

Multiple Linear Regression: Model Checking and Diagnostics

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Today's Lecture

- Model checking and diagnostics
- Variable transformations

Model selection vs. model checking

Assume $y|\mathbf{x} = f(\mathbf{x}) + \epsilon$

- model checking asks whether the ϵ match the assumed form, whether there are systematic and diagnosable (and fixable!) deviations from assumed model structure.
- model selection (coming soon!) focuses on how you construct $f(\cdot)$;

Model checking: possible challenges

Two major areas of concern

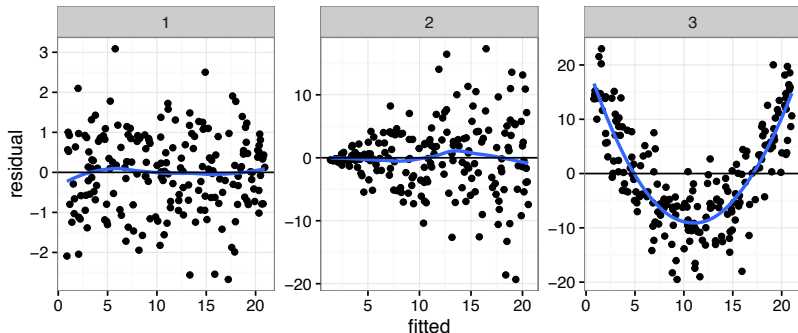
- Global lack of fit, or general breakdown of model assumptions
 - ▶ Linearity
 - ▶ Unbiased, uncorrelated errors $E(\epsilon|x) = E(\epsilon) = 0$
 - ▶ Constant variance $Var(y|x) = Var(\epsilon|x) = \sigma^2$
 - ▶ Independent errors
 - ▶ Normality of errors
- Effect of influential points and outliers

Model checking: possible solutions and strategies

- Global lack of fit, or general breakdown of model assumptions
 - ▶ Residual analysis – QQ plots, residual plots against fitted values and predictors
 - ▶ Adjusted variable plots
- Effect of influential points and outliers
 - ▶ Measure of leverage, influence, outlying-ness

Residual plots: verifying assumptions

Which assumptions (if any) do these plots show violations of?



Assumption violations are not often this obvious
(but sometimes they are!).

QQ-plots for checking Normality of residuals

QQ plot defined

QQ-plot stands for quantile-quantile plot, and is used to compare two distributions. If the two distributions are the same, then each point (which represents a quantile from each distribution) should lie along a line.

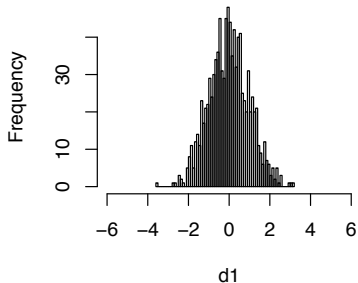
For a single (x, y) point

- x = a specific quantile for the $N(0,1)$ distribution
- y = the same quantile from the sample of data

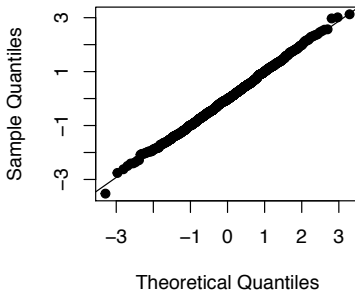
example: Gaussian or Normal(0,1) distribution

```
d1 <- rnorm(1000)
layout(matrix(1:2, nrow=1))
hist(d1, breaks=50, xlim=c(-6, 6))
qqnorm(d1, pch = 19)
qqline(d1)
```

