

Linear, Generalized, and Mixed/Multilevel models - an introduction with R

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Introduction to linear models

Modern statistics are easier than this

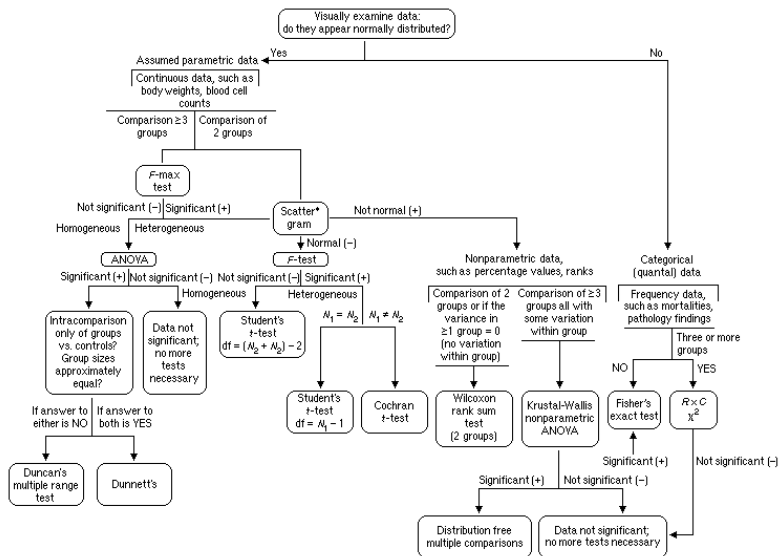
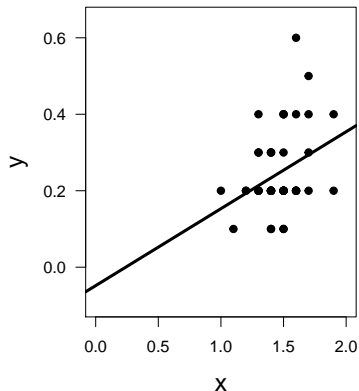


Figure 1:

Our overarching regression framework

$$y_i = a + bx_i + \varepsilon_i$$

$$\varepsilon_i \sim N(0, \sigma^2)$$



Data

y = response variable

x = predictor

Parameters

a = intercept

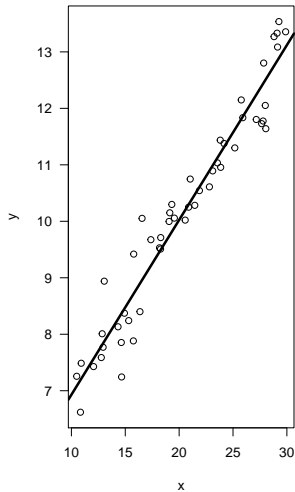
b = slope

σ = residual variation

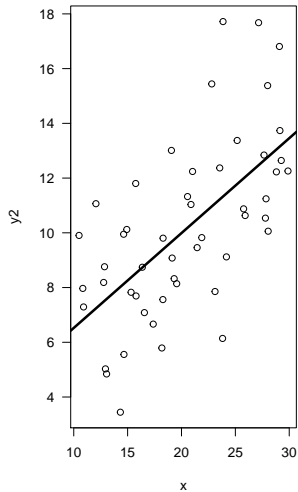
ε = residuals

Residual variation (error)

small



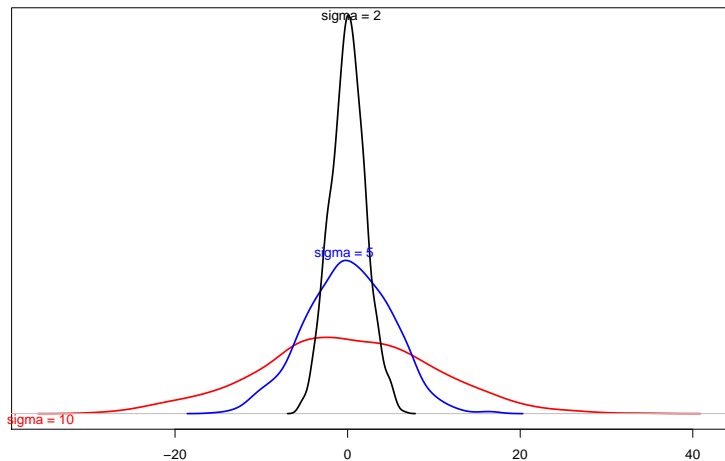
large



Residual variation

$$\varepsilon_i \sim N(0, \sigma^2)$$

Distribution of residuals



In a Normal distribution

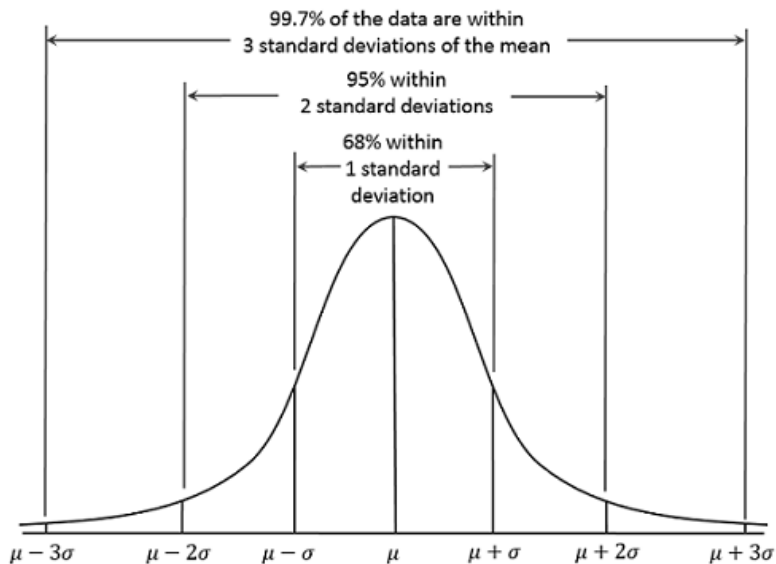


Figure 2:

Linear models

Example dataset: forest trees

- Go to <https://tinyurl.com/treesdata>

```
trees <- read.csv("data-raw/trees.csv")  
head(trees)
```

	plot	dbh	height	sex	dead
1	4	29.68	36.1	male	0
2	5	33.29	42.3	male	0
3	2	28.03	41.9	female	0
4	5	39.86	46.5	female	0
5	1	47.94	43.9	female	0
6	1	10.82	26.2	male	0

Example dataset: forest trees

- ▶ Go to <https://tinyurl.com/treesdata>
- ▶ Download zip file and uncompress (within your project folder!)

```
trees <- read.csv("data-raw/trees.csv")  
head(trees)
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	plot	dbh	height	sex	dead
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Questions

- ▶ What is the relationship between DBH and height?

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- ▶ Do taller trees have bigger trunks?

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- ▶ Do taller trees have bigger trunks?
- ▶ Can we predict height from DBH? How well?

Always plot your data first!

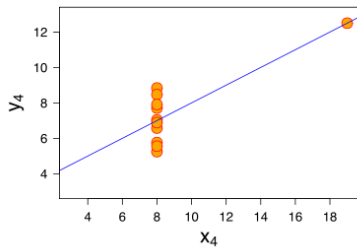
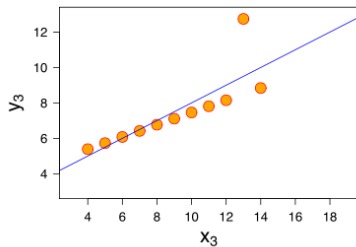
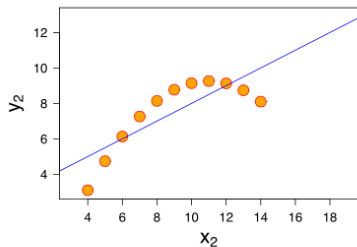
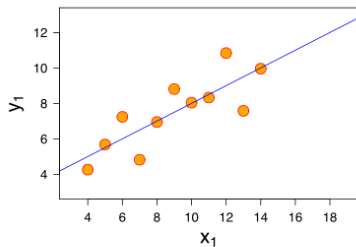
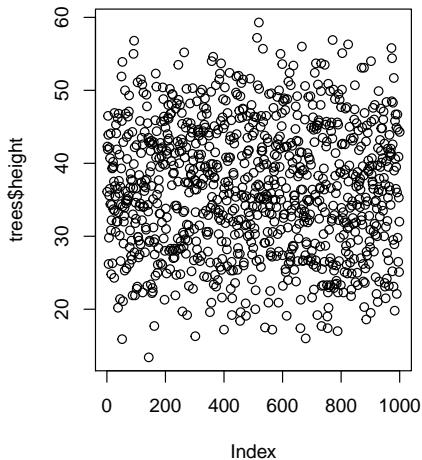


Figure 3:

Exploratory Data Analysis (EDA)

Outliers

```
plot(trees$height)
```



Outliers impact on regression

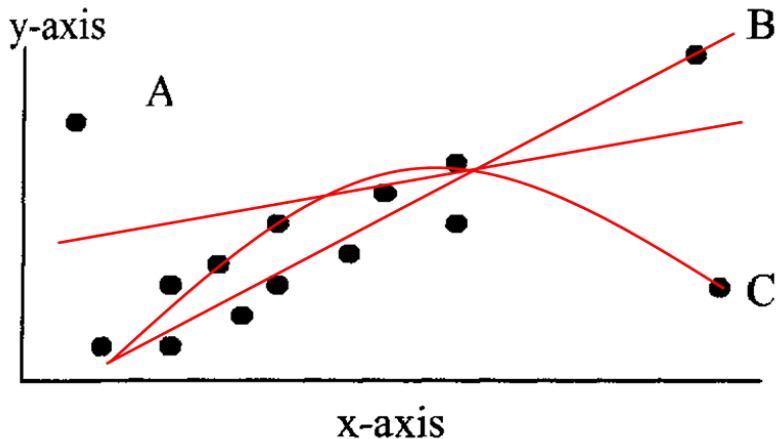
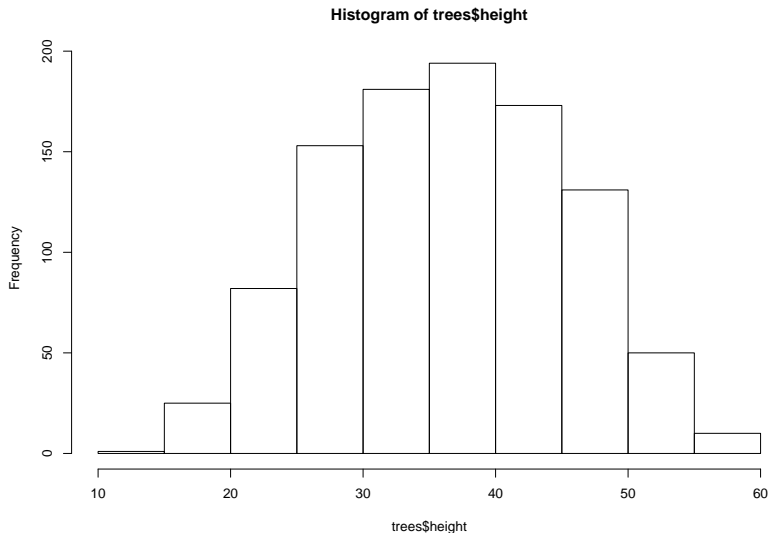


Figure 4:

See <http://rpsychologist.com/d3/correlation/>

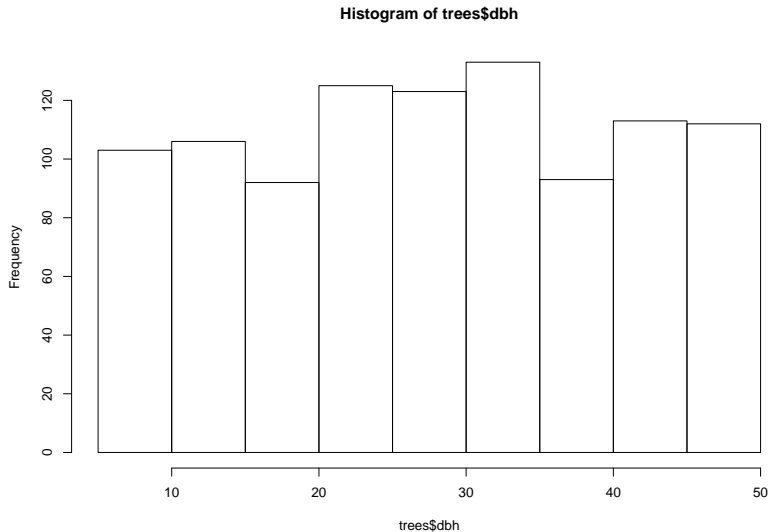
Histogram of response variable

```
hist(trees$height)
```



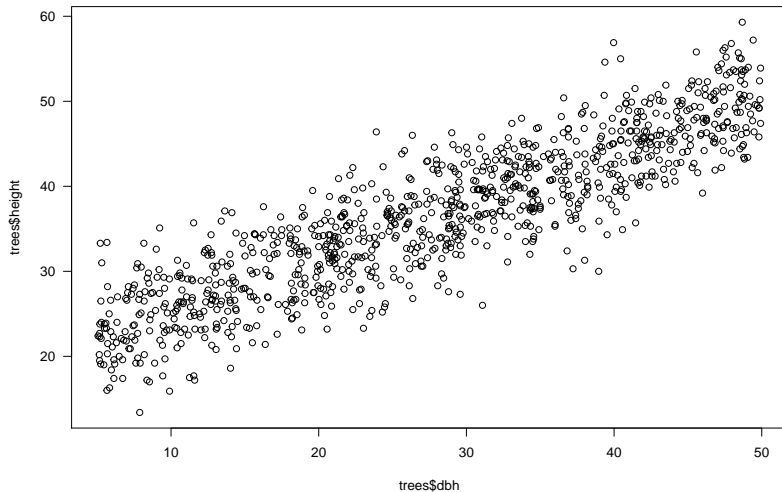
Histogram of predictor variable

```
hist(trees$dbh)
```



Scatterplot

```
plot(trees$dbh, trees$height, las = 1)
```



Now fit model

Hint: `lm`

Now fit model

Hint: `lm`

```
m1 <- lm(height ~ dbh, data = trees)
```

What does this mean?

Call:

```
lm(formula = height ~ dbh, data = trees)
```

Residuals:

Min	1Q	Median	3Q	Max
-13.3270	-2.8978	0.1057	2.7924	12.9511

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	19.33920	0.31064	62.26	<2e-16 ***
dbh	0.61570	0.01013	60.79	<2e-16 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 4.093 on 998 degrees of freedom

Multiple R-squared: 0.7874, Adjusted R-squared: 0.7871

F-statistic: 3695 on 1 and 998 DF, p-value: < 2.2e-16

Retrieving model coefficients

```
coef(m1)
```

(Intercept)	dbh
19.3391968	0.6157036

Confidence intervals

```
confint(m1)
```

	2.5 %	97.5 %
(Intercept)	18.7296053	19.948788
dbh	0.5958282	0.635579

Using effects package

