

The source of this bias can be shown formally

We estimated this

\[ Y_i = \beta_0^* + \beta_1^* X_i + \varepsilon_i^* \]

leaving out \( Z_i \). Suppose we ran an auxiliary regression of \( Z_i \) on \( X_i \):

\[ Z_i = \gamma_0 + \gamma_1 X_i + \nu_i \]

We could substitute this back into the true model,

\[ Y_i = \beta_0^{\text{true}} + \beta_1^{\text{true}} X_i + \beta_2^{\text{true}} Z_i + \varepsilon_i^{\text{true}} \]

\[ Y_i = \beta_0^{\text{true}} + \beta_1^{\text{true}} X_i + \beta_2^{\text{true}} (\gamma_0 + \gamma_1 X_i + \nu_i) + \varepsilon_i^{\text{true}} \]

\[ Y_i = (\beta_0^{\text{true}} + \beta_2^{\text{true}} \gamma_0) + (\beta_1^{\text{true}} + \beta_2^{\text{true}} \gamma_1) X_i + (\varepsilon_i^{\text{true}} + \beta_2^{\text{true}} \nu_i) \]
Omitted variable bias

The parenthetical terms below are what we recover when we run LS omitting $Z_i$

\[
Y_i = (\beta_0^{\text{true}} + \beta_2^{\text{true}} \gamma_0) + (\beta_1^{\text{true}} + \beta_2^{\text{true}} \gamma_1) X_i + (\varepsilon_i^{\text{true}} + \beta_2^{\text{true}} \nu_i)
\]

\[
Y_i = \beta_0^* + \beta_1^* X_i + \varepsilon_i^*
\]

The estimate we get of $\beta_1$ is:

\[
E(\hat{\beta}_1) = \beta_1^{\text{true}} + \beta_2^{\text{true}} \gamma_1
\]

\[
= \beta_1^{\text{true}} + \beta_2^{\text{true}} \left( \frac{\sum_i (X_i - \bar{X})(Z_i - \bar{Z})}{\sum_i(X_i - \bar{X})^2} \right)
\]

Which is unbiased only if $\beta_2^{\text{true}} = 0$ or $\text{corr}(X_i, Z_i) = 0$. 
Omitted variable bias

Thus there are only two conditions under which we can safely omit variable $Z$ from our model:

\[ \beta \text{ is really zero; } Z \text{ has no effect on } Y \text{ in the “true” model} \]

or

The correlation of $Z$ with the included $X$’s is zero

we should include any $Z$ that is correlated with both $Y$ and some included $X$

(A major exception – post-treatment variables – discussed below)

This is why the complaint “You should have controlled for . . .” carries so much weight in criticizing empirical research.

Specification arguably the major concern in most observational research (along with selection & endogeneity)
Omitted variable bias

Does this mean we should include the kitchen sink in the regression?

Is there a penalty to including irrelevant variables?

Yes, but it is smaller. Lose efficiency in two ways:

- Lost degrees of freedom
- Lost variance in relevant covariates after conditioning on irrelevant ones

So is the kitchen sink safer?

Kevin Clarke: you only solve OVB with “right” specification. If there are countervailing biases, adding a subset of omitted variables could make things worse

My view: in observational research, show robustness to specification choice. Try to break your findings, then report how easily they are broken
Omitted variable bias

Is there any other reason why we should omit variables?

Generally, you should omit “post-treatment” variables or intermediate steps in a (presumed) causal process.

If you want to model $X \rightarrow Y$, and one path between $X$ and $Y$ is $X \rightarrow W \rightarrow Y$, then omit $W$ or risk $W$ masking the underlying effect of $X$.

Consider this hypothesis:

*Post-college education makes people more likely to vote Democratic*

Suppose we test this by regressing vote choice in 2004 on years of education.

A critic objects that we should have controlled for party ID, which is correlated with both our response and covariate. Is he right?

The same logic says to include vote intention 10 minutes prior to voting!

We can validly omit variables $W$ which lie along the causal chain from affect $X$ to $Y$’s if we want $\beta X$ to absorb the impact of $X$ through $W$. 
Omitted variable bias

Is there any other reason why we should omit variables?

Generally, you should omit “post-treatment” variables or intermediate steps in a (presumed) causal process.

If you want to model $X \rightarrow Y$, and one path between $X$ and $Y$ is $X \rightarrow W \rightarrow Y$, then omit $W$ or risk $W$ masking the underlying effect of $X$.

Note an implication:

To get a good estimate of the relationship between $X$ and $Y$ from a regression, we may need to craft the rest of the regression around that causal story.