Histogram of d1



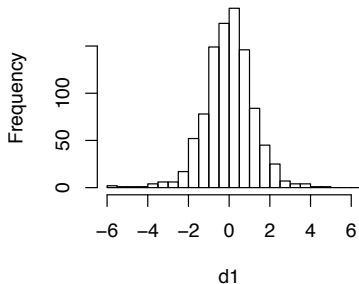
Normal Q-Q Plot



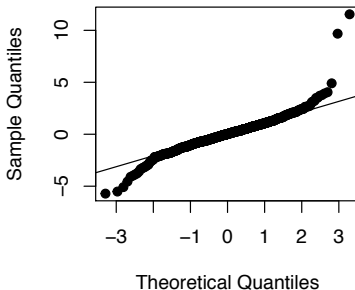
example: Student's T-distribution with 6 d.f.

```
d1 <- rt(1000, df=5)
layout(matrix(1:2, nrow=1))
hist(d1, breaks=50, xlim=c(-6, 6))
qqnorm(d1, pch = 19)
qqline(d1)
```

Histogram of d1



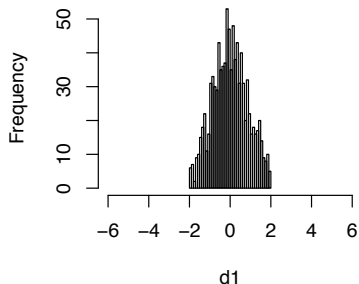
Normal Q-Q Plot



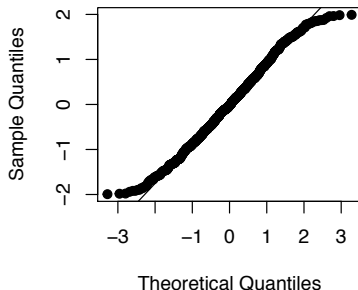
example: Truncated Gaussian

```
d1 <- rnorm(1000)
d1 <- subset(d1, abs(d1)<2)
layout(matrix(1:2, nrow=1))
hist(d1, breaks=50, xlim=c(-6, 6))
qqnorm(d1, pch = 19)
qqline(d1)
```

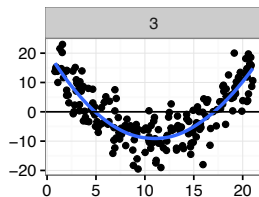
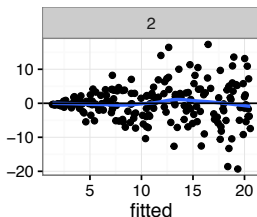
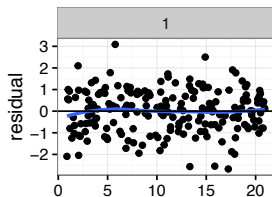
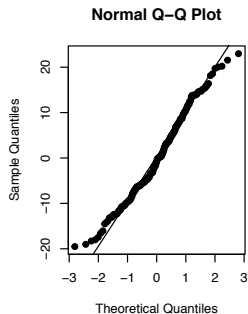
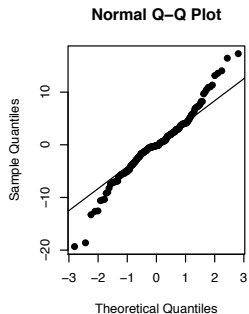
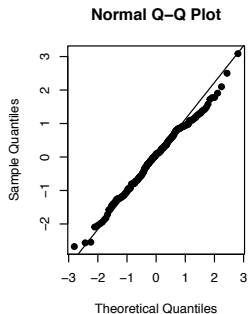
Histogram of d1



Normal Q-Q Plot



QQ-plots for our three fits from earlier



Recall: Lung Data Example

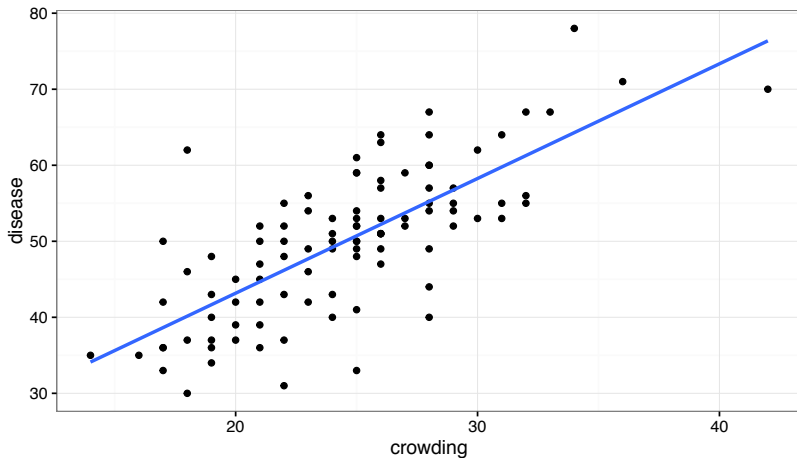
99 observations on patients who have sought treatment for the relief of respiratory disease symptoms.