```
library(effects)  
summary(allEffects(m1))
```

model: height ~ dbh

dbh effect

dbh

	5	20	30	40	50
	22.41771	31.65327	37.81030	43.96734	50.12438

Lower 95 Percent Confidence Limits

dbh

	5	20	30	40	50
	21.89682	31.35487	37.55287	43.61733	49.61669

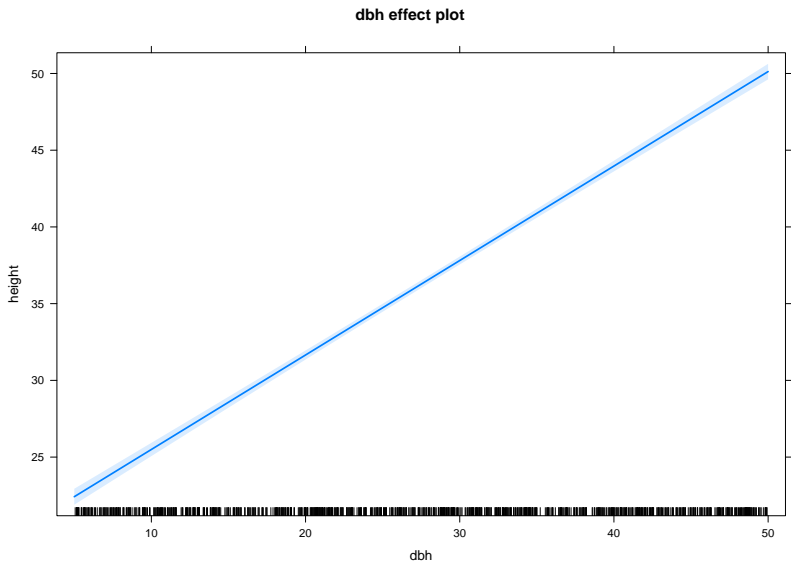
Upper 95 Percent Confidence Limits

dbh

	5	20	30	40	50
	22.93861	31.95167	38.06774	44.31735	50.63207

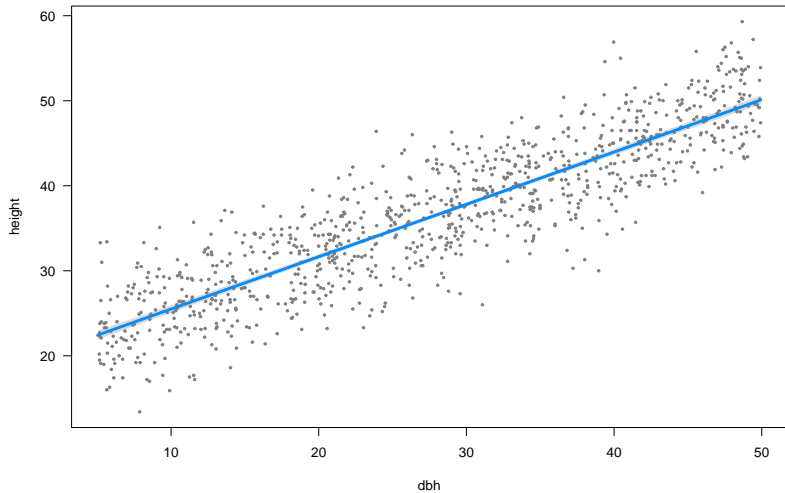
Plot effects

```
plot(allEffects(m1))
```



Plot model (visreg)

```
library(visreg)  
visreg(m1)
```



Linear model assumptions

- ▶ Linearity (transformations, GAM...)

Linear model assumptions

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- ▶ Residuals:

Linear model assumptions

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- ▶ Residuals:
 - ▶ Independent

Linear model assumptions

- ▶ Linearity (transformations, GAM...)
- ▶ Residuals:
 - ▶ Independent
 - ▶ Equal variance

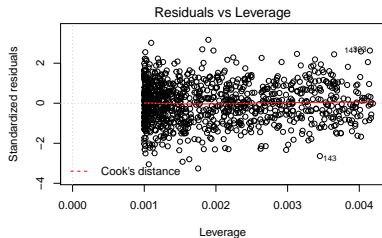
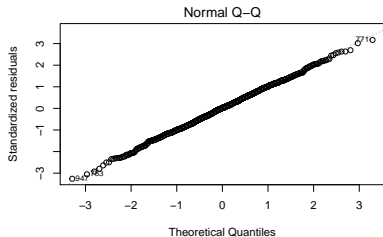
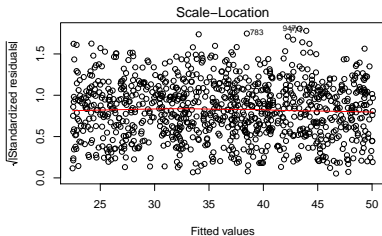
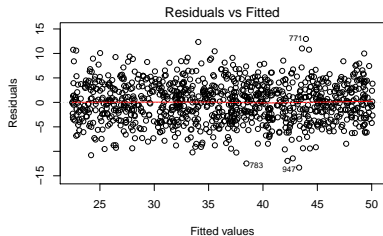
Linear model assumptions

- ▶ Linearity (transformations, GAM...)
- ▶ Residuals:
 - ▶ Independent
 - ▶ Equal variance
 - ▶ Normal

Linear model assumptions

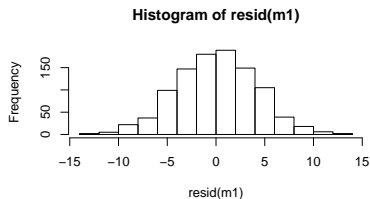
- ▶ Linearity (transformations, GAM...)
- ▶ Residuals:
 - ▶ Independent
 - ▶ Equal variance
 - ▶ Normal
- ▶ No measurement error in predictors

Model checking: residuals



Are residuals normal?

```
hist(resid(m1))
```



```
lm(formula = height ~ dbh, data = trees)
      coef.est coef.se
(Intercept) 19.34    0.31
      dbh      0.62    0.01
---
n = 1000, k = 2
residual sd = 4.09, R-Squared = 0.79
```

SD of residuals = 4.09 coincides with estimate of σ .

How good is the model in predicting tree height?

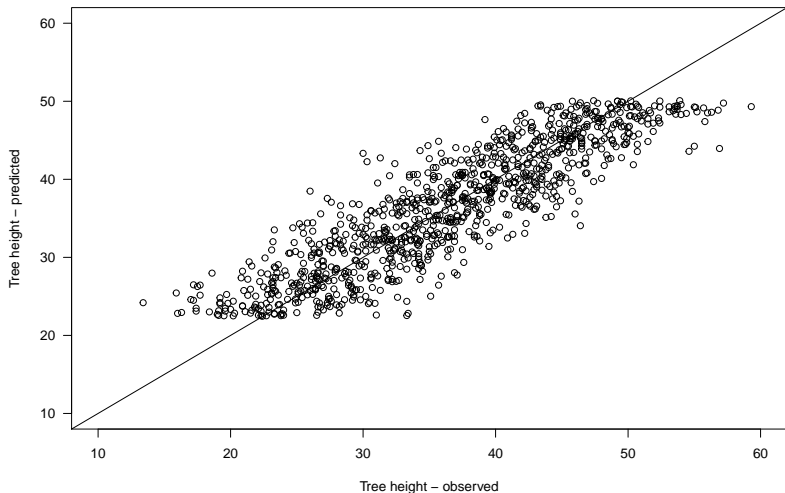
fitted gives predictions for each observation

```
trees$height.pred <- fitted(m1)  
head(trees)
```

	plot	dbh	height	sex	dead	height.pred
1	4	29.68	36.1	male	0	37.61328
2	5	33.29	42.3	male	0	39.83597
3	2	28.03	41.9	female	0	36.59737
4	5	39.86	46.5	female	0	43.88114
5	1	47.94	43.9	female	0	48.85603
6	1	10.82	26.2	male	0	26.00111

Calibration plot: Observed vs Predicted values

```
plot(trees$height, trees$height.pred, xlab = "Tree height - obse
```



Using fitted model for prediction

Q: Expected tree height if DBH = 39 cm?

```
new.dbh <- data.frame(dbh = c(39))  
predict(m1, new.dbh, se.fit = TRUE)
```

```
$fit
```

```
1
```

```
43.35164
```

```
$se.fit
```

```
[1] 0.1715514
```

```
$df
```

```
[1] 998
```

```
$residual.scale
```

```
[1] 4.092629
```

Important functions

- ▶ `plot`

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- ▶ `plot`
- ▶ `summary`

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- ▶ `plot`
- ▶ `summary`
- ▶ `coef`

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Important functions

- ▶ `plot`
- ▶ `summary`
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- ▶ `confint`
- ▶ `fitted`
- ▶ `resid`
- ▶ `allEffects`

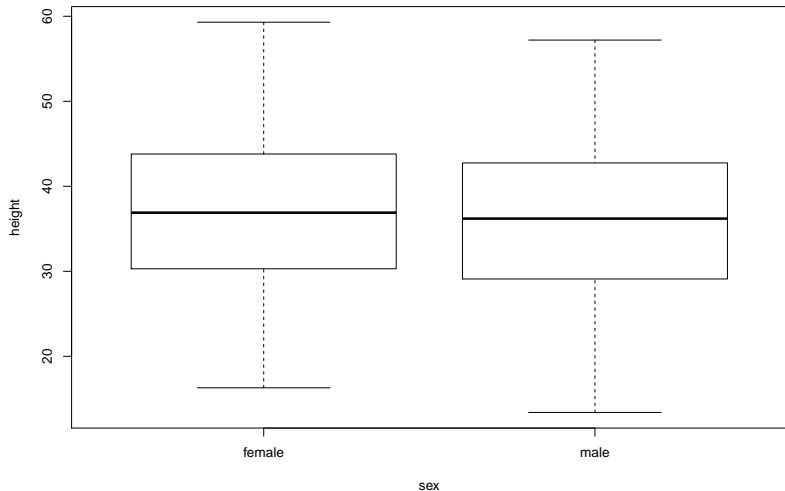
Important functions

- ▶ `plot`
- ▶ `summary`
- ▶ `coef`
- ▶ `confint`
- ▶ `fitted`
- ▶ `resid`
- ▶ `allEffects`
- ▶ `predict`

Categorical predictors (factors)

Q: Does tree height vary with sex?

```
plot(height ~ sex, data = trees)
```



Model height ~ sex

```
m2 <- lm(height ~ sex, data = trees)
```

Call:

```
lm(formula = height ~ sex, data = trees)
```

Residuals:

Min	1Q	Median	3Q	Max
-22.6881	-6.7881	-0.0097	6.7261	22.3687

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	36.9312	0.3981	92.778	<2e-16 ***
sexmale	-0.8432	0.5607	-1.504	0.133

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 8.865 on 998 degrees of freedom

Multiple R-squared: 0.002261, Adjusted R-squared: 0.001261

F-statistic: 2.261 on 1 and 998 DF, p-value: 0.133

Linear model with categorical predictors

$$y_i = a + bx_i + \varepsilon_i$$

$$y_i = a + b_{male} + \varepsilon_i$$

Model height ~ sex

```
m2 <- lm(height ~ sex, data = trees)
```

Call:

```
lm(formula = height ~ sex, data = trees)
```

Residuals:

Min	1Q	Median	3Q	Max
-22.6881	-6.7881	-0.0097	6.7261	22.3687

Coefficients:

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Effects: Height ~ sex

Compare CIs

```
summary(allEffects(m2))
```

```
model: height ~ sex
```

```
sex effect
```

```
sex
```

```
female      male
```

```
36.93125 36.08810
```

```
Lower 95 Percent Confidence Limits
```

```
sex
```

```
female      male
```

```
36.15012 35.31319
```

```
Upper 95 Percent Confidence Limits
```

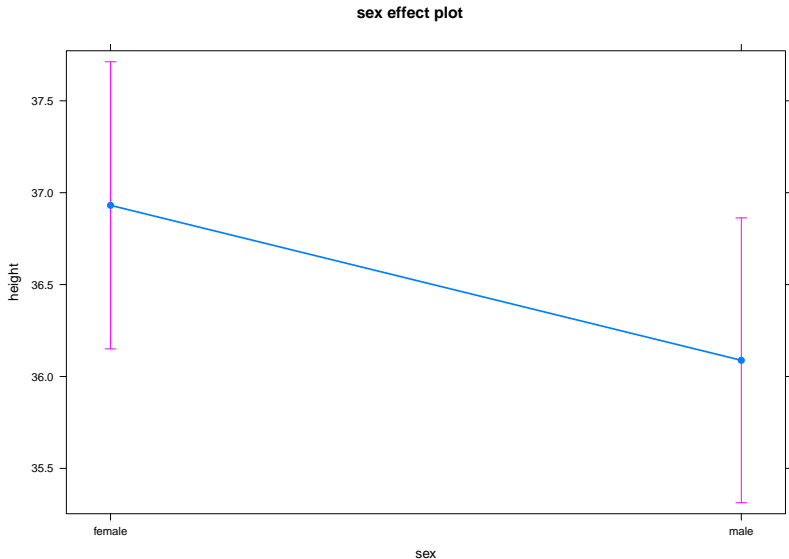
```
sex
```

```
female      male
```

```
37.71238 36.86300
```

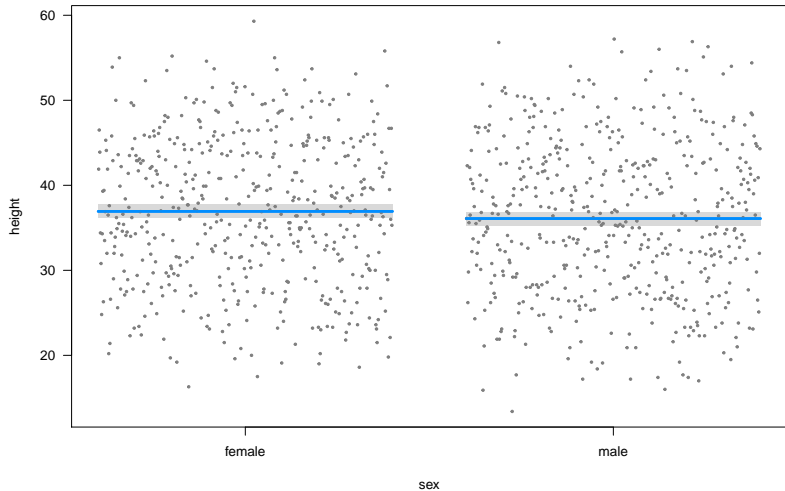
Plot

```
plot(allEffects(m2))
```

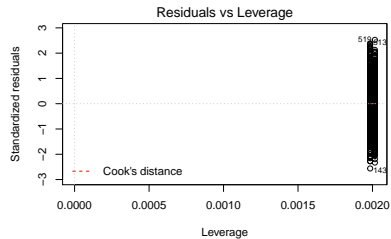
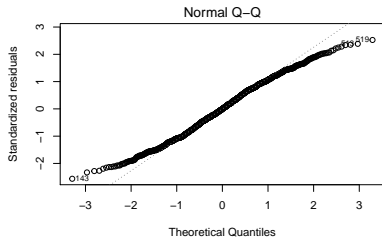
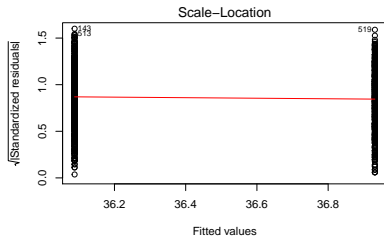
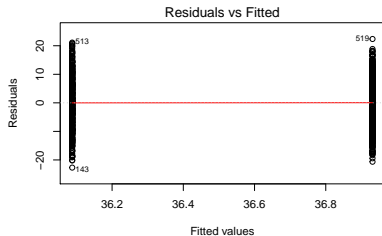


Plot (visreg)