But now suppose we do want to understand the relationship between short-run intentions and voting.

We’d need to specify the model differently.

So testing multiple hypotheses might require multiple specifications, even in the same paper.
What makes linear regression “linear”?  
Any thoughts?  

As long as this remains a valid statement:  

\[ Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \ldots + \varepsilon \]

the regression is linear  

But we can make any algebraic substitutions we want  

To see this, replace the big $X$’s of the above equation with functions of small $x$’s  

This is a trick from high school algebra:  
I’m just renaming complex algebraic expressions with simple names
Transformations of covariates

Suppose \( X_1 = x_1^2 \) and \( X_2 = x_1 \). Then, by algebraic substitution, the DGP is:

\[
Y = \beta_0 + \beta_1 x_1^2 + \beta_2 x_1 + \ldots + \varepsilon
\]

Sample output (note we’re leaving out \( x_1 \) to make a point; normally we need it, too):

Call:
\[\text{lm(formula = y} \sim \text{I(x^2))}\]

Residuals:

<table>
<thead>
<tr>
<th></th>
<th>Min</th>
<th>1Q</th>
<th>Median</th>
<th>3Q</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>-4.3876</td>
<td>-0.1713</td>
<td>0.2031</td>
<td>0.3971</td>
<td>0.7495</td>
</tr>
</tbody>
</table>

Coefficients:

|                  | Estimate  | Std. Error | t value  | Pr(>|t|) |
|------------------|-----------|------------|----------|----------|
| (Intercept)      | 0.9008332 | 0.0310244  | 29.036   | <2e-16   *** |
| I(x^2)           | -0.0006920 | 0.0007175  | -0.965   | 0.335    |
The same regression; two views

Left is the regression in the original, untransformed scale of $x$

Right is the regression as R sees it, in the transformed scale $x^2 = X$

Linear regression is always linear in the transformed covariate

It may be very curvilinear in the scale we care about
Transformations of covariates

Suppose $X_1 = x_1^3$, $X_2 = x_1^2$, $X_3 = x_1$.

This is a cubic polynomial specification:

$$Y = \beta_0 + \beta_1 x_1^3 + \beta_2 x_1^2 + \beta_3 x_1 + \ldots + \varepsilon$$

Or even add a quartic term, $X_4 = x_1^4$. Then,

$$Y = \beta_0 + \beta_1 x_1^3 + \beta_2 x_1^2 + \beta_3 x_1 + \beta_4 x_1^4 + \ldots + \varepsilon$$
Sample output for a cubic (3rd order) polynomial:

Call:
\( \text{lm(formula = y \sim I(x^3) + I(x^2) + I(x))} \)

Residuals:

\[
\begin{array}{cccccc}
\text{Min} & \text{1Q} & \text{Median} & \text{3Q} & \text{Max} \\
-4.3835 & -0.1706 & 0.2006 & 0.3983 & 0.7536 \\
\end{array}
\]

Coefficients:

\[
\begin{array}{cccccc}
\text{Estimate} & \text{Std. Error} & \text{t value} & \text{Pr(>|t|)} \\
(\text{Intercept}) & 0.8840553 & 0.0830528 & 10.644 & <2e-16 *** \\
I(x^3) & 0.0002337 & 0.0011134 & 0.210 & 0.834 \\
I(x^2) & -0.0044455 & 0.0168552 & -0.264 & 0.792 \\
I(x) & 0.0166821 & 0.0721171 & 0.231 & 0.817 \\
\end{array}
\]

None of the coefficients are significant, but they collectively explain a lot of variance. (How is this possible?)

With polynomials (and with other interactions) \( t \)-tests of individual parameters are less interesting than CIs around \( \hat{Y} \)
These are two different regressions.

Left is a cubic (3rd order) polynomial specification. It has 2 bends

Right is a quartic (4th order) polynomial. It has 3 bends

Each polynomial order we add puts another bend in the line
If we include \( n \) terms, there will be a bend for every observation

Called “curve-fitting”: a perfect (\& perfectly useless) model

Always have a theoretical reason to include polynomial terms

Seldom is more than a quadratic justified by theory

If you include polynomial terms, you need to interpret the result graphically
Warnings about polynomial fits:

- High order polynomials will always fit the sample well, but seldom fit the population well (curve-fitting)

- Extrapolation from polynomial or interactive specifications is dangerous. These functional forms behave wildly outside the known data
Polynomial overfitting experiment: generate 10 obs from the “true” model:

\[ Y = x + \varepsilon, \quad \varepsilon \sim N(0, 3) \]

and fit these data using different polynomials of \( x \).