The variables are:

- `disease` measure of disease severity (larger values indicates more serious condition).
- `education` highest grade completed
- `crowding` measure of crowding of living quarters (larger values indicate more crowding)
- `airqual` measure of air quality at place of residence (larger number indicates poorer quality)
- `nutrition` nutritional status (larger number indicates better nutrition)
- `smoking` smoking status (1 if smoker, 0 if non-smoker)

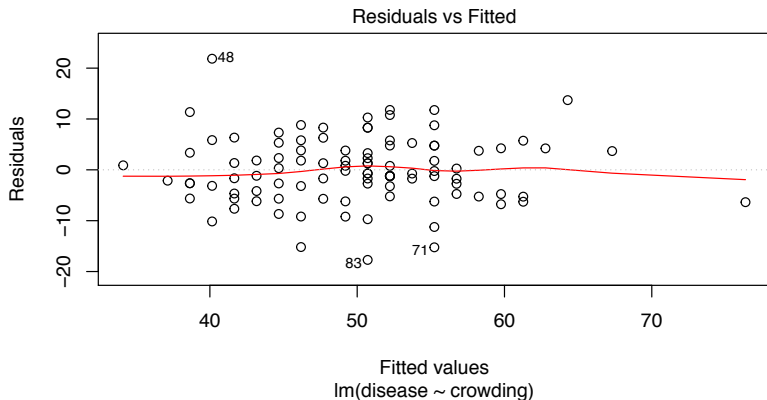
Typical regression plot: fitted line

```
ggplot(data, aes(crowding, disease)) +  
  geom_point() + geom_smooth(method="lm", se=FALSE)
```



Typical residual plot: fitted vs. residuals

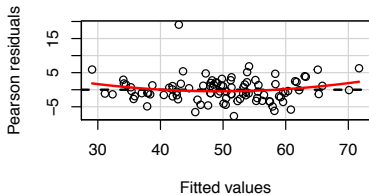
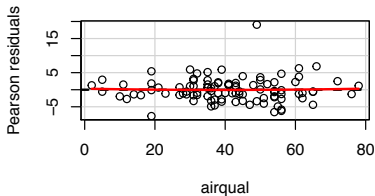
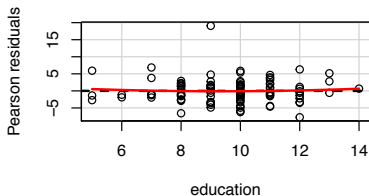
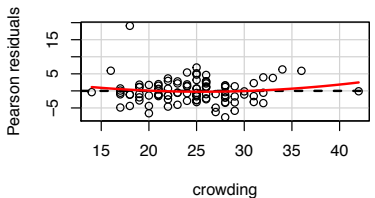
```
slr1 <- lm(disease ~ crowding, data=data)
plot(slr1, which=1)
```



But this is more complicated with MLR: how do we visualize adjusted multivariable relationships?

Predictor vs. residual plots

```
library(car)
mlr1 <- lm(disease ~ crowding + education + airqual, data=data)
residualPlots(mlr1, tests=FALSE)
```



Checking model structure: adjusted variable plots!

- You can plot residuals against each of the predictors, or plot outcomes against predictors, BUT...
- Keep in mind the MLR uses adjusted relationships; scatterplots don't show that adjustment!

Adjusted variable plots (partial regression plots, added variable plots) can be useful.