```
visreg(m2)
```

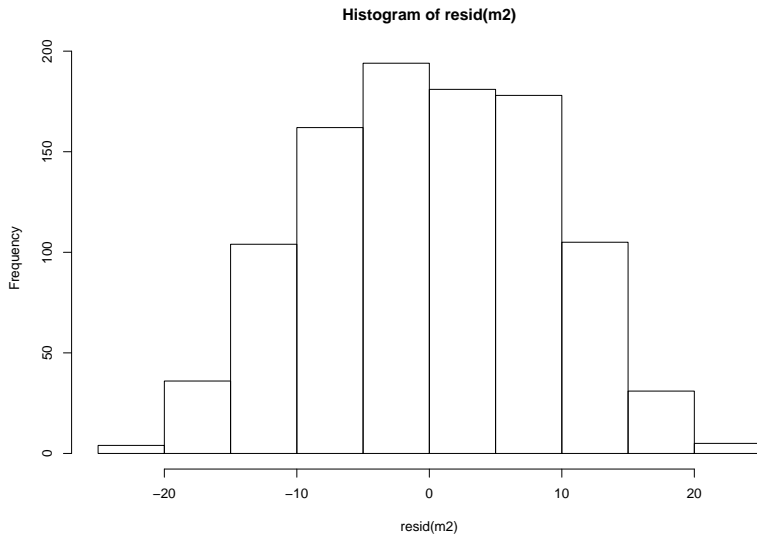


Model checking: residuals



Model checking: residuals

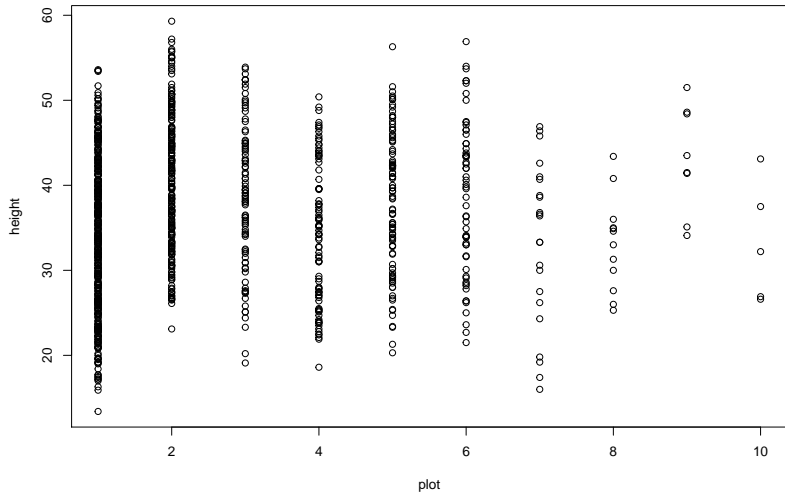
```
hist(resid(m2))
```



Q: Does height differ among field plots?

Plot data first

```
plot(height ~ plot, data = trees)
```



Linear model with categorical predictors

$$y_i = a + bx_i + \varepsilon_i$$

$$y_i = a + b_{plot2} + c_{plot3} + d_{plot4} + e_{plot5} + \dots + \varepsilon_i$$

Model Height ~ Plot

All right here?

```
m3 <- lm(height ~ plot, data = trees)
```

Call:

```
lm(formula = height ~ plot, data = trees)
```

Residuals:

	Min	1Q	Median	3Q	Max
	-22.4498	-6.7049	0.0709	6.7537	23.0640

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	35.4636	0.4730	74.975	< 2e-16 ***
plot	0.3862	0.1413	2.733	0.00639 **

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 8.842 on 998 degrees of freedom

Multiple R-squared: 0.007429, Adjusted R-squared: 0.006435

Plot is a factor!

```
trees$plot <- as.factor(trees$plot)
```

Model Height ~ Plot

Call:

```
lm(formula = height ~ plot, data = trees)
```

Residuals:

	Min	1Q	Median	3Q	Max
	-20.4416	-6.9004	0.0379	6.3051	19.7584

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)	
(Intercept)	33.8416	0.4266	79.329	< 2e-16	***
plot2	6.3411	0.7126	8.899	< 2e-16	***
plot3	4.9991	0.9828	5.086	4.36e-07	***
plot4	0.5329	0.9872	0.540	0.58949	
plot5	4.3723	0.9425	4.639	3.97e-06	***
plot6	4.7601	1.1709	4.065	5.18e-05	***
plot7	-0.7416	1.8506	-0.401	0.68871	
plot8	-0.6832	2.4753	-0.276	0.78258	
plot9	9.1709	3.0165	3.040	0.00243	**
plot10	-0.5816	3.8013	-0.153	0.87843	

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 8.446 on 990 degrees of freedom

Multiple R-squared: 0.1016, Adjusted R-squared: 0.09344

F-statistic: 12.44 on 9 and 990 DF, p-value: < 2.2e-16

Estimated tree heights for each site

```
summary(allEffects(m3))
```

```
model: height ~ plot
```

```
plot effect
```

```
plot
```

	1	2	3	4	5	6	7	8
	33.84158	40.18265	38.84066	34.37444	38.21386	38.60167	33.10000	33.15833
	9	10						
	43.01250	33.26000						

```
Lower 95 Percent Confidence Limits
```

```
plot
```

	1	2	3	4	5	6	7	8
	33.00444	39.06264	37.10317	32.62733	36.56463	36.46190	29.56629	28.37367
	9	10						
	37.15251	25.84764						

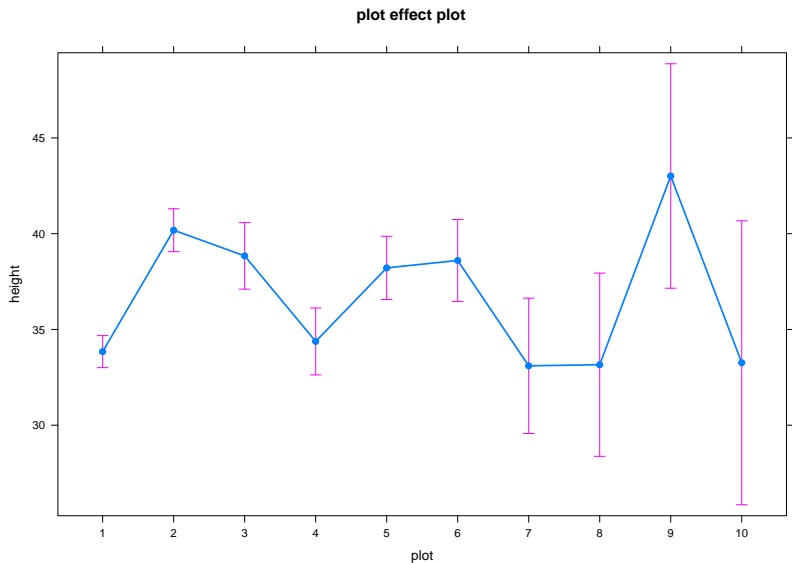
```
Upper 95 Percent Confidence Limits
```

```
plot
```

	1	2	3	4	5	6	7	8
	34.67872	41.30265	40.57814	36.12156	39.86309	40.74143	36.63371	37.94299
	9	10						
	48.87249	40.67236						

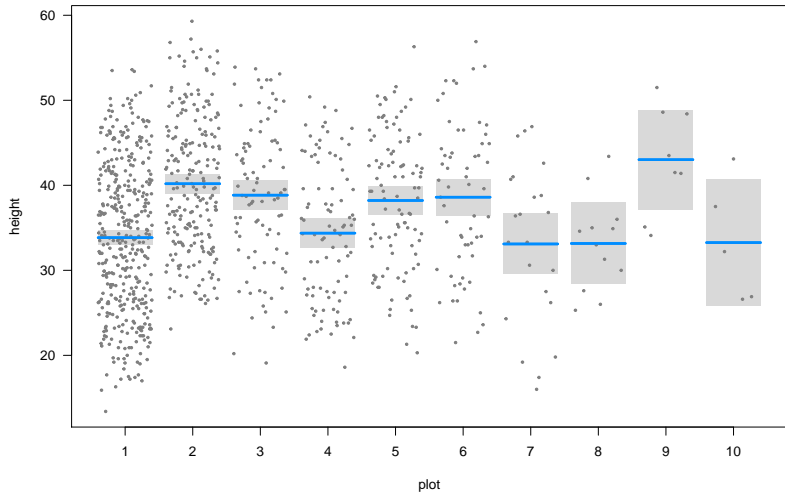
Plot

```
plot(allEffects(m3))
```

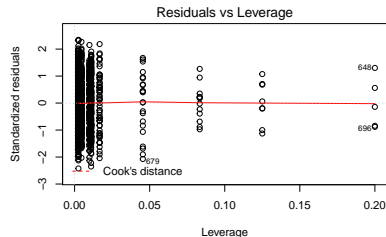
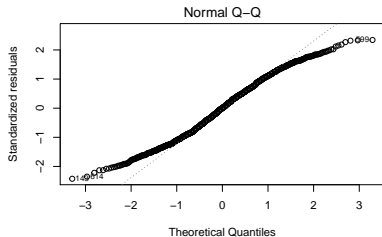
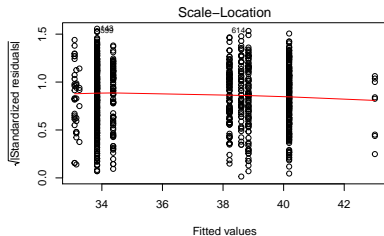
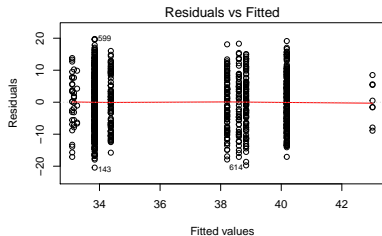


Plot (visreg)

```
visreg(m3)
```



Model checking: residuals



Combining continuous and categorical predictors

Predicting tree height based on dbh and site

$$y_i = a + bx_i + \varepsilon_i$$

$$y_i = a + b_{plot2} + c_{plot3} + d_{plot4} + e_{plot5} + \dots + k \cdot DBH_i + \varepsilon_i$$

Predicting tree height based on dbh and site

Call:

```
lm(formula = height ~ plot + dbh, data = trees)
```

Residuals:

	Min	1Q	Median	3Q	Max
	-10.1130	-1.9885	0.0582	2.0314	11.3320

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	16.699037	0.260565	64.088	< 2e-16 ***
plot2	6.504303	0.256730	25.335	< 2e-16 ***
plot3	4.357457	0.354181	12.303	< 2e-16 ***
plot4	1.934650	0.356102	5.433	6.98e-08 ***
plot5	3.637432	0.339688	10.708	< 2e-16 ***
plot6	4.204511	0.421906	9.966	< 2e-16 ***
plot7	-0.176193	0.666772	-0.264	0.7916
plot8	-5.312648	0.893603	-5.945	3.82e-09 ***
plot9	5.437049	1.087766	4.998	6.84e-07 ***
plot10	2.263338	1.369986	1.652	0.0988 .
dbh	0.617075	0.007574	81.473	< 2e-16 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 3.043 on 989 degrees of freedom

Multiple R-squared: 0.8835, Adjusted R-squared: 0.8823

F-statistic: 750 on 10 and 989 DF, p-value: < 2.2e-16

Estimated tree heights for each site

```
summary(allEffects(multreg))
```

```
model: height ~ plot + dbh
```

```
plot effect
```

```
plot
```

	1	2	3	4	5	6	7	8
	33.90437	40.40868	38.26183	35.83902	37.54181	38.10889	33.72818	28.59173
	9	10						
	39.34142	36.16771						

```
Lower 95 Percent Confidence Limits
```

```
plot
```

	1	2	3	4	5	6	7	8
	33.60276	40.00512	37.63569	35.20858	36.94739	37.33787	32.45495	26.86438
	9	10						
	37.22831	33.49623						

```
Upper 95 Percent Confidence Limits
```

```
plot
```

	1	2	3	4	5	6	7	8
	34.20599	40.81223	38.88798	36.46947	38.13622	38.87990	35.00141	30.31907
	9	10						
	41.45454	38.83919						

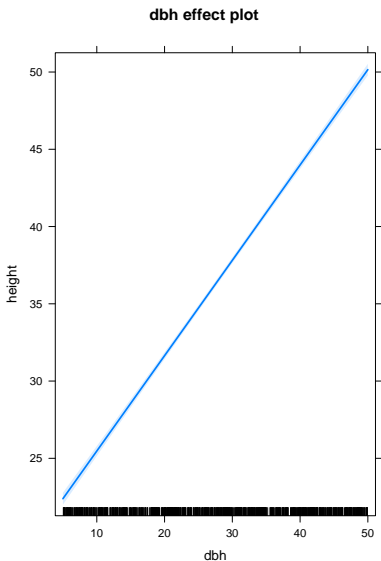
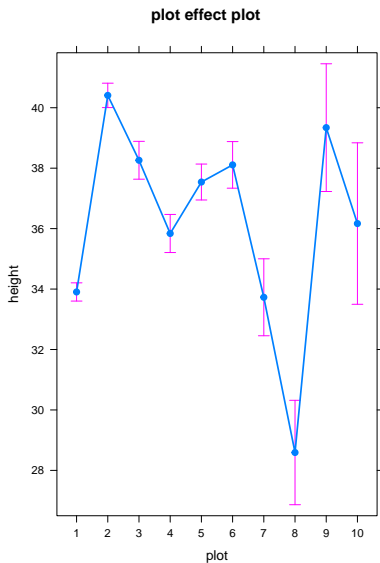
```
dbh effect
```

```
dbh
```

	5	20	30	40	50
	22.38634	31.64246	37.81321	43.98396	50.15471

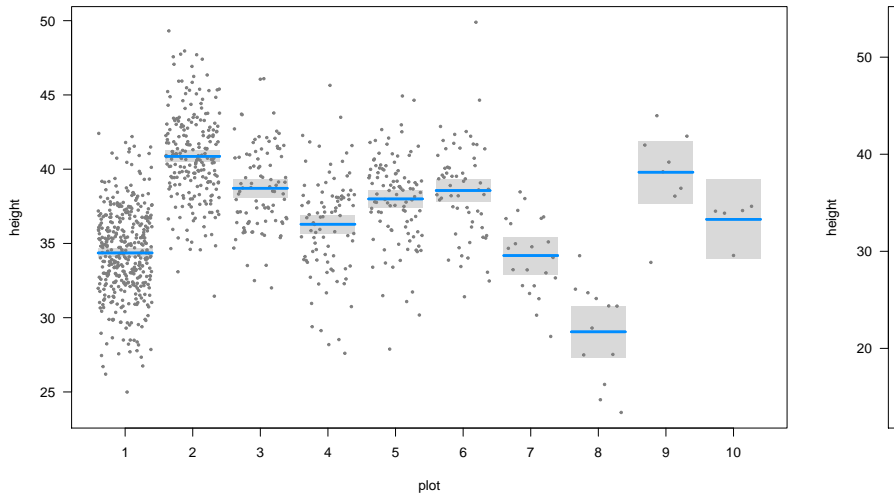
Plot