We will show the fit of the model to the original data on the left
We will also draw new “out of sample data” from the same true model:

$$Y_{OOS} = x_{OOS} + \varepsilon_{OOS}, \quad \varepsilon \sim N(0, 3)$$

and use the model as fitted on the original dataset to predict out of sample cases.

We will show the fit of the old model to the out-of-sample data on the right.
Above is the fit from a quadratic specification of $x$, ie, we estimated:

$$Y = \hat{\beta}_1 x + \hat{\beta}_2 x^2 + \hat{\epsilon}$$

Note that we omitted the constant for didactic reasons.
Above is the fit from a cubic specification of $x$; that is, we estimated:

$$Y = \hat{\beta}_1 x + \hat{\beta}_2 x^2 + \hat{\beta}_3 x^3 + \hat{\epsilon}$$

How many polynomials can we add and still find $\hat{\beta}$?

What will happen to the fit in and out of sample as we add polynomials?
Above is the fit from a quartic specification of $x$; that is, we estimated:

$$Y = \hat{\beta}_1 x + \hat{\beta}_2 x^2 + \hat{\beta}_3 x^3 + \hat{\beta}_4 x^4 + \hat{\epsilon}$$

On the left, we see the model finds a curious non-linearity, by which low and high $x$ suppress $y$, but middle values of $x$ increase $y$. Do you trust this finding?
In small samples, outliers can easily create the illusion of complex curves relating $x$ and $y$.

We need a lot of data to discern if such curves are more than spurious.

(And so we probably need a strong theory, too, to justify the data collection)
What happens as we approach a tenth-order polynomial?
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What happens as we approach a tenth-order polynomial?
When the number of parameters in the model equals the number of observations, least squares is able to fit a line through every datapoint.

This means the fit will be “perfect”: no error.

And out of sample, it will be completely useless, and worse than guessing that $y$ simply equals its sample mean in every case.


Two lessons:

1. Beware curve-fitting:

Significance tests may suggest adding polynomials, but that isn’t enough to justify their inclusion
2. Beware good fits in-sample unless they fit well out of sample, too

Every goodness of fit measure has an out-of-sample counterpart

Out of sample goodness of fit is much more important than in sample
More Transformations of covariates

Suppose \( X_1 = \log(x_1) \). Then,

\[
Y = \beta_0 + \beta_1 \log(x_1) + \beta_2 X_2 + \ldots + \varepsilon
\]
These are the same regression

Left is on the original, untransformed scale

Right is on the transformed, log scale

Log transformations for effects that diminish in per unit potency as $x$ increases
Transformations of covariates

We could keep going, combining transformations and interactions to get very nonlinear models.

Suppose \( X_1 = x_1 \times x_2 \) and \( X_2 = x_2 \). Then,

\[
Y = \beta_0 + \beta_1 x_1 \times x_2 + \beta_2 x_2 + \ldots + \varepsilon
\]

Suppose \( X_1 = x_1 \times \log(x_2) \times \sqrt{x_3} \) and \( X_2 = x_2/(x_1 + x_2^2) \). Then,

\[
Y = \beta_0 + \beta_1 (x_1 \times \log(x_2) \times \sqrt{x_3}) + \beta_2 x_2/(x_1 + x_2^2) + \ldots + \varepsilon
\]

Without strong theoretical support, these models will be silly and produce unreliable results with little predictive power.
Transformations of the response variable

We could even replace $Y$

This is a good idea if you think $Y$ is not a linear function of the regressors, but $g(y)$ is a linear function of them.

Usually, this is the case for counts, e.g., or money.

Raising a government budget from $1$ million to $10$ million may be “just as hard” as raising it from $10$ to $100$ million.

If $X$’s affect the order of magnitude of $Y$, you should log $Y$. 
Transformations of the response variable

If $Y = \log(y)$, then,

$$\log(y) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \ldots + \varepsilon$$

Now all $X$’s have diminishing effect

In fact, level changes in $X$ yield percentage changes in $Y$

If you log both $X$ and $Y$, then % changes in $X$ cause % changes in $Y$
Transformations of the response variable

What if $Y$ is bounded but continuous?

Suppose it ranges between 0 and 1 (but doesn’t include these values)?

Then we need to “stretch it out” to range from $-\infty$ to $\infty$

The logit transformation does this: $Y = \log(y/(1 - y))$.

$$\log \left( \frac{y}{1 - y} \right) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \ldots + \varepsilon$$

This model doesn’t work if the original data include 0s or 1s

It is not “the logit model”, which is a non-linear model of 0s and 1s only

It is a linear regression with a logit transformed response variable
Note the curve is S-shaped.