Adjusted (or added) variable plots

- Regress y on everything but x_j ; take residuals $r_{y|-x_j}$
- Regress x_j on everything but x_j ; take residuals $r_{x_j|-x_j}$
- Regress $r_{y|-x_j}$ on $r_{x_j|-x_j}$; slope of this line will match β_j in the full MLR
- Plot of $r_{y|-x_j}$ against $r_{x_j|-x_j}$ shows the “adjusted” relationship
- This figure can be used to diagnose violations of linearity in MLR models.

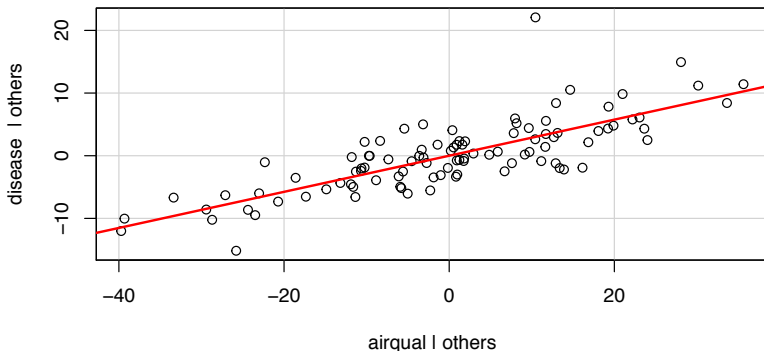
AV plots

```
coef(mlr1)
```

```
## (Intercept)    crowding    education    airqual  
## -7.7505215    1.3127837    1.4376563    0.2880687
```

```
avPlot(mlr1, variable="airqual")
```

Added-Variable Plot: airqual



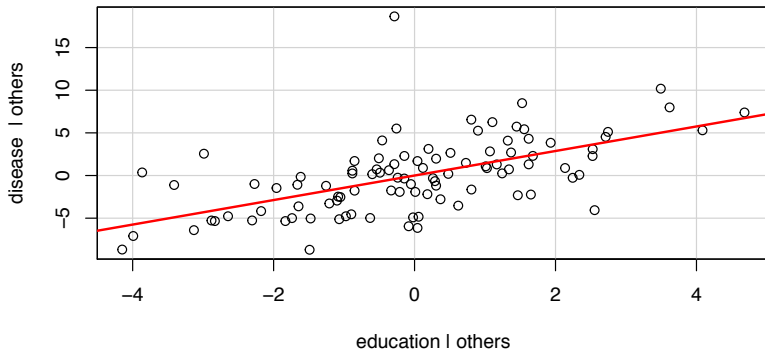
AV plots

```
coef(mlr1)
```

```
## (Intercept)    crowding    education    airqual  
## -7.7505215    1.3127837    1.4376563    0.2880687
```

```
avPlot(mlr1, variable="education")
```

Added-Variable Plot: education



Model checking: possible solutions

- Global lack of fit, or general breakdown of model assumptions
 - ▶ Residual analysis – QQ plots, residual plots against fitted values and predictors
 - ▶ Adjusted variable plots
- Effect of influential points and outliers
 - ▶ Measure of leverage, influence, outlying-ness

Isolated points

Points can be isolated in three ways

- Leverage point – outlier in x , measured by hat matrix
- Outlier – outlier in y , measured by residual
- Influential point – a point that largely affects β
 - ▶ Deletion influence; $|\hat{\beta} - \hat{\beta}_{(-i)}|$
 - ▶ Basically, a high-leverage outlier

Quantifying leverage

We measure leverage (the “distance” of \mathbf{x}_i from the distribution of \mathbf{x}) using