```
plot(allEffects(multreg))
```

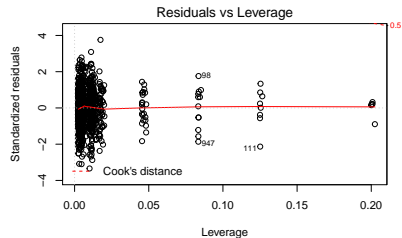
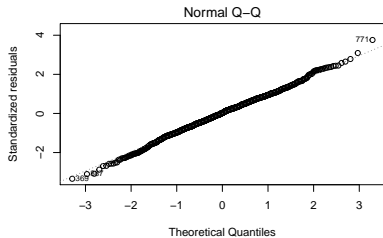
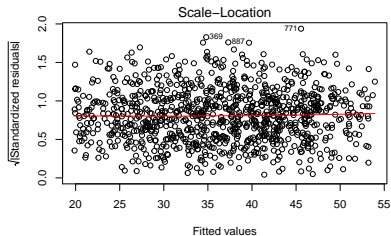
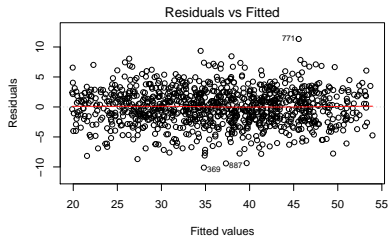


Plot (visreg)

```
visreg(multreg)
```



Model checking: residuals



Extra exercises

- ▶ paperplanes: How does flight distance differ with age, gender or paper type?

Extra exercises

- ▶ paperplanes: How does flight distance differ with age, gender or paper type?
- ▶ mammal sleep: Are sleep patterns related to diet?

Extra exercises

- ▶ paperplanes: How does flight distance differ with age, gender or paper type?
- ▶ mammal sleep: Are sleep patterns related to diet?
- ▶ iris: Predict petal length \sim petal width and species

Generalised Linear Models: Logistic regression

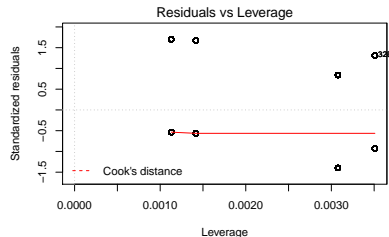
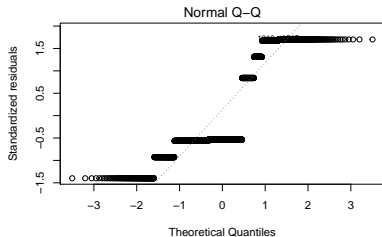
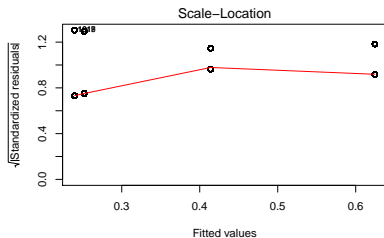
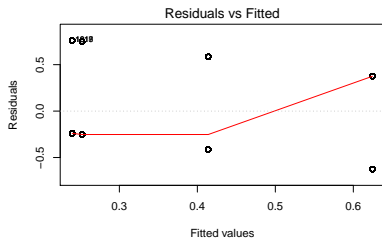
Q: Survival of passengers on the Titanic ~ Class

Read titanic_long.csv dataset.

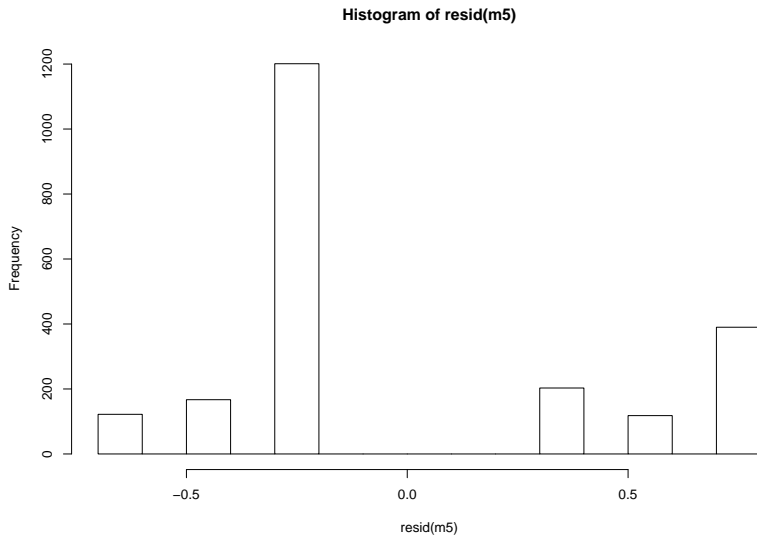
	class	age	sex	survived
1	first	adult	male	1
2	first	adult	male	1
3	first	adult	male	1
4	first	adult	male	1
5	first	adult	male	1
6	first	adult	male	1

Let's fit linear model:

```
m5 <- lm(survived ~ class, data = titanic)
```



Weird residuals!



What if your residuals are clearly non-normal? | And variance not constant (heteroscedasticity)?

- ▶ Binary variables (0/1)

What if your residuals are clearly non-normal? | And variance not constant (heteroscedasticity)?

- ▶ Binary variables (0/1)
- ▶ Counts (0, 1, 2, 3, ...)

Generalised Linear Models

1. **Response variable** - distribution family

Generalised Linear Models

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 - ▶ Bernoulli - Binomial

Generalised Linear Models

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- ▶ Poisson

Generalised Linear Models

1. **Response variable** - distribution family

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- ▶ Poisson
- ▶ Gamma

Generalised Linear Models

1. **Response variable** - distribution family

- ▶ Bernoulli - Binomial
- ▶ Poisson
- ▶ Gamma
- ▶ etc

Generalised Linear Models

1. **Response variable** - distribution family
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2. **Predictors** (continuous or categorical)

Generalised Linear Models

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3. **Link function**

Generalised Linear Models

1. **Response variable** - distribution family
 - ▶ Bernoulli - Binomial
 - ▶ Poisson
 - ▶ Gamma
 - ▶ etc
2. **Predictors** (continuous or categorical)
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 - ▶ Gaussian: identity

Generalised Linear Models

1. **Response variable** - distribution family

- ▶ Bernoulli - Binomial
- ▶ Poisson
- ▶ Gamma
- ▶ etc

2. **Predictors** (continuous or categorical)

3. **Link function**

- ▶ Gaussian: identity
- ▶ Binomial: logit, probit

Generalised Linear Models

1. **Response variable** - distribution family

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- ▶ Gamma
- ▶ etc

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3. **Link function**

- ▶ Gaussian: identity
- ▶ Binomial: logit, probit
- ▶ Poisson: log...

Generalised Linear Models

1. **Response variable** - distribution family

- ▶ Bernoulli - Binomial
- ▶ Poisson
- ▶ Gamma
- ▶ etc

2. **Predictors** (continuous or categorical)

3. **Link function**

- ▶ Gaussian: identity
- ▶ Binomial: logit, probit
- ▶ Poisson: log...
- ▶ See family.

The modelling process

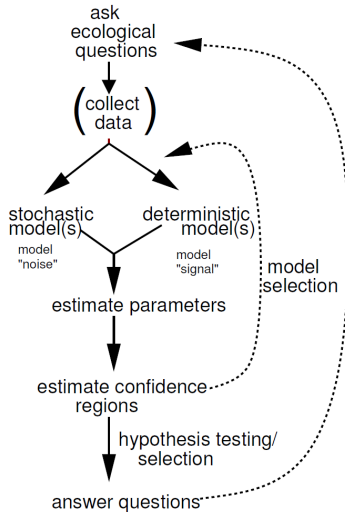


Figure 1.5 Flow of the modeling process.

Figure 5:

Bernoulli - Binomial distribution (Logistic regression)

- Response variable: Yes/No (e.g. survival, sex, presence/absence)

$$\text{logit}(p) = \ln \left(\frac{p}{1-p} \right)$$

Then

$$\text{Pr}(\text{alive}) = a + bx$$

$$\text{logit}(\text{Pr}(\text{alive})) = a + bx$$

$$\text{Pr}(\text{alive}) = \text{invlogit}(a + bx) = \frac{e^{a+bx}}{1 + e^{a+bx}}$$

Bernoulli - Binomial distribution (Logistic regression)

- ▶ Response variable: Yes/No (e.g. survival, sex, presence/absence)
- ▶ Link function: `logit` (others possible, see family).

$$\text{logit}(p) = \ln \left(\frac{p}{1-p} \right)$$

Then

$$\text{Pr}(\text{alive}) = a + bx$$

$$\text{logit}(\text{Pr}(\text{alive})) = a + bx$$

$$\text{Pr}(\text{alive}) = \text{invlogit}(a + bx) = \frac{e^{a+bx}}{1 + e^{a+bx}}$$

Back to survival of Titanic passengers

How many passengers travelled in each class?

Back to survival of Titanic passengers

How many passengers travelled in each class?

```
tapply(titanic$survived, titanic$class, length)
```

crew	first	second	third
885	325	285	706

Back to survival of Titanic passengers

How many passengers travelled in each class?

```
tapply(titanic$survived, titanic$class, length)
```

crew	first	second	third
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How many survived?

Back to survival of Titanic passengers

How many passengers travelled in each class?

```
tapply(titanic$survived, titanic$class, length)
```

crew	first	second	third
885	325	285	706

How many survived?

```
tapply(titanic$survived, titanic$class, sum)
```

crew	first	second	third
212	203	118	178

Back to survival of Titanic passengers

How many passengers travelled in each class?

```
tapply(titanic$survived, titanic$class, length)
```

crew	first	second	third
885	325	285	706

How many survived?

```
tapply(titanic$survived, titanic$class, sum)
```

crew	first	second	third
212	203	118	178

What proportion survived in each class?

```
as.numeric(tapply(titanic$survived, titanic$class, mean))
```

```
[1] 0.2395480 0.6246154 0.4140351 0.2521246
```

Back to survival of Titanic passengers (dplyr)

Passenger survival according to class

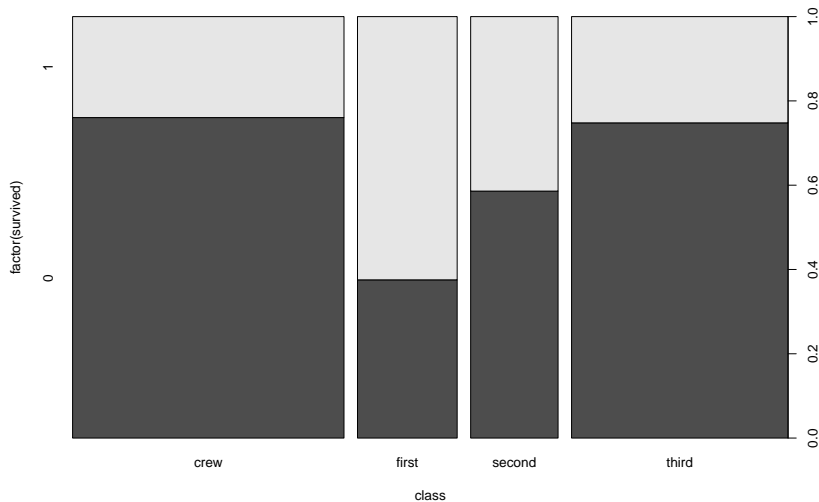
```
library(dplyr)
titanic %>%
  group_by(class, survived) %>%
  summarise(count = n())
```

```
# A tibble: 8 x 3
# Groups:   class [?]
  class survived count
  <fctr>    <int> <int>
1  crew         0   673
2  crew         1   212
3 first        0   122
4 first        1   203
5 second       0   167
6 second       1   118
7 third        0   528
8 third        1   178
```

```
Or summarise(group_by(titanic, class, survived), count =
n())
```

Or graphically...

```
plot(factor(survived) ~ class, data = titanic)
```



Fitting GLMs in R: glm

```
tit.glm <- glm(survived ~ class, data=titanic, family=binomial)
```

Call:

```
glm(formula = survived ~ class, family = binomial, data = titanic)
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-1.3999	-0.7623	-0.7401	0.9702	1.6906

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-1.15516	0.07876	-14.667	< 2e-16 ***
classfirst	1.66434	0.13902	11.972	< 2e-16 ***
classecond	0.80785	0.14375	5.620	1.91e-08 ***
classtthird	0.06785	0.11711	0.579	0.562

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Interpreting logistic regression output

Parameter estimates (logit-scale)

(Intercept)	classfirst	classecond	classtthird
-1.15515905	1.66434399	0.80784987	0.06784632

We need to back-transform: apply *inverse logit*

Crew probability of survival:

```
plogis(coef(tit.glm)[1])
```

```
(Intercept)  
0.239548
```

Looking at the data, the proportion of crew who survived is

```
[1] 0.239548
```

Q: Probability of survival for 1st class passengers?

```
plogis(coef(tit.glm)[1] + coef(tit.glm)[2])
```

```
(Intercept)  
0.6246154
```

Needs to add intercept (baseline) to the parameter estimate. Again this value matches the data:

```
sum(titanic$survived[titanic$class == "first"]) /  
  nrow(titanic[titanic$class == "first", ])
```

```
[1] 0.6246154
```

Model interpretation using effects package

```
library(effects)  
allEffects(tit.glm)
```

```
model: survived ~ class
```

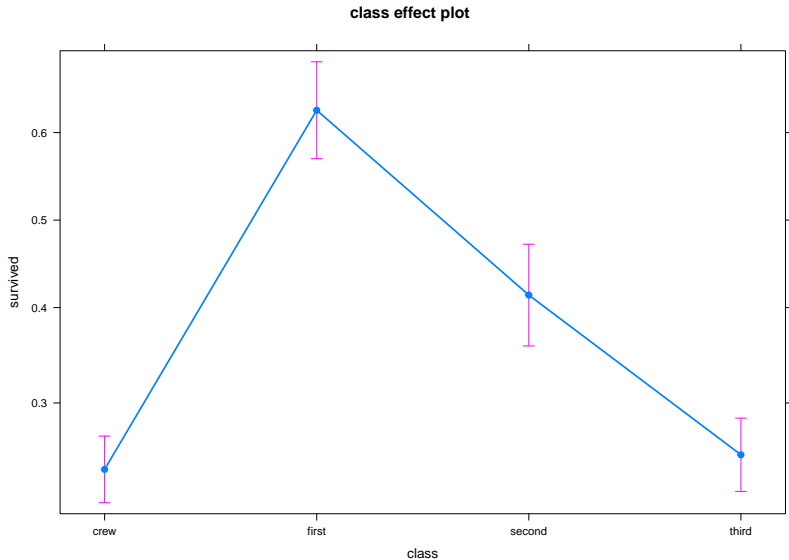
```
class effect
```

```
class
```

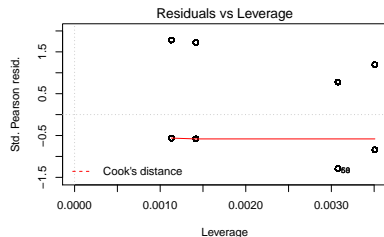
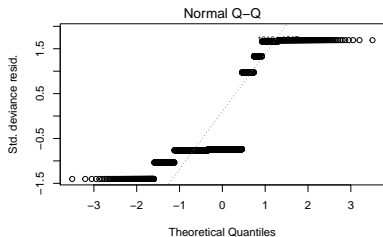
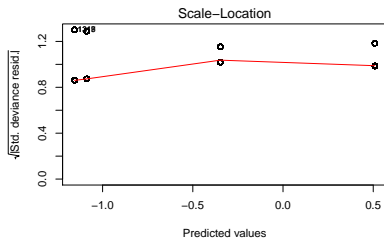
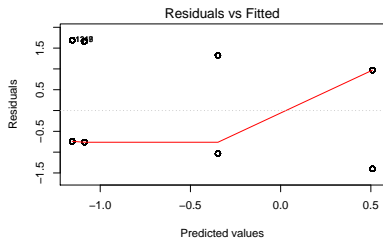
	crew	first	second	third
	0.2395480	0.6246154	0.4140351	0.2521246

Effects plot

```
plot(allEffects(tit.glm))
```



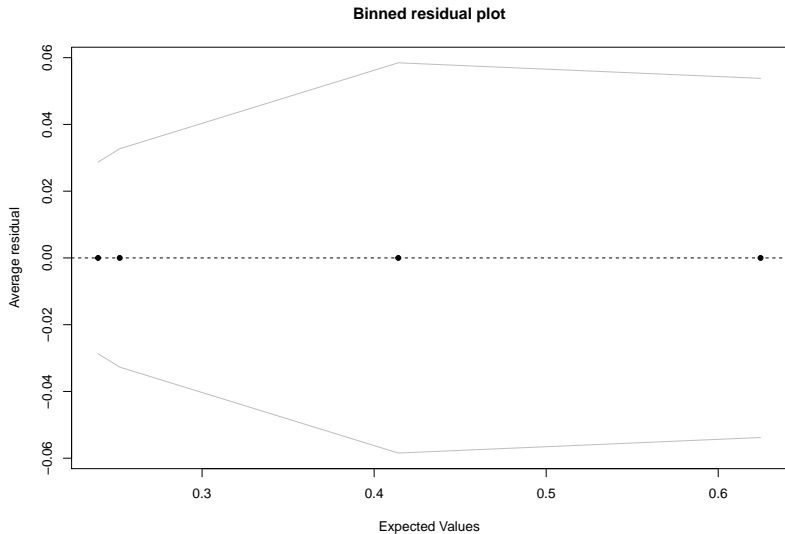
Logistic regression: model checking



Not very useful.