We could rescale it to work for any bounds, not just \((0, 1)\)

That is, just transform \(y\) to \(y^* = \frac{y-a}{b-a}\),
then run the regression of \(\text{logit}(y^*)\) on your covariates

This works generally, with the caveat that for any bounds \((a, b)\),
none of the data can be exactly \(a\) or \(b\)
Transformations of the response variable

All of the above models are examples of linear regression

They are all linear in the parameters

They can all be estimated by least squares

What specifications can’t we estimate?

We can’t use LS to estimate models that are non-linear in the parameters

Example of a model that is non-linear in the parameters:

\[ Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \beta_1 \beta_2 X_3 + \ldots + \varepsilon \]

No amount of algebra can turn the above into a linear model

There are advanced methods to deal with this, e.g., non-linear least squares

Doesn’t come up that often, because so many specifications are linear in the parameters

Lots of flexibility hidden in linear regression
So just what is linear regression again?

We have expanded linear regression to encompass any model for unbounded continuous functions $f(\cdot)$ and arbitrary functions $g_k(\cdot)$:

$$y_i = \beta_0 + \beta_1 x_{1i} + \ldots + \varepsilon_i$$

$$f(y_i) = \beta_0 + \beta_1 g_1(x_{1i}) + \ldots + \varepsilon_i$$

$$f(y_i) = \beta_0 + \sum_{k=1}^{K} \beta_k g_k(x_{ki}) + \ldots + \varepsilon_i$$

“Linearity” in linear regression just refers to the fact that the effect of a one unit change in $g(x_k)$ on $f(y)$ is $\beta_k$.

But the relationship between $x_k$ and $y$ themselves could be very non-linear, as a result of $f(\cdot)$ and $g_k(\cdot)$. 
Interpreting models with transformed covariates or responses

Various methods:

- For logged response, $\hat{\beta}$ is the % change in $Y$ for a level change in $X$
- For logged covariate, $\hat{\beta}$ is the level change in $Y$ for a % changes in $X$
- For both sides logged, $\hat{\beta}$ is the % change in $Y$ for % changes in $X$, known as the elasticity of $Y$ with respect to $X$
- For polynomial coefficients, make a plot of $\hat{Y}$ as $X$ varies: don’t try to interpret each coefficient separately!
- For interaction like $X \times Z$, make one or more plots of $\hat{Y}$ under different combinations of $X$ and $Z$: don’t try to interpret the coefficients on $X$, $Z$, and $X \times Z$ separately!
Interpreting models with transformed covariates or responses

So how do we get $\hat{Y}$ for these models?

Could do it by hand fairly easily

But what if we want confidence intervals around $\hat{Y}$ too?

Use predict().

Key is setting newdata input correctly

If lm() did the transformations and interactions

  ie, you used the I() command in the model formula

  and you used * or : to make the interactions

Then predict() will construct interactions and polynomials from their base terms as needed

If not, you need to give predict properly constructed interactions and polynomials
What we’ve learned about linear regression so far

Now we know how to:

1. Specify a regression model
2. Estimate that model
3. Interpret our findings

But how do we know if our findings are any good?
That we used the right specification?
That our model explained the data well or poorly?

Need to learn one more skill:

4. Select models with good fit
Running example: Life Expectancy

We will consider a simple linear regression today.

The data are 138 countries observed in 1985.

The response variable is average life expectancy in years.

We consider four covariates:
- `gdpcap85`: Per capita GDP in 1985, thousands of international dollars.
- `school`: Average years of education.
- `civlib5`: Low = 1 to high = 7 scale of civil liberties.
- `wartime`: Percent of recent history spent at war.

Data are from Barro & Lee.

Example contrived for pedagogical use, not a serious effort at demography.
Running example: Life Expectancy

```r
lm(formula = lifeexp ~ gdpcap85 + school + civlib5 + wartime,
    na.action = "na.omit")
```

Coefficients:

|                | Estimate  | Std. Error | t value | Pr(>|t|) |
|----------------|-----------|------------|---------|----------|
| (Intercept)    | 52.9671697 | 3.0379356  | 17.435  | <2e-16   *** |
| gdpcap85       | 0.0003216  | 0.0002551  | 1.260   | 0.211    |
| school         | 2.3866392  | 0.4102266  | 5.818   | 9e-08    *** |
| civlib5        | -0.7522863 | 0.4861266  | -1.548  | 0.125    |
| wartime        | 1.0936902  | 5.5354286  | 0.198   | 0.844    |

---

Signif. codes:  0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Residual standard error: 5.43 on 90 degrees of freedom
Multiple R-Squared: 0.7509,    Adjusted R-squared: 0.7399
F-statistic: 67.84 on 4 and 90 DF,  p-value: < 2.2e-16
Running example: Life Expectancy

Note that initially, only education has a significant effect

Surprising?