$$h_{ii} = \mathbf{x}_i^T (\mathbf{X}^T \mathbf{X})^{-1} \mathbf{x}_i$$

where h_{ii} is the $(i, i)^{th}$ entry of the hat matrix. Where, recall

$$\mathbf{H} = \mathbf{X}(\mathbf{X}^T \mathbf{X})^{-1} \mathbf{X}^T$$

Quantifying Leverage via the Hat Matrix

Note that

$$\sum_i h_{ii} \stackrel{\text{def}}{=} \text{tr}(\mathbf{H}) = p$$

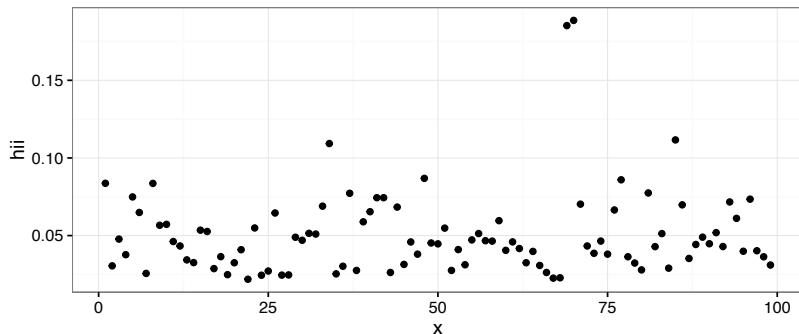
where p is the total number of independent predictors (i.e. β s) in your model (including a β_0 if you have one).

What counts as “big” leverage?

- Average leverage is p/n
- Typical rules of thumb are $2p/n$ or $3p/n$
- Leverage plots can be useful as well

Example Leverage plot with lung data

```
mlr <- lm(disease ~ nutrition+ airqual + crowding + smoking,  
          data=data)  
hii <- hatvalues(mlr)  
x <- 1:length(hii)  
qplot(x, hii, geom="point")
```



Example Leverage plot with lung data

Can be useful to investigate specific points.

```
cols <- c("disease", "crowding", "education", "airqual")
summary(data[,cols])
```

##	disease	crowding	education	airqual
##	Min. :30.00	Min. :14.00	Min. : 5.000	Min. : 2.00
##	1st Qu.:42.50	1st Qu.:21.00	1st Qu.: 8.000	1st Qu.:31.00
##	Median :51.00	Median :25.00	Median :10.000	Median :41.00
##	Mean :49.92	Mean :24.47	Mean : 9.566	Mean :40.92
##	3rd Qu.:55.00	3rd Qu.:28.00	3rd Qu.:11.000	3rd Qu.:54.00
##	Max. :78.00	Max. :42.00	Max. :14.000	Max. :78.00

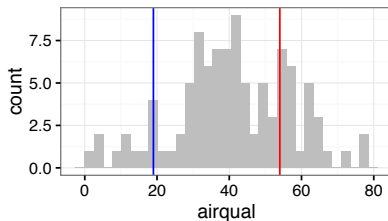
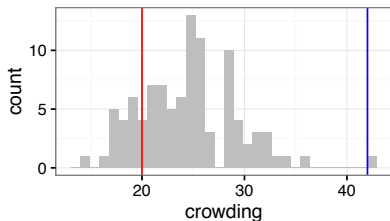

```
(d <- data[which(hii>.15), cols])
```

##	disease	crowding	education	airqual	
##	69	39	20	8	54
##	70	70	42	12	19

Example Leverage plot with lung data

Can be useful to investigate specific points.

```
library(gridExtra)
p1 <- ggplot(data) + geom_histogram(aes(x=crowding), fill="grey") +
  geom_vline(xintercept=d[1,"crowding"], color="red") +
  geom_vline(xintercept=d[2,"crowding"], color="blue")
p2 <- ggplot(data) + geom_histogram(aes(x=airqual), fill="grey") +
  geom_vline(xintercept=d[1,"airqual"], color="red") +
  geom_vline(xintercept=d[2,"airqual"], color="blue")
grid.arrange(p1, p2, ncol=2)
```



Outliers

- When we refer to “outliers” we typically mean “points that don’t have the same mean structure as the rest of the data”
- Residuals give an idea of “outlying-ness”, but we need to standardize somehow
- We can use the fact that $\text{Var}(\hat{\epsilon}_i) = \sigma^2(1 - h_{ii}) \dots$

Outliers

The *standardized* residual is given by

$$\hat{\epsilon}_i^* = \frac{\hat{\epsilon}_i}{\sqrt{\text{Var}(\hat{\epsilon}_i)}} = \frac{\hat{\epsilon}_i}{\hat{\sigma}\sqrt{(1-h_{ii})}}$$

The *Studentized* residual is given by

$$t_i = \frac{\hat{\epsilon}_{(-i)}}{\hat{\sigma}_{(-i)}\sqrt{(1-h_{ii})}} = \hat{\epsilon}_i^* \left(\frac{n-p}{n-p-\hat{\epsilon}_i^{*2}} \right)^{1/2}$$

Studentized residuals follow a t_{n-p-1} distribution.