Binned residual plots for logistic regression

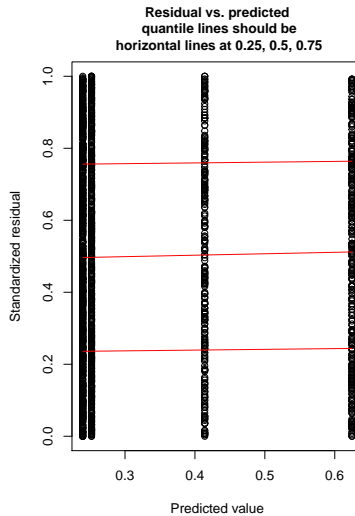
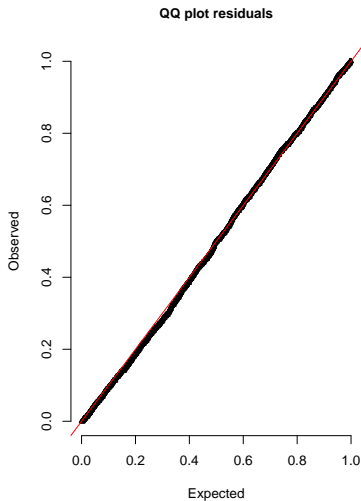
```
predvals <- predict(tit.glm, type="response")  
arm::binnedplot(predvals, titanic$survived - predvals)
```



Residual diagnostics with DHARMA

```
library(DHARMA)
simulateResiduals(tit.glm, plot = TRUE)
```

DHARMA scaled residual plots



Recapitulating

1. Import data: `read.table` or `read.csv`

Recapitulating

1. Import data: `read.table` or `read.csv`
2. Check data: `summary`

Recapitulating

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Recapitulating

1. Import data: `read.table` or `read.csv`
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4. Fit model: `glm`. Don't forget to specify `family`!

Recapitulating

1. Import data: `read.table` or `read.csv`
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5. Examine models: `summary`

Recapitulating

1. Import data: `read.table` or `read.csv`
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5. Examine models: `summary`
6. Use `plogis` to apply back-transformation (*invlogit*) to parameter estimates (`coef`). Alternatively, use `allEffects` from `effects` package.

Recapitulating

1. Import data: `read.table` or `read.csv`
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5. Examine models: `summary`
6. Use `plogis` to apply back-transformation (*invlogit*) to parameter estimates (`coef`). Alternatively, use `allEffects` from `effects` package.
7. Plot model: `plot(allEffects(model))`. Or use `visreg`.

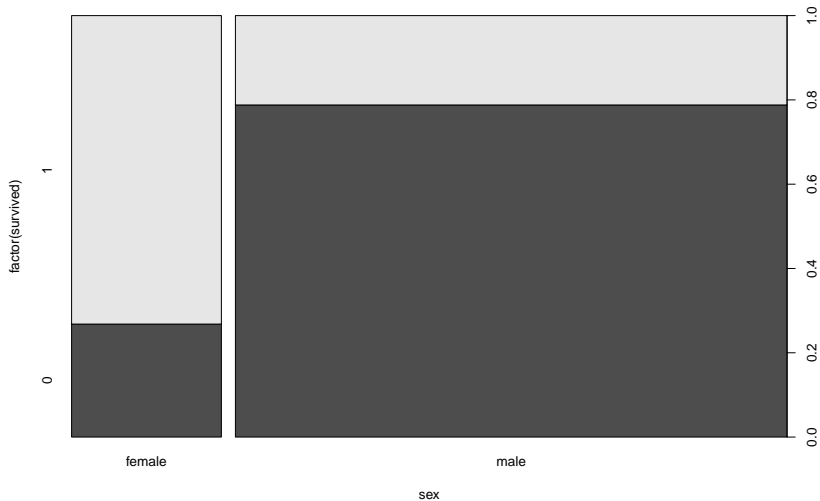
Recapitulating

1. Import data: `read.table` or `read.csv`
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4. Fit model: `glm`. Don't forget to specify family!
5. Examine models: `summary`
6. Use `plogis` to apply back-transformation (*invlogit*) to parameter estimates (`coef`). Alternatively, use `allEffects` from `effects` package.
7. Plot model: `plot(allEffects(model))`. Or use `visreg`.
8. Examine residuals: use `arm::binnedplot` or `DHARMA::simulateResiduals`.

Q: Did men have higher survival than women?

Plot first

```
plot(factor(survived) ~ sex, data = titanic)
```



Fit model

```
tit.sex <- glm(survived ~ sex, data = titanic, family = binomial)
```

Call:

```
glm(formula = survived ~ sex, family = binomial, data = titanic)
```

Deviance Residuals:

	Min	1Q	Median	3Q	Max
	-1.6226	-0.6903	-0.6903	0.7901	1.7613

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	1.0044	0.1041	9.645	<2e-16 ***
sexmale	-2.3172	0.1196	-19.376	<2e-16 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 2769.5 on 2200 degrees of freedom

Effects

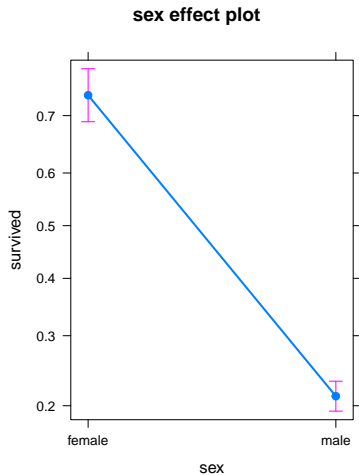
```
model: survived ~ sex
```

```
sex effect
```

```
sex
```

```
female    male
```

```
0.7319149 0.2120162
```



Q: Did women have higher survival because they travelled more in first class?

Let's look at the data

tapply

```
tapply(titanic$survived, list(titanic$class, titanic$sex), sum)
```

	female	male
crew	20	192
first	141	62
second	93	25
third	90	88

Mmmm...

Fit model with both factors (interactions)

```
tit.sex.class <- glm(survived ~ class * sex, data = titanic, fam
```

```
glm(formula = survived ~ class * sex, family = binomial, data =
```

```
      coef.est coef.se
```

```
(Intercept)      1.90    0.62
```

```
classfirst       1.67    0.80
```

```
classecond       0.07    0.69
```

```
classthird      -2.06    0.64
```

```
sexmale         -3.15    0.62
```

```
classfirst:sexmale -1.06    0.82
```

```
classecond:sexmale -0.64    0.72
```

```
classthird:sexmale  1.74    0.65
```

```
---
```

```
  n = 2201, k = 8
```

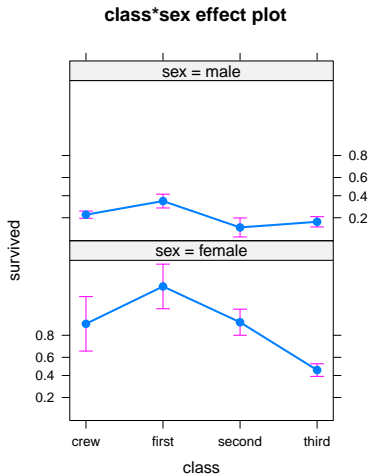
```
  residual deviance = 2163.7, null deviance = 2769.5 (difference
```

Effects

```
model: survived ~ class * sex
```

```
class*sex effect
```

	sex	
class	female	male
crew	0.8695652	0.2227378
first	0.9724138	0.3444444
second	0.8773585	0.1396648
third	0.4591837	0.1725490



So, women had higher probability of survival than men, even within the same class.

Logistic regression for proportion data

Read Titanic data in different format

Read Titanic_prop.csv data.

X	Class	Sex	Age	No
Min. : 1.00	1st :4	Female:8	Adult:8	Min. : 0.00
1st Qu.: 4.75	2nd :4	Male :8	Child:8	1st Qu.: 0.00
Median : 8.50	3rd :4			Median : 8.50
Mean : 8.50	Crew:4			Mean : 93.12
3rd Qu.:12.25				3rd Qu.: 96.25
Max. :16.00				Max. :670.00

Yes
Min. : 0.00
1st Qu.: 9.50
Median : 14.00
Mean : 44.44
3rd Qu.: 75.25
Max. :192.00

These are the same data, but summarized (see Freq variable).

Use cbind(n.success, n.failures) as response

```
prop.glm <- glm(cbind(Yes, No) ~ Class, data = tit.prop, family
```

Call:

```
glm(formula = cbind(Yes, No) ~ Class, family = binomial, data =
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-9.6404	-0.2915	1.5698	5.0366	10.1516

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	0.5092	0.1146	4.445	8.79e-06 ***
Class2nd	-0.8565	0.1661	-5.157	2.51e-07 ***
Class3rd	-1.5965	0.1436	-11.114	< 2e-16 ***
ClassCrew	-1.6643	0.1390	-11.972	< 2e-16 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Effects

```
model: cbind(Yes, No) ~ Class
```

```
Class effect
```

```
Class
```

	1st	2nd	3rd	Crew
	0.6246154	0.4140351	0.2521246	0.2395480

Compare with former model based on raw data:

```
model: survived ~ class
```

```
class effect
```

```
class
```

	crew	first	second	third
	0.2395480	0.6246154	0.4140351	0.2521246

Same results!

Logistic regression with continuous predictors

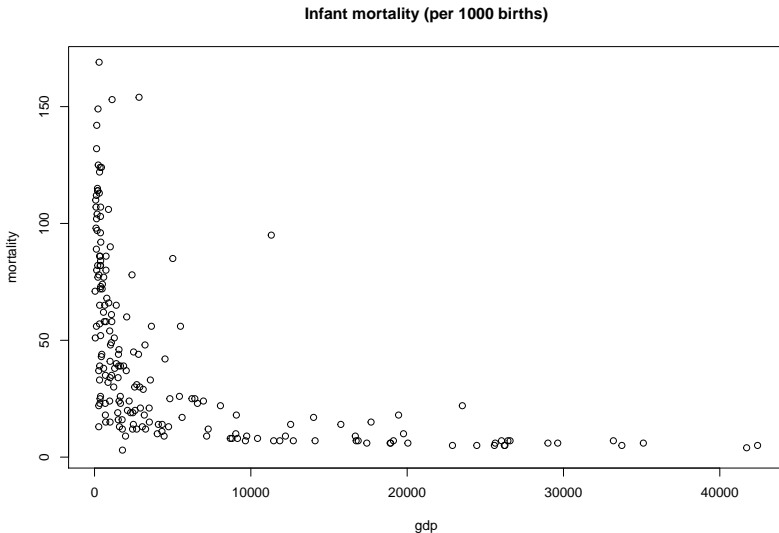
Example dataset: GDP and infant mortality

Read UN_GDP_infantmortality.csv.

	country	mortality	gdp
Afghanistan	: 1	Min. : 2.00	Min. : 36
Albania	: 1	1st Qu.: 12.00	1st Qu.: 442
Algeria	: 1	Median : 30.00	Median : 1779
American.Samoa	: 1	Mean : 43.48	Mean : 6262
Andorra	: 1	3rd Qu.: 66.00	3rd Qu.: 7272
Angola	: 1	Max. : 169.00	Max. : 42416
(Other)	: 201	NA's : 6	NA's : 10

EDA

```
plot(mortality ~ gdp, data = gdp, main = "Infant mortality (per
```



Fit model

```
gdp.glm <- glm(cbind(mortality, 1000 - mortality) ~ gdp,  
               data = gdp, family = binomial)
```

Call:

```
glm(formula = cbind(mortality, 1000 - mortality) ~ gdp, family =  
     data = gdp)
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-9.2230	-3.5163	-0.5697	2.4284	13.5849

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-2.657e+00	1.311e-02	-202.76	<2e-16 ***
gdp	-1.279e-04	3.458e-06	-36.98	<2e-16 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Effects

```
allEffects(gdp.glm)
```

```
model: cbind(mortality, 1000 - mortality) ~ gdp
```

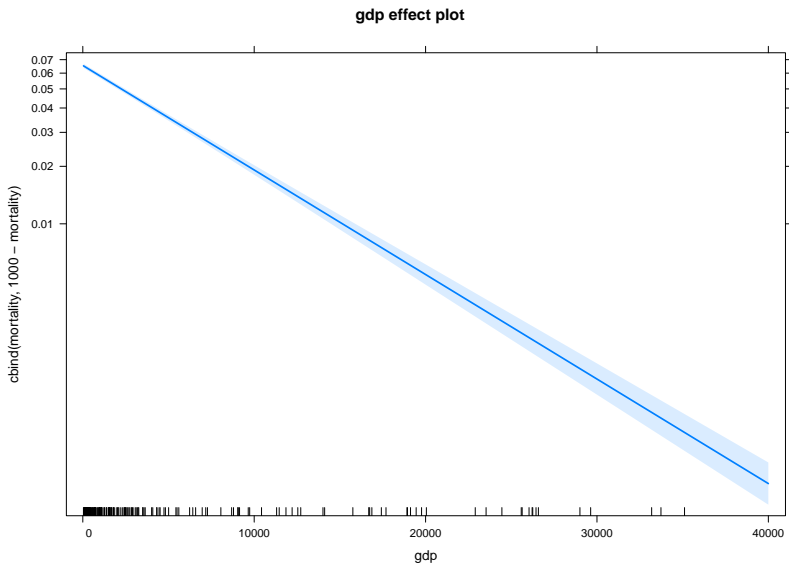
```
gdp effect
```

```
gdp
```

	40	10000	20000	30000	40000
	0.0652177296	0.0191438829	0.0054028095	0.0015096074	0.0004206154