Perhaps we have badly fit the model?
Two questions in model fitting

Question 1: Are the relationships I think I see really there?

Question 2: Do they explain much of what is happening?

Somewhat overlapping questions

Often helpful to tackle them together

How? There are many ways to answer these questions

You will need to use several to avoid being fooled
Are the relationships I think I see really there?

Look to:

- standard errors & associated tests
- patterns in the residuals
- warning signs for outliers
Standard errors and associated tests

You already know the $t$-test

But what if you want to test more than one parameter at a time?

$F$-test: joint tests of whether some or all the parameters of the model are 0

Recall the analysis of variance breakdown of what a regression “explains”:

Total Sum of Squares: $\sum_{i=1}^{n} (y_i - \bar{y})^2 = (y - \bar{y})' (y - \bar{y})$

Residual Sum of Squares: $\sum_{i=1}^{n} (y_i - \hat{y}_i)^2 = \varepsilon' \varepsilon$

Regression Sum of Squares: $\sum_{i=1}^{n} (\hat{y}_i - \bar{y})^2 = (X\beta - \bar{y})' (X\beta - \bar{y})$

Recall that $R^2 = \frac{\text{Regression Sum of Squares}}{\text{Total Sum of Squares}}$ or the proportion of variance explained by the model
Standard errors and associated tests

Suppose our “null” hypothesis is $\beta_1 = \beta_2 = 0$.

This hypothesis imposes restrictions on $q = 2$ parameters of a model with $k$ parameters total estimated on $n$ observations.

The $F$-test associated with this null is:

$$F_0 = \frac{\text{Regression Sum of Squares}/q}{\text{Residual Sum of Squares}/(n - k - 1)}$$

$$= \frac{n - k - 1}{q} \times \frac{R^2_{\text{unrestricted}} - R^2_{\text{restricted}}}{1 - R^2_{\text{unrestricted}}}$$

Under the null, $F_0$ is follows an $F$ distribution with $q, n - k - 1$ degrees of freedom.

We can reject this null even when we can’t be sure which of $\beta_1$ or $\beta_2$ is non-zero.

We don’t know which is explaining variance in $Y$, but we know that together, they explained enough variance to reject the null.
Standard errors and associated tests

A special case of the $F$-test asks whether every $\beta$ is zero simultaneously.

In other words, do any of our covariates explain anything?

Reported as part of the regression summary:

F-statistic: 67.84 on 4 and 90 DF, p-value: < 2.2e-16

In this case, we could choose to reject the null that every coefficient is 0

This is the same as saying some parameter(s) is/are not zero (but which?)

Seldom interesting. Virtually always rejected

Useful in rare cases where we want a regression to find null results; e.g., if there shouldn’t be a relationship between $y$ and a set of covariates
Examining residuals

We noted earlier that residuals from a least squares regression should have no pattern. In particular, the variance in $\varepsilon_i$ should not be a function of $Y$ or $X$. 
Examining residuals for heteroskedasticity

This is what we hope to see: white noise

No relationship between $X$ (or $Y$) and the mean or variance of the residuals
Examining residuals for heteroskedasticity

Often, however, we find a megaphone shape, or some other change in variance

So why do we care? So what do we do about it?
Weighted least squares

Besides biasing SEs, heteroskedasticity makes LS inefficient. Why?

Observations with higher variance in errors contain less information

Observations with lower variance tend to be very close to the LS line

We’ll get better (more efficient) estimates if we give more weight to the latter

Needed:

• A measure of $\text{Var}(\varepsilon_i) = \sigma_i^2$ for each observation

• A method for weighting observations in least squares estimation

• In particular, we want to minimize the weighted sum of squares

$$\sum_{i=1}^{n} w_i \varepsilon_i^2 = \varepsilon' W \varepsilon$$

where $W$ is a diagonal matrix with weights $w_i$ on the diagonal
Weighted least squares

The solution is simple: just add weight terms to the estimator

$$\hat{\beta}_{WLS} = (X'WX)^{-1}X'Wy$$

where $W$ is a diagonal matrix with weights $w_i$ on the diagonal

What are the weights? They are (proportional to) the standard error for each $y_i$

Ideally, the weights are defined such that

$$\varepsilon_i \sim \mathcal{N}(0, \sigma^2_i)$$
$$\sigma^2_i = 1/w_i^2$$

→ the larger the weight $w_i$, the smaller the variance of $\varepsilon_i$, the more information in $i$

In R, just add the argument `lm(..., weights)"
Weighted least squares

Hypothetical example: Modeling school spending

- Small districts have more variable budgets

  A single shock (a charitable gift; a new disabled student) can shift the mean

- Big districts have stable budgets

  Shocks average out over many schools

- In a model of district spending per pupil, small districts should get lower weight

- Signal to noise ratio higher in large districts
Adjusting standard errors for heteroskedasticity

Suppose we diagnose, or suspect, heteroskedasticity, but have no weights.
We cannot use WLS, and thus rely on the less efficient LS estimates.
But we can try to get standard errors approaching the WLS se’s.