Influence

Intuitively, “influence” is a combination of outlying-ness and leverage. More specifically, we can measure the “deletion influence” of each observation: quantify how much $\hat{\beta}$ changes if an observation is left out.

- $|\hat{\beta} - \hat{\beta}_{(-i)}|$
- Cook's distance is

$$\begin{aligned} D_i &= \frac{(\hat{\beta} - \hat{\beta}_{(-i)})^T (\mathbf{X}^T \mathbf{X}) (\hat{\beta} - \hat{\beta}_{(-i)})}{p \hat{\sigma}^2} \\ &= \frac{(\hat{\mathbf{y}} - \hat{\mathbf{y}}_{(-i)})^T (\hat{\mathbf{y}} - \hat{\mathbf{y}}_{(-i)})}{p \hat{\sigma}^2} \\ &= \frac{1}{p} \hat{\epsilon}_i^2 \frac{h_{ii}}{1 - h_{ii}} \end{aligned}$$

Handy R functions

Suppose you fit a linear model in R;

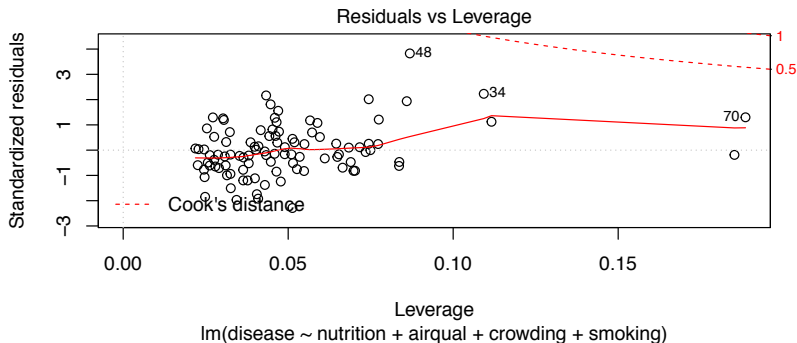
- `hatvalues` gives the diagonal elements of the hat matrix h_{ii} (leverages)
- `rstandard` gives the standardized residuals
- `rstudent` gives the studentized residuals
- `cooks.distance` gives the Cook's distances

Built-in R plots for `lm` objects

You can also use the `plot.lm()` function to look at leverage, outlying-ness, and influence all together. Recall that

$$D_i = \frac{1}{p} \hat{\epsilon}_i^2 \frac{h_{ij}}{1 - h_{ij}}$$

```
plot(mlr, which=5)
```

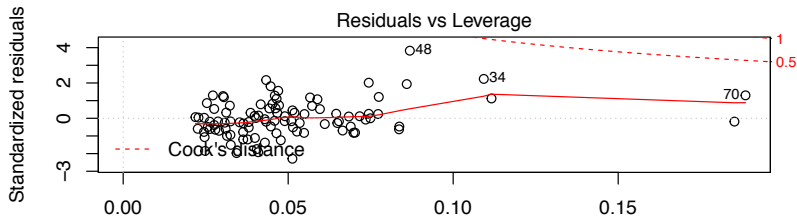


Model checking summary

You are looking for...

- Points that show worrisome level of influence \implies sensitivity analysis!
- Systematic departures from model assumptions \implies transformations, different model structure
- Unrealistic outliers \implies check your data!

No points show worrisome influence in this lung data analysis, although observation 70 showed up in both of our analyses.