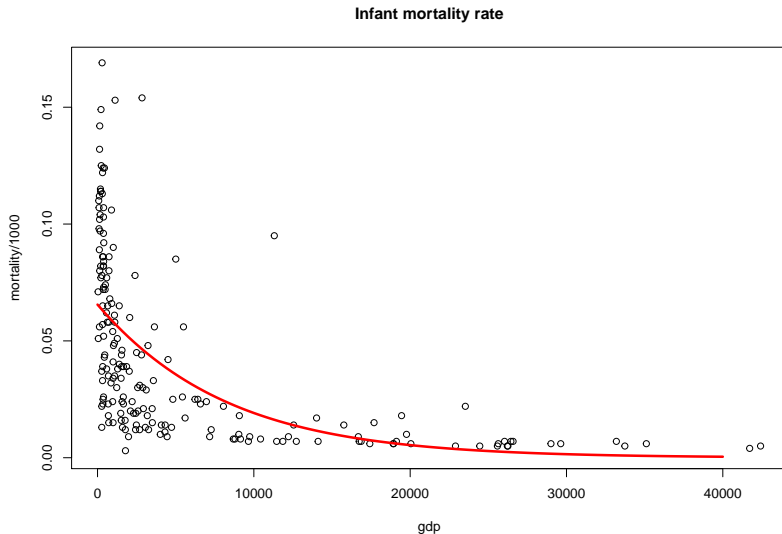
Effects plot

```
plot(allEffects(gdp.glm))
```



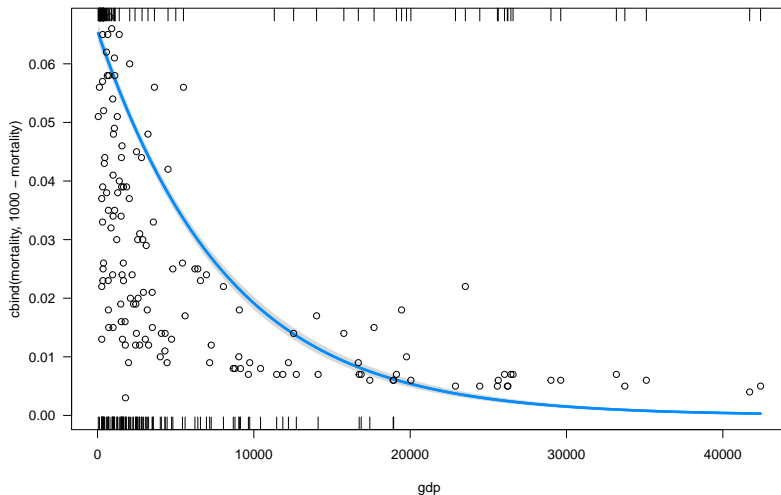
Plot model and data

```
plot(mortality/1000 ~ gdp, data = gdp, main = "Infant mortality  
curve(plogis(coef(gdp.glm)[1] + coef(gdp.glm)[2]*x), from = 0, t
```



Plot model using visreg:

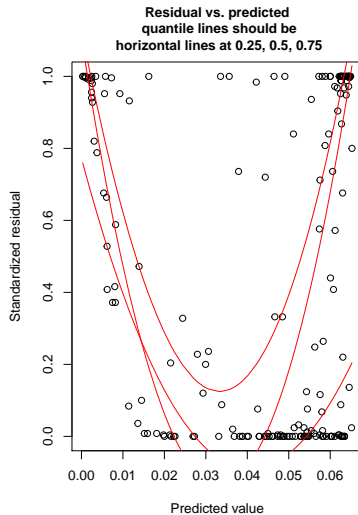
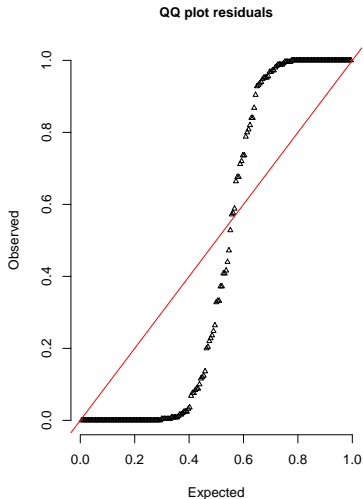
```
visreg(gdp.glm, scale = "response")  
points(mortality/1000 ~ gdp, data = gdp)
```



Residuals diagnostics with DHARMa

```
simulateResiduals(gdp.glm, plot = TRUE)
```

DHARMa scaled residual plots



Overdispersion

Testing for overdispersion (DHARMa)

```
simres <- simulateResiduals(gdp.glm, refit = TRUE)  
testOverdispersion(simres)
```

DHARMa nonparametric overdispersion test via comparison to simulation under H_0 = fitted model

```
data:  simres  
dispersion = 20.761, p-value < 2.2e-16  
alternative hypothesis: overdispersion
```

Overdispersion in logistic regression with proportion data

```
gdp.overdisp <- glm(cbind(mortality, 1000 - mortality) ~ gdp,  
                    data = gdp, family = quasibinomial)
```

Call:

```
glm(formula = cbind(mortality, 1000 - mortality) ~ gdp, family =  
    data = gdp)
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-9.2230	-3.5163	-0.5697	2.4284	13.5849

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	-2.657e+00	5.977e-02	-44.465	< 2e-16 ***
gdp	-1.279e-04	1.577e-05	-8.111	5.96e-14 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for quasibinomial family taken to be 20.79)

Mean estimates do not change after accounting for overdispersion

```
model: cbind(mortality, 1000 - mortality) ~ gdp
```

```
gdp effect
```

```
gdp
```

	40	10000	20000	30000	40000
	0.0652177296	0.0191438829	0.0054028095	0.0015096074	0.0004206154

```
model: cbind(mortality, 1000 - mortality) ~ gdp
```

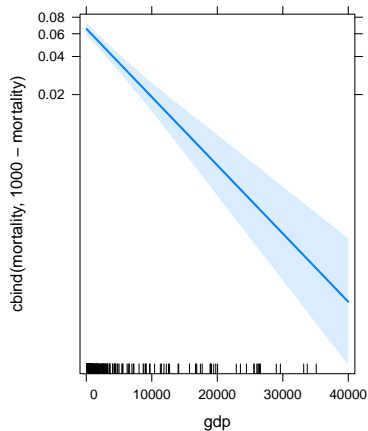
```
gdp effect
```

```
gdp
```

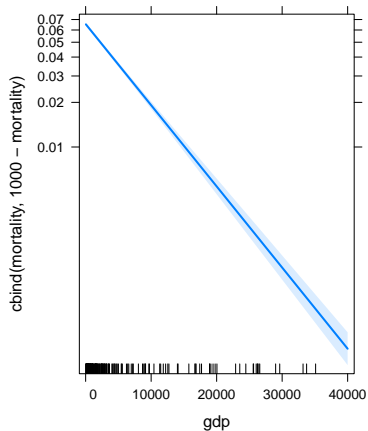
	40	10000	20000	30000	40000
	0.0652177296	0.0191438829	0.0054028095	0.0015096074	0.0004206154

But standard errors (uncertainty) do!

gdp effect plot

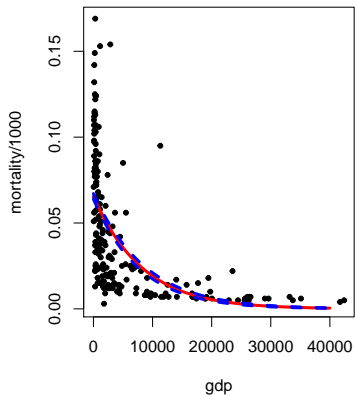


gdp effect plot

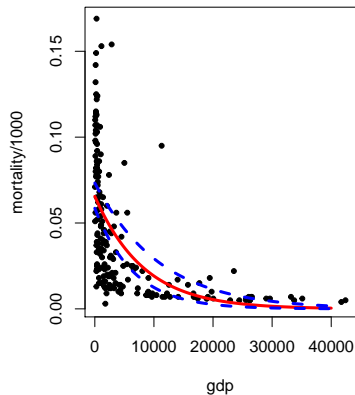


Plot model and data

Binomial



Quasibinomial



Overdispersion

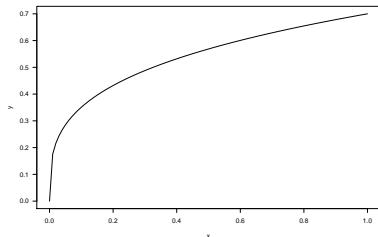
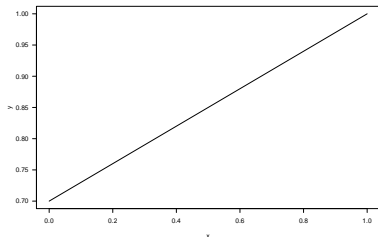
Whenever you fit logistic regression to **proportion** data, check family quasibinomial.

Think about the shape of relationships

$$y \sim x + z$$

Really? Not everything has to be linear! Actually, it often is not.

Think about shape of relationship. See chapter 3 in Bolker's book.



GLM for count data: Poisson regression

Types of response variable

- ▶ Gaussian: 1m

Types of response variable

- ▶ Gaussian: `lm`
- ▶ Bernoulli / Binomial: `glm (family binomial / quasibinomial)`

Types of response variable

- ▶ Gaussian: `lm`
- ▶ Bernoulli / Binomial: `glm (family binomial / quasibinomial)`
- ▶ Counts: `glm (family poisson / quasipoisson)`

Poisson regression

- ▶ Response variable: Counts (0, 1, 2, 3...) - discrete

Then

$$\log(N) = a + bx$$

$$N = e^{a+bx}$$

Poisson regression

- ▶ Response variable: Counts (0, 1, 2, 3...) - discrete
- ▶ Link function: \log

Then

$$\log(N) = a + bx$$

$$N = e^{a+bx}$$

Example dataset: Seedling counts in quadrats

```
seedl <- read.csv("data-row/seedlings.csv")
```

X	count	row	col
Min. : 1.00	Min. :0.00	Min. :1	Min. : 1.0
1st Qu.:13.25	1st Qu.:1.00	1st Qu.:2	1st Qu.: 3.0
Median :25.50	Median :2.00	Median :3	Median : 5.5
Mean :25.50	Mean :2.14	Mean :3	Mean : 5.5
3rd Qu.:37.75	3rd Qu.:3.00	3rd Qu.:4	3rd Qu.: 8.0
Max. :50.00	Max. :7.00	Max. :5	Max. :10.0

light	area
Min. : 2.571	Min. :0.25
1st Qu.:26.879	1st Qu.:0.25
Median :47.493	Median :0.50
Mean :47.959	Mean :0.62
3rd Qu.:67.522	3rd Qu.:1.00
Max. :99.135	Max. :1.00

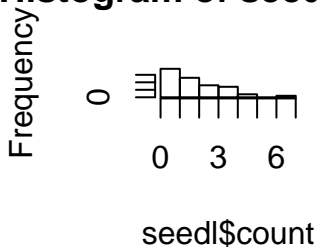
EDA

```
table(seed1$count)
```

```
0  1  2  3  4  5  7  
7 12 13  8  7  2  1
```

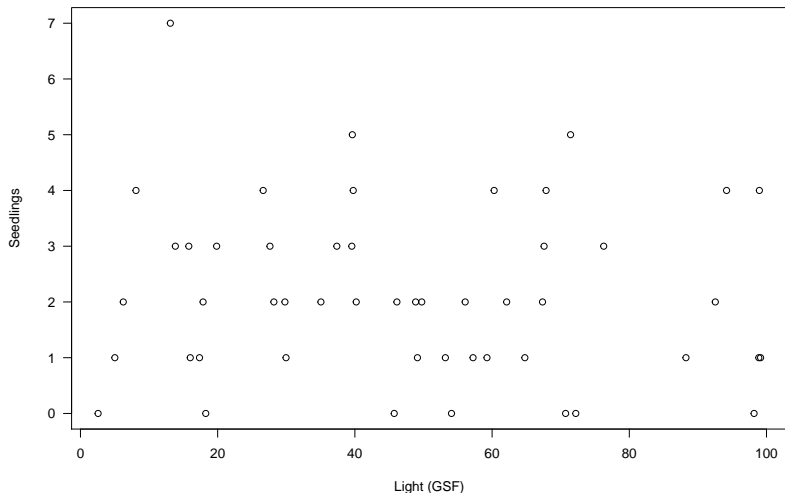
```
hist(seed1$count)
```

Histogram of seed1\$count



Q: Relationship between Nseedlings and light?

```
plot(seedl$light, seedl$count, las = 1, xlab = "Light (GSF)", ylab = "Seedlings")
```



Let's fit model (Poisson regression)

```
seedl.glm <- glm(count ~ light, data = seedl, family = poisson)
summary(seedl.glm)
```

Call:

```
glm(formula = count ~ light, family = poisson, data = seedl)
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-2.1906	-0.8466	-0.1110	0.5220	2.4577

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	0.881805	0.188892	4.668	3.04e-06 ***
light	-0.002576	0.003528	-0.730	0.465

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for poisson family taken to be 1)

Interpreting Poisson regression output

Parameter estimates (log scale):

```
coef(seedl.glm)
```

(Intercept)	light
0.881805022	-0.002575656

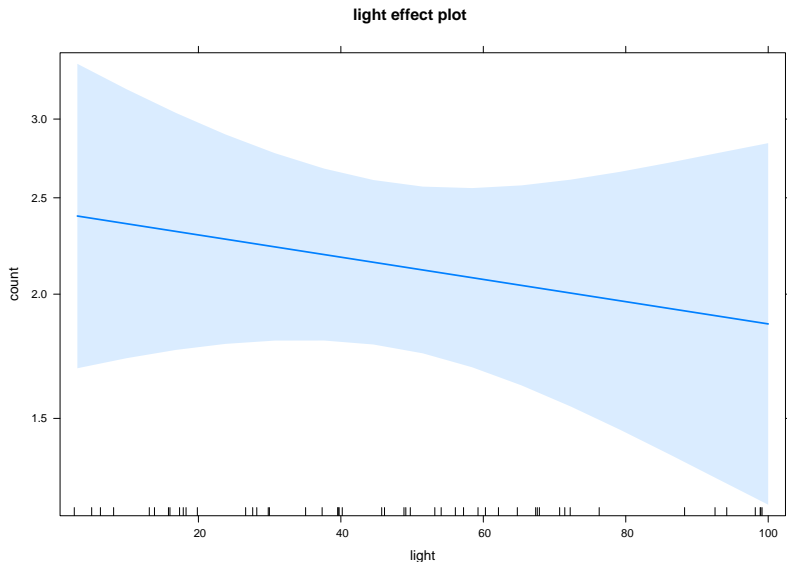
We need to back-transform: apply the inverse of the logarithm

```
exp(coef(seedl.glm))
```