Recall the standard errors from LS are the square roots of the diagonal elements of

\[ \hat{V}(\hat{\beta}) = \hat{\sigma}^2 (X'X)^{-1} \]

So if \( \sigma^2 \) varies by \( i \), these will be wrong.

A “heteroskedasticity robust” formula for the Var-Cov matrix is:

\[ \hat{V}(\hat{\beta}) = (X'X)^{-1} (\sum_i \hat{e}_i x_i' x_i) (X'X)^{-1} \]
Adjusting standard errors for heteroskedasticity

\[ \hat{V}(\hat{\beta}) = (X'X)^{-1}\left(\sum_i \hat{\varepsilon}_i x_i'x_i\right)(X'X)^{-1} \]

SE’s calculated from this equation are known by many names:

- White standard errors
- robust standard errors
- sandwich standard errors
- heteroskedasticity consistent standard errors

Always a second best approach.

But very popular (used in most poli sci papers that use LS)
Adjusting standard errors for heteroskedasticity

To get robust standard errors in R:

```r
res <- lm(lifeexp~gdpcap85+school+civlib5+wartime,
          na.action="na.omit")

vc <- hccm(res) # library car must be loaded

se.corrected <- sqrt(diag(vc))
    # se's are the square roots of diagonal elements of
    # the variance-covariance matrix of the parameters
```

# Note: you’ll need to calculate your own t-tests & CIs

If the robust SEs are very similar to the usual SEs, don’t bother reporting them

(Catch-22: If robust se’s are very different, LS is inefficient anyway, and you should look for weights or ways to model the heteroskedasticity)
The life expectancy example

Is heteroskedasticity a problem in our example?

Let’s look at the residuals
Life expectancy residuals against fitted Y

Whoa! That's not (just) heteroskedasticity!

The residuals appear correlated with the fit itself. (Why is that bad?)
Other patterns in the residuals

Non-constant variance isn’t the only pattern we could see in residuals.

What if the residuals don’t appear to have mean 0 for all levels of $X$?

E.g., what if they are rising in $X$, or show curvature?

This is evidence of *misspecification*.

In particular, curvature in the residuals indicates missing transformations of $X$.

Let’s plot our life expectancy residuals against each covariate.
Life expectancy residuals against Civil Liberties

Not much sign of a pattern in the mean or variance here.
This looks like heteroskedasticity, but is it?

Perhaps there just are too few high war cases to see the full spread
Clearly something amiss. What does this plot show?

How might we fix it?
Remember that GDP wasn’t even significant.

But there *does* seem to be a relation between GDP and the residual.

So why didn’t this show up in our model? What did we miss?
Other patterns in the residuals

A pattern in the mean of the residuals is evidence of specification bias

How do we eliminate the bias?

Include transformations of the regressor!

E.g., if you see a curve, try adding $X^2$ or $\log X$

Let’s respecify the Life expectancy regression to include logs of GDP and school
Respecified model

```
lm(formula = lifeexp ~ I(log(gdpcap85)) + I(log(school)) + civlib5 + wartime, na.action = "na.omit")
```

Coefficients:

|                      | Estimate | Std. Error | t value | Pr(>|t|) |
|----------------------|----------|------------|---------|----------|
| (Intercept)          | 13.1893  | 5.4439     | 2.423   | 0.0174 * |
| I(log(gdpcap85))     | 5.3730   | 0.6867     | 7.825   | 9.35e-12 *** |
| I(log(school))       | 6.0431   | 0.9040     | 6.685   | 1.88e-09 *** |
| civlib5              | -0.1843  | 0.3084     | -0.598  | 0.5517   |
| wartime              | -0.2174  | 3.6894     | -0.059  | 0.9531   |

---

Signif. codes:  0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Residual standard error: 3.627 on 90 degrees of freedom
Multiple R-Squared: 0.8889, Adjusted R-squared: 0.8839
F-statistic: 180 on 4 and 90 DF, p-value: < 2.2e-16
Respecified model: residuals

The se’s indicate a much better fit regarding GDP and schools

But was this the “right” transformation?