Back to the outline

- Model checking and diagnostics
- Variable transformations

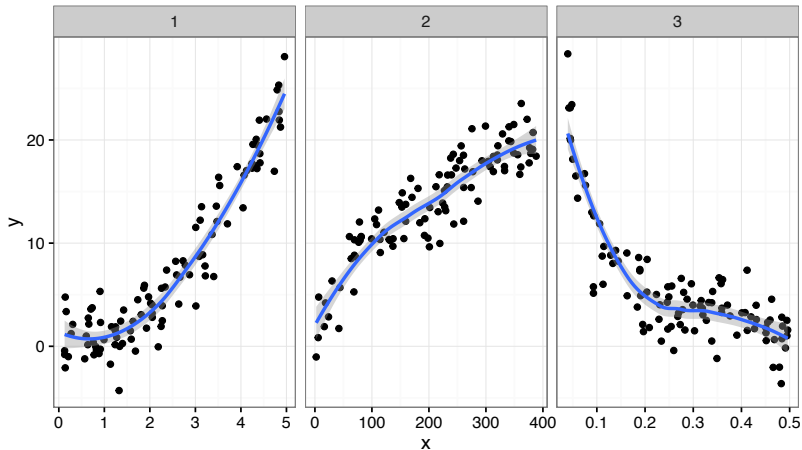
Overview of variable transformations

The problems

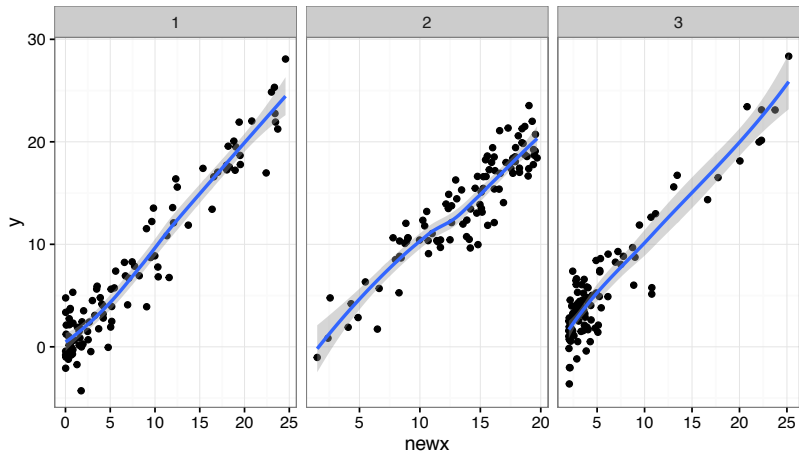
- Non-linearity between X and Y \longrightarrow transform X
- Skewed distribution of X s/points with high leverage \longrightarrow transform X
- Non-constant variance \longrightarrow transform Y

Transforming your X variables

Transforming predictor variables can help with constant-variance non-linear relationships.



Transforming your X variables



β interpretations with transformed X s

Transforming predictor variables can help with non-linearities, but can make coefficient interpretations hard.

Possible solutions

- Interpret β s qualitatively across a region of interest: “We found strong evidence for an inverse association, where values of Y were inversely proportional to X across the observed range (a, b) .”
- Occasionally, a “one unit change in X ” can be meaningful: e.g. $\log_a X$. A one unit change in $\log_a X$ indicates a a -fold increase in X .

β interpretations with transformed X s

- Transforming predictor variables can help with non-linearities, but can make coefficient interpretations hard.
- Can also use polynomials, splines (more soon!).

Transforming Y s for non-constant variance

What to do ...

- Nothing; just use least squares and bootstrap
- Use weighted LS, GLS (Biostat Methods 3?)
- Use a variance stabilizing transformation
- Consider a generalized linear model (more soon)

Box-Cox Transformations

Outcome is raised to the λ power:

$$y_i^\lambda = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \epsilon_i$$

- Estimate λ , a new parameter, by maximum likelihood.
- Some well-known choices of λ : 2, -1, 1/2
- By definition, when $\lambda = 0$, we specify $y_i^\lambda = \log_e y_i$

Today's big ideas

- Model checking
- Variable transformations
- Next up: inference about MLR parameters