(Intercept)	light
2.4152554	0.9974277

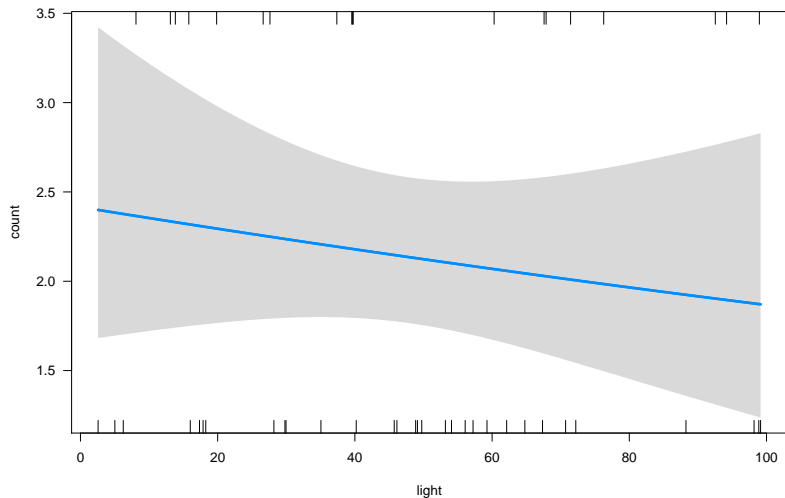
So what's the relationship between Nseedlings and light?

```
plot(allEffects(seed1.glm))
```

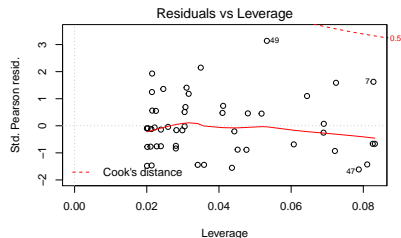
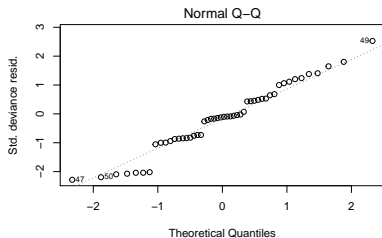
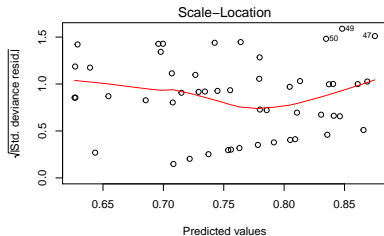
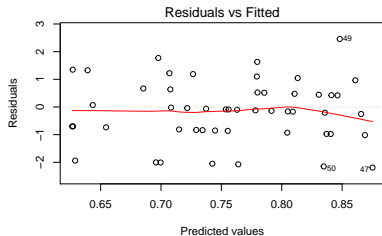


Using visreg

```
visreg(seed1.glm, scale = "response")
```

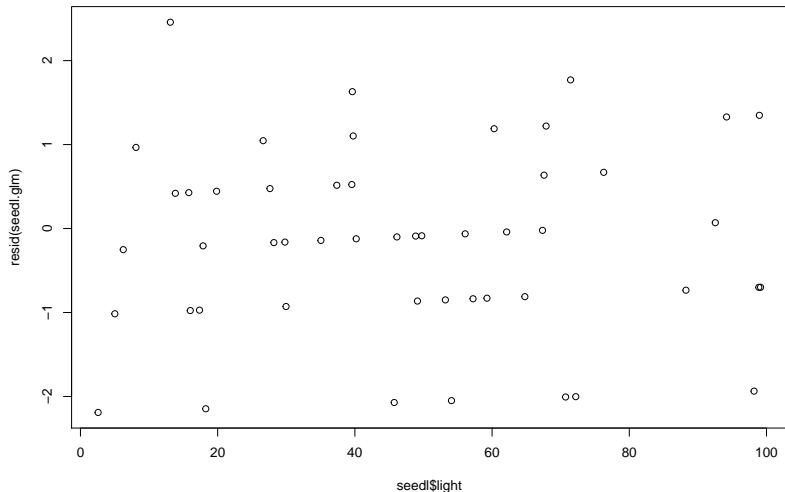


Poisson regression: model checking



Is there pattern of residuals along predictor?

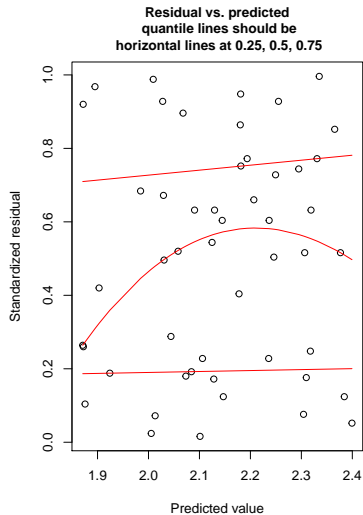
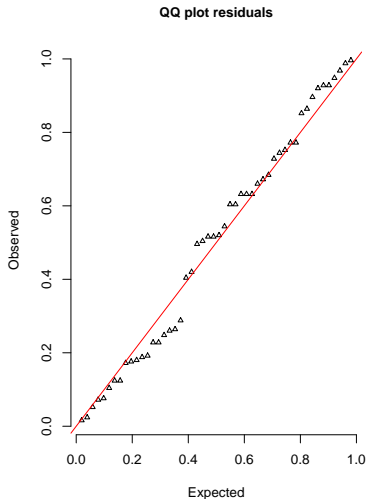
```
plot(seedl$light, resid(seedl.glm))
```



Residuals diagnostics with DHARMA

```
simulateResiduals(seed1.glm, plot = TRUE)
```

DHARMA scaled residual plots



Poisson regression: Overdispersion

Always check overdispersion with count data

```
simres <- simulateResiduals(seed1.glm, refit = TRUE)  
testOverdispersion(simres)
```

DHARMA nonparametric overdispersion test via comparison to
simulation under H_0 = fitted model

```
data:  simres  
dispersion = 1.1574, p-value = 0.196  
alternative hypothesis: overdispersion
```

Accounting for overdispersion in count data

Use family quasipoisson

Call:

```
glm(formula = count ~ light, family = quasipoisson, data = seedl
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-2.1906	-0.8466	-0.1110	0.5220	2.4577

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	0.881805	0.201230	4.382	6.37e-05 ***
light	-0.002576	0.003758	-0.685	0.496

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for quasipoisson family taken to be 1.1349

Null deviance:	63.029	on 49	degrees of freedom
Residual deviance:	62.492	on 48	degrees of freedom

Mean estimates do not change after accounting for overdispersion

```
model: count ~ light
```

```
light effect
```

```
light
```

	3	30	50	70	100
	2.396665	2.235657	2.123408	2.016794	1.866826

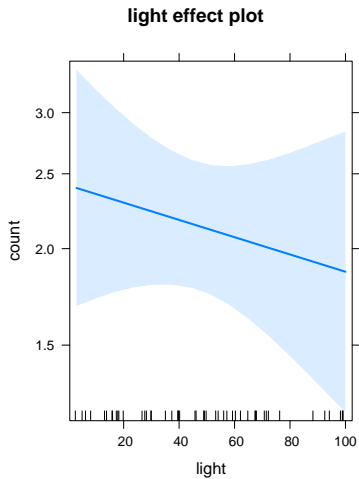
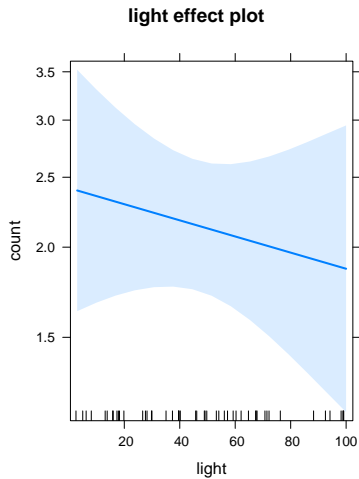
```
model: count ~ light
```

```
light effect
```

```
light
```

	3	30	50	70	100
	2.396665	2.235657	2.123408	2.016794	1.866826

But standard errors may change



What if survey plots have different area?

Avoid regression of ratios

seedlings/area \sim light

J. R. Statist. Soc. A (1993)
156, Part 3, pp. 379–392

Spurious Correlation and the Fallacy of the Ratio Standard Revisited

By RICHARD A. KRONMAL†

Figure 6:

Use offset to standardise response variables in GLMs

```
seedl.offset <- glm(count ~ light, offset = seedl$area, data = seedl, family = poisson)
summary(seedl.offset)
```

Call:

```
glm(formula = count ~ light, family = poisson, data = seedl,
     offset = seedl$area)
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-2.6926	-0.8532	0.1491	0.5211	3.1051

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	0.299469	0.185468	1.615	0.106
light	-0.004498	0.003441	-1.307	0.191

(Dispersion parameter for poisson family taken to be 1)

Null deviance: 70.263 on 49 degrees of freedom

Note estimates now referred to area units

```
exp(coef(seedl.offset))
```

(Intercept)	light
1.3491422	0.9955123

Mixed / Multilevel models

Example dataset: trees

- Data on 1000 trees from 10 plots.

```
head(trees)
```

	plot	dbh	height	sex	dead	dbh.c
1	2	38.85	37.8	female	0	13.85
2	4	26.05	38.1	female	0	1.05
3	5	42.66	50.2	female	0	17.66
4	2	20.72	30.1	female	0	-4.28
5	4	21.83	34.0	female	0	-3.17
6	4	8.23	21.9	male	0	-16.77

Example dataset: trees

- ▶ Data on 1000 trees from 10 plots.
- ▶ Trees per plot: 4 - 392.

```
head(trees)
```

	plot	dbh	height	sex	dead	dbh.c
1	2	38.85	37.8	female	0	13.85
2	4	26.05	38.1	female	0	1.05
3	5	42.66	50.2	female	0	17.66
4	2	20.72	30.1	female	0	-4.28
5	4	21.83	34.0	female	0	-3.17
6	4	8.23	21.9	male	0	-16.77

Q: What's the relationship between tree diameter and height?

A simple linear model

```
lm.simple <- lm(height ~ dbh, data = trees)
```

Call:

```
lm(formula = height ~ dbh, data = trees)
```

Residuals:

Min	1Q	Median	3Q	Max
-13.7384	-4.7652	0.4759	4.2931	13.5282

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	13.18767	0.41476	31.80	<2e-16 ***
dbh	0.60967	0.01351	45.14	<2e-16 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 5.549 on 998 degrees of freedom

Multiple R-squared: 0.6712, Adjusted R-squared: 0.6709

F-statistic: 2038 on 1 and 998 DF, p-value: < 2.2e-16

Remember our model structure

$$y_i \sim N(\mu_i, \sigma^2)$$

$$\mu_i = \alpha + \beta x_i$$

In this case:

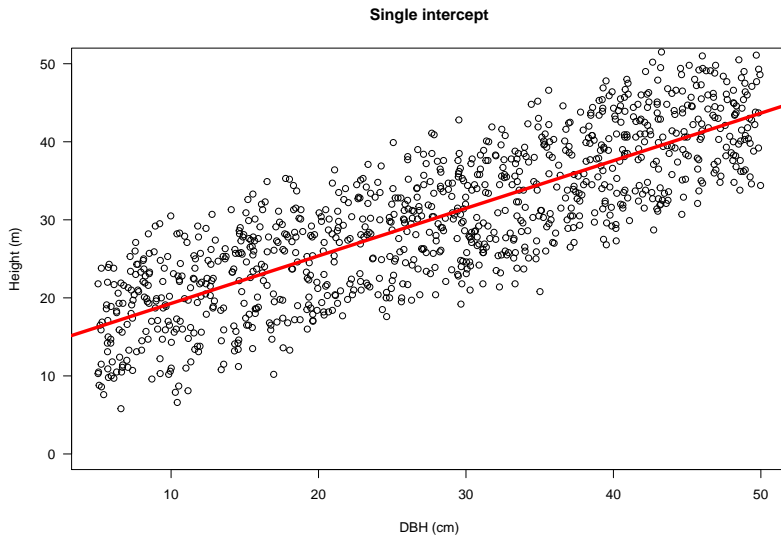
$$Height_i \sim N(\mu_i, \sigma^2)$$

$$\mu_i = \alpha + \beta DBH_i$$

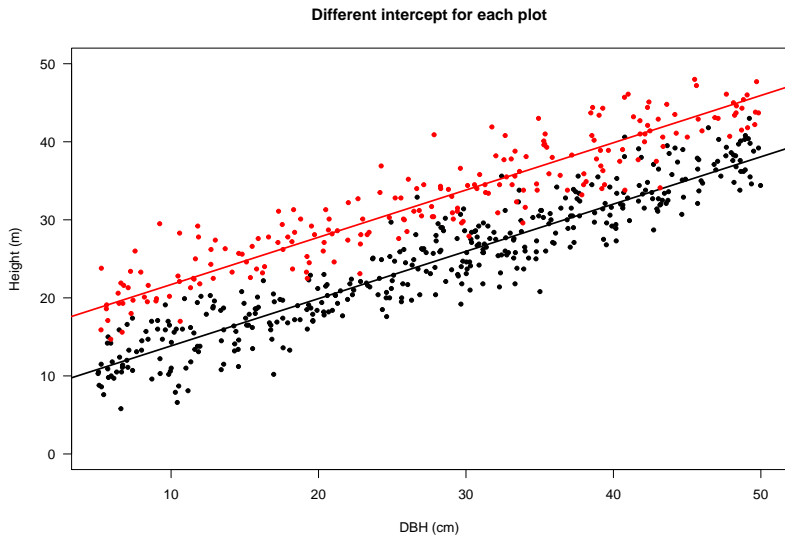
α : expected height when $DBH = 0$

β : how much height increases with every unit increase of DBH

There is only one intercept



What if allometry varies among plots?