Let’s run through the residual diagnostics again, to see if the new specification eliminates the patterns we found
Life expectancy residuals against Civil Liberties

![Graph showing life expectancy residuals against Civil Liberties]
Life expectancy residuals against Wartime

Diagram showing the relationship between life expectancy residuals and wartime. The x-axis represents wartime, ranging from 0.0 to 0.4, while the y-axis represents the residuals, ranging from -10 to 10. The data points are scattered across the plot, indicating a trend in the residuals with respect to wartime.
Life expectancy residuals against School

res.$\$resid

school

-10 −5 0 5 10

2 4 6 8 10 12
Life expectancy residuals against GDP

Much better overall, though perhaps a bit of underfitting at the high end

Let’s look at residuals against $\hat{Y}$
A pattern still shows up at the highest fitted values

Possibly a result of limits to medical technology

Further testing of transformations of $X$, and perhaps $Y$, might help
Is heteroskedasticity messing up the standard errors?

There may still be some (mild) heteroskedasticity here

We can check the robust SEs:

```r
vc <- hccm(res.l)
se.corrected <- sqrt(diag(vc))
```

<table>
<thead>
<tr>
<th></th>
<th>Est</th>
<th>SE</th>
<th>Robust SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>13.19</td>
<td>5.44</td>
<td>5.38</td>
</tr>
<tr>
<td>log(gdpcap85)</td>
<td>5.37</td>
<td>0.69</td>
<td>0.67</td>
</tr>
<tr>
<td>log(school)</td>
<td>6.04</td>
<td>0.90</td>
<td>0.84</td>
</tr>
<tr>
<td>civlib5</td>
<td>-0.18</td>
<td>0.31</td>
<td>0.29</td>
</tr>
<tr>
<td>wartime</td>
<td>-0.22</td>
<td>3.69</td>
<td>3.78</td>
</tr>
</tbody>
</table>

In this case, it makes little difference which SEs we use

Caveat: robust SEs are not perfect fixes, and may be wrong in small samples
What about transforming the response?

How do you know when to transform $Y$, instead of $X$?

Theory: if you think all covariates should have diminishing effects as $Y$ rises

Evidence: Histogram of residuals shows non-Normality amenable to a transformative “fix”
Other patterns in the residuals

Histograms of the residuals are also useful

The histogram should look approximately Normal

If it appears skewed, e.g., to the right, try logging $Y$
Does my model explain much of what is happening?

The “coefficient of determination”, $R^2$

The standard error of the regression, $\sigma^2$

Out of sample tests

Cross-validation
Much ado about $R^2$

Many regression tables in journals report $R^2$

$R^2$ is the proportion of variance in $Y$ explained by the regression model

$$R^2 = \frac{\text{Explained sum of squares}}{\text{Total sum of squares}} = 1 - \frac{\text{Residual sum of squares}}{\text{Total sum of squares}} = \frac{\sum \hat{y}^2}{\sum y^2} = 1 - \frac{\sum \hat{\epsilon}^2}{\sum y^2}$$

$R^2$ is bounded by 0 and 1

$R^2 = 0$: model has no explanatory power

$R^2 = 1$: model perfectly predicts every obs without error

Conventional wisdom: higher $R^2$ is “better”
King (“How not to lie with statistics”) on what $R^2$ does & doesn’t say

- $R^2$ shows the *proportion of variance explained* by the covariates, compared to a model with no covariates

- $R^2$ indirectly reports the scatter around the regression line ($\hat{\sigma}^2$ directly reports the amount of scatter)

- $R^2$'s comparable only for models with same observations and response variable
  (In the life expectancy example, it’s good when adding substantively interesting variables raises $R^2$ noticably)

- Maximizing $R^2$ is usually perverse. More covariates always raise $R^2$.

- The most useful model is seldom the one with the highest $R^2$.
  (Uninteresting high $R^2$ models:
   $Y$ regressed on itself. Vote choice regressed on vote intention.)

- $R^2$ is not an “estimate”. (It can't be significant or non-significant.)

- $R^2$ seldom of substantive interest, unlike $\hat{\sigma}^2$, or the se of $\hat{\beta}$. 
The standard error of the regression

\( \hat{\sigma} \) is at least as useful to report as \( R^2 \)

It tells us how much the fitted values, \( \hat{y} \), miss the true \( y \) on average

Thus it is on the scale of \( y \). Easy to work into a substantive conclusion

How is this a measure of goodness of fit?

As \( \sigma \) goes down, we make better predictions

\( \hat{y} \) gets closer to the true \( y \)
### Fitting the life expectancy model

<table>
<thead>
<tr>
<th>Covariates</th>
<th>Goodness of Fit</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$R^2$</td>
<td>ser ($\sigma$)</td>
</tr>
<tr>
<td>GDP, Sch, CL, War</td>
<td>0.75</td>
<td>5.43</td>
</tr>
<tr>
<td>log(GDP), log(Sch), CL, War</td>
<td>0.88</td>
<td>3.63</td>
</tr>
</tbody>
</table>

$R^2$ is a useful statistic here (same data & response)

What does it mean?

What does the $\sigma$ mean?

Is the second model a better fit?
More persuasive tests of fit

The most persuasive evidence of model fit is successful prediction

Suppose we estimate a model of presidential approval ratings, using data over 1950—1995.

We could ask: How well does the model fit the 1950—1995 data?

Could also ask: How well does the model predict 1996—2005 data?

*Out-of-sample* tests are powerful checks against spurious findings

If we intend the model to apply to the out of sample data, this is a better test

The usual caveat applies: the best fitting model is not necessarily the most interesting

But if our model fits as well to the out of sample data as the in sample, much greater confidence we have found something real

(Both samples could have outliers, heteroskedasticity, or specification error, so we still have to check for these)
Out of sample goodness of fit

How to do an out-of-sample test:

1. Fit the model on the training sample, \( \{X_{\text{training}}, y_{\text{training}}\} \), obtain \( \hat{\beta}_{\text{training}} \)

2. Calculate the fitted \( \hat{y}_{\text{test}} \) for the test sample, \( \{X_{\text{test}}, y_{\text{test}}\} \)

3. Compare fit for training sample to our ability to predict test sample.

For example, compare \( \hat{\sigma}_{\text{training}} \), the standard error of the residuals from the training regression, to the standard error of the “residuals” from the test predictions:

\[
\text{std dev}(y_{\text{test}} - X_{\text{test}}\hat{\beta}_{\text{training}})
\]

(This is the average prediction error)

A very good predictive model will have about the same error out of sample as in sample. (Rare to find)
Cross-validation

What if we’re short on data, and can’t afford to sequester half of it for an out of sample GoF test? Not a problem: just use cross-validation

1. Select all but $1/k$th of the data, $\{y_{training}, X_{training}\}$. Leave-one-out cross validation ($k = n$) is especially good.

2. Regress $y_{training}$ on $X_{training}$ to obtain $\beta_{training}$

3. Use $\hat{\beta}_{training}$ and $X_{test}$ to form predictions $\hat{y}_{test}$

4. Compare $\hat{y}_{test}$ with the true $y_{test}$, by calculating the average prediction error:

$$\text{std dev}(y_{test} - X_{test}\hat{\beta}_{training})$$

5. Repeat steps 1–4 $k$ times, and average the squared average prediction errors. The square root is the $k$-fold cross-validation predictive error.

The best predictive model, according to cross-validation, will minimize this average predictive error.
Cross-validation

R can do cross-validation for you:

```r
library(boot)
data <- data.frame(lifeexp, gdpcap85, school, civlib5, wartime,
        row.names=country)
data <- na.omit(data)

res.glm1 <- glm(lifeexp~gdpcap85+school+civlib5+wartime,
        na.action="na.omit")
    # Note the use of glm rather than lm

cv.err <- cv.glm(data, res.glm1, k=5)
    # This runs the cross-validation

cv.err$delta
        # Report the CV prediction error

# And repeat for each model
```
### Cross-validation

<table>
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<tr>
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<th>Goodness of Fit</th>
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<tr>
<td>GDP, Sch, CL, War</td>
<td>0.75</td>
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<td>32.3</td>
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<tr>
<td>log(GDP), log(Sch), CL, War</td>
<td>0.88</td>
<td>3.63</td>
<td>13.8</td>
</tr>
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The CV prediction error is much higher than $\hat{\sigma}$

Not a surprise

When we try to predict “new” data from a model, we usually find it works less well

Getting the specification “right” makes a huge difference for correct prediction

Missing life expectancy by 32 years is horrific.
Missing by 14 is just okay.

Thinking we would only miss by 3.63 was overconfident.
Cross-validation

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<tr>
<td>log(GDP), log(Sch), log(CL), War, War$^2$</td>
<td>0.90</td>
<td>3.48</td>
<td>12.1</td>
</tr>
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Could we do better? Try to eliminate remaining curvature in the residuals?

Try a model with curves in each covariate

(Why did I use the square of War?)

A slight improvement. Shows up in CV, so it might even be real.

But getting GDP and School right was crucial