Fitting a varying intercepts model with `lm`

```
lm(formula = height ~ factor(plot) + dbh, data = trees)
```

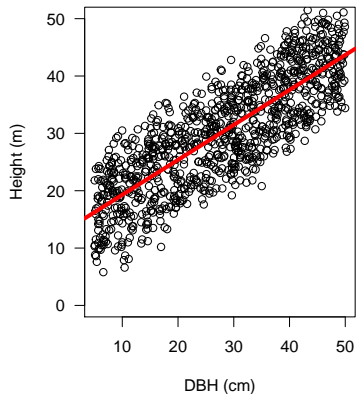
	coef.est	coef.se
(Intercept)	7.79	0.24
factor(plot)2	7.86	0.24
factor(plot)3	7.95	0.32
factor(plot)4	11.48	0.33
factor(plot)5	11.05	0.32
factor(plot)6	11.55	0.43
factor(plot)7	7.41	0.63
factor(plot)8	3.05	0.97
factor(plot)9	9.73	1.45
factor(plot)10	-0.14	0.92
dbh	0.61	0.01

n = 1000, k = 11

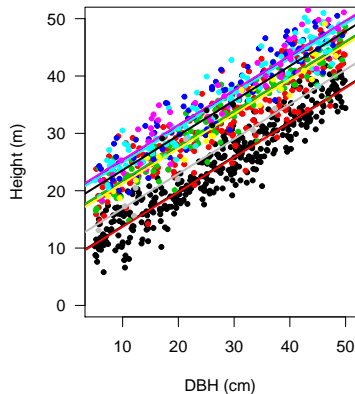
residual sd = 2.89, R-Squared = 0.91

Single vs varying intercept

Pooling all plots



Different intercept for each plot



Mixed models enable us to account for variability

► Varying intercepts

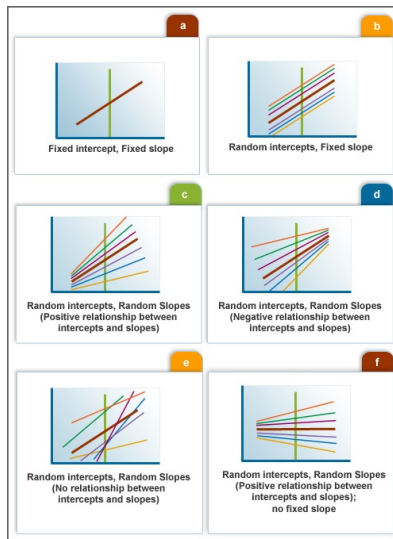


Figure 7:

Mixed models enable us to account for variability

- ▶ Varying intercepts
- ▶ Varying slopes

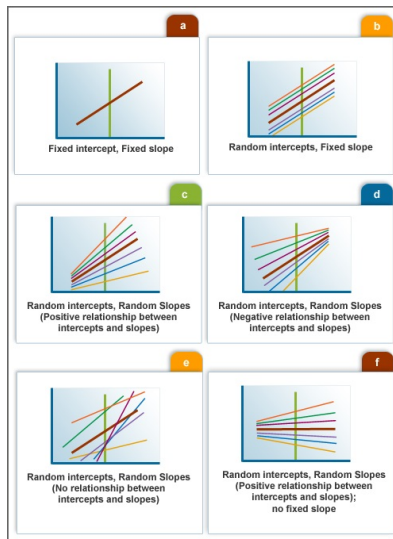


Figure 7:

Mixed model with varying intercepts

$$y_i = a_j + bx_i + \varepsilon_i$$

$$a_j \sim N(0, \tau^2)$$

$$\varepsilon_i \sim N(0, \sigma^2)$$

En nuestro ejemplo:

$$\text{Height}_i = \text{plot}_j + b\text{DBH}_i + \varepsilon_i$$

$$\text{plot}_j \sim N(0, \tau^2)$$

$$\varepsilon_i \sim N(0, \sigma^2)$$

Mixed models estimate varying parameters (intercepts and/or slopes) with pooling among levels (rather than considering them fully independent)

Hence there's gradient between

- ▶ **complete pooling**: Single overall intercept.

Hence there's gradient between

- ▶ **complete pooling:** Single overall intercept.
 - ▶ `lm (height ~ dbh)`

Hence there's gradient between

- ▶ **complete pooling:** Single overall intercept.
 - ▶ `lm (height ~ dbh)`
- ▶ **no pooling:** One *independent* intercept for each plot.

Hence there's gradient between

- ▶ **complete pooling:** Single overall intercept.
 - ▶ `lm (height ~ dbh)`
- ▶ **no pooling:** One *independent* intercept for each plot.
 - ▶ `lm (height ~ dbh + factor(plot))`

Hence there's gradient between

- ▶ **complete pooling**: Single overall intercept.
 - ▶ `lm (height ~ dbh)`
- ▶ **no pooling**: One *independent* intercept for each plot.
 - ▶ `lm (height ~ dbh + factor(plot))`
- ▶ **partial pooling**: Inter-related intercepts.

Hence there's gradient between

- ▶ **complete pooling**: Single overall intercept.
 - ▶ `lm (height ~ dbh)`
- ▶ **no pooling**: One *independent* intercept for each plot.
 - ▶ `lm (height ~ dbh + factor(plot))`
- ▶ **partial pooling**: Inter-related intercepts.
 - ▶ `lmer(height ~ dbh + (1 | plot))`

Random vs Fixed effects?

1. Fixed effects constant across individuals, random effects vary.

http://andrewgelman.com/2005/01/25/why_i_dont_use/

Random vs Fixed effects?

1. Fixed effects constant across individuals, random effects vary.
2. Effects are fixed if they are interesting in themselves; random if interest in the underlying population.

http://andrewgelman.com/2005/01/25/why_i_dont_use/

Random vs Fixed effects?

1. Fixed effects constant across individuals, random effects vary.
2. Effects are fixed if they are interesting in themselves; random if interest in the underlying population.
3. Fixed when sample exhausts the population; random when the sample is small part of the population.

http://andrewgelman.com/2005/01/25/why_i_dont_use/

Random vs Fixed effects?

1. Fixed effects constant across individuals, random effects vary.
2. Effects are fixed if they are interesting in themselves; random if interest in the underlying population.
3. Fixed when sample exhausts the population; random when the sample is small part of the population.
4. Random effect if it's assumed to be a realized value of random variable.

http://andrewgelman.com/2005/01/25/why_i_dont_use/

Random vs Fixed effects?

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2. Effects are fixed if they are interesting in themselves; random if interest in the underlying population.
3. Fixed when sample exhausts the population; random when the sample is small part of the population.
4. Random effect if it's assumed to be a realized value of random variable.
5. Fixed effects estimated using least squares or maximum likelihood; random effects estimated with shrinkage.

http://andrewgelman.com/2005/01/25/why_i_dont_use/

What is a random effect, really?

1. Varies by group

Random effects are estimated with partial pooling, while fixed effects are not (infinite variance).

What is a random effect, really?

1. Varies by group
2. Variation estimated with probability model

Random effects are estimated with partial pooling, while fixed effects are not (infinite variance).

Shrinkage improves parameter estimation

Especially for groups with low sample size

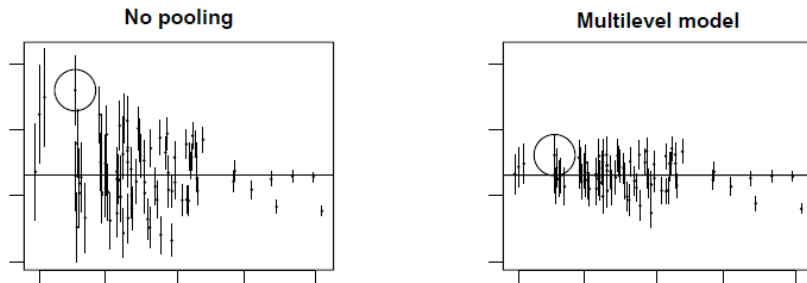


Figure 8:

From Gelman & Hill p. 253

Fitting mixed/multilevel models

```
library(lme4)
mixed <- lmer(height ~ dbh + (1|plot), data = trees)
```

```
lmer(formula = height ~ dbh + (1 | plot), data = trees)
      coef.est coef.se
(Intercept) 14.80    1.44
dbh          0.61    0.01
```

Error terms:

Groups	Name	Std.Dev.
plot	(Intercept)	4.45
Residual		2.89

number of obs: 1000, groups: plot, 10
AIC = 5015.6, DIC = 4996.4
deviance = 5002.0

Retrieve model coefficients

```
coef(mixed)
```

```
$plot
```

	(Intercept)	dbh
1	7.798373	0.6056549
2	15.647613	0.6056549
3	15.735397	0.6056549
4	19.253661	0.6056549
5	18.819467	0.6056549
6	19.306574	0.6056549
7	15.197908	0.6056549
8	11.016485	0.6056549
9	17.265447	0.6056549
10	7.940715	0.6056549

```
attr(,"class")
```

```
[1] "coef.mer"
```

Broom: model estimates in tidy form

```
library(broom)
tidy(mixed)
```

	term	estimate	std.error	statistic	gr
1	(Intercept)	14.7981641	1.437421280	10.29494	fi
2	dbh	0.6056549	0.007040079	86.02956	fi
3	sd_(Intercept).plot	4.4535702	NA	NA	p
4	sd_Observation.Residual	2.8852942	NA	NA	Resid

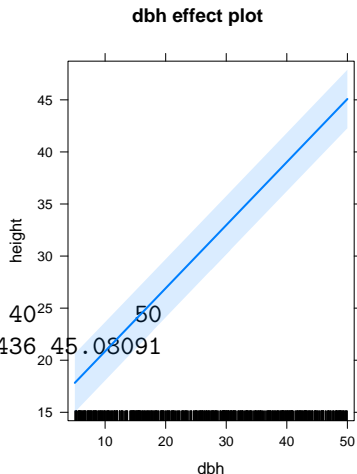
Visualising model: allEffects

```
model: height ~ dbh
```

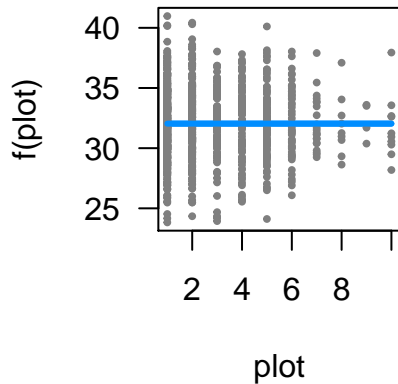
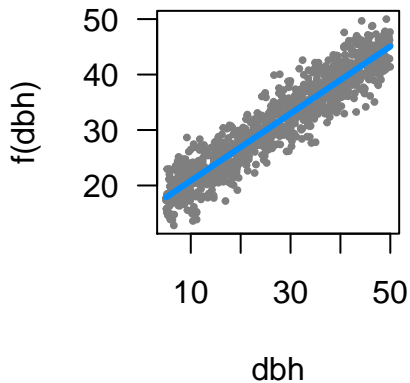
```
dbh effect
```

```
dbh
```

	5	20	30	40	50
	17.82644	26.91126	32.96781	39.02436	45.08091

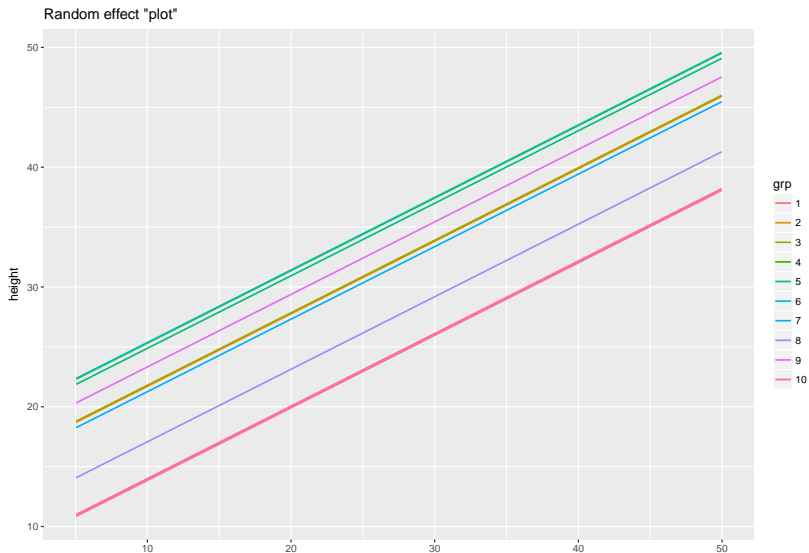


Visualising model: visreg



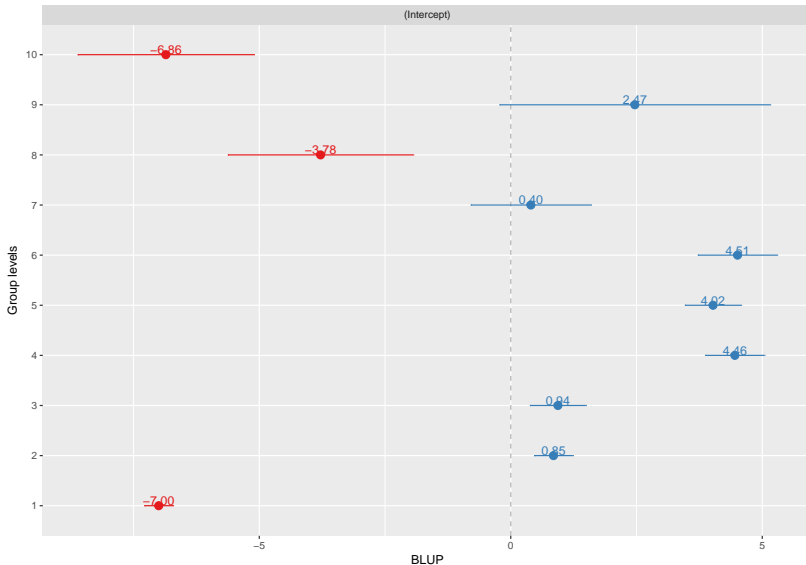
Visualising model: sjPlot

```
library(sjPlot)
sjp.lmer(mixed, type = "ri.slope")
```



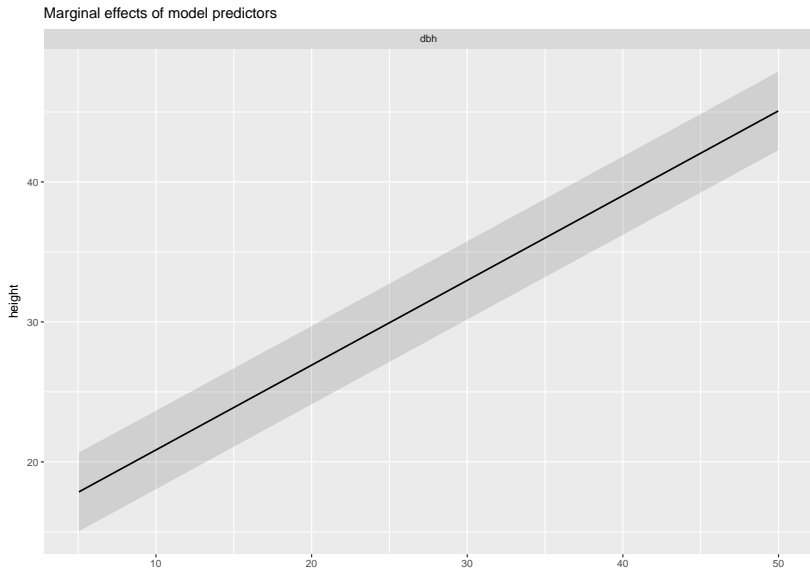
Visualising model: sjPlot

```
sjp.lmer(mixed)
```



Visualising model: sjPlot

```
sjp.lmer(mixed, type = "eff", show.ci = TRUE)
```

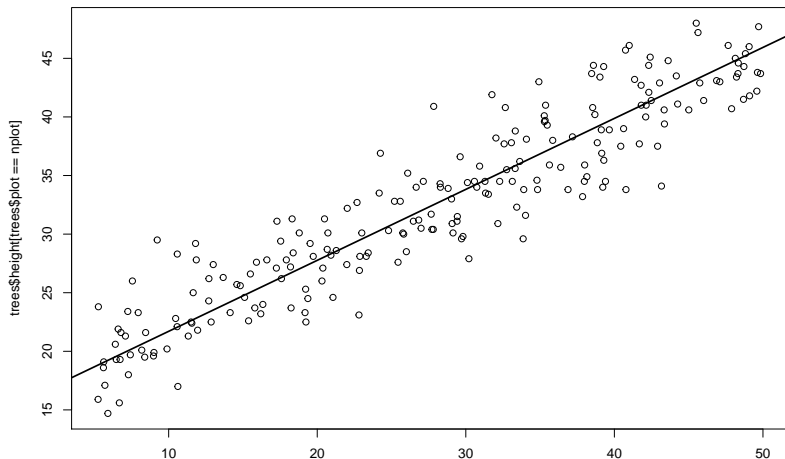


Using merTools to understand fitted model

```
library(merTools)
shinyMer(mixed)
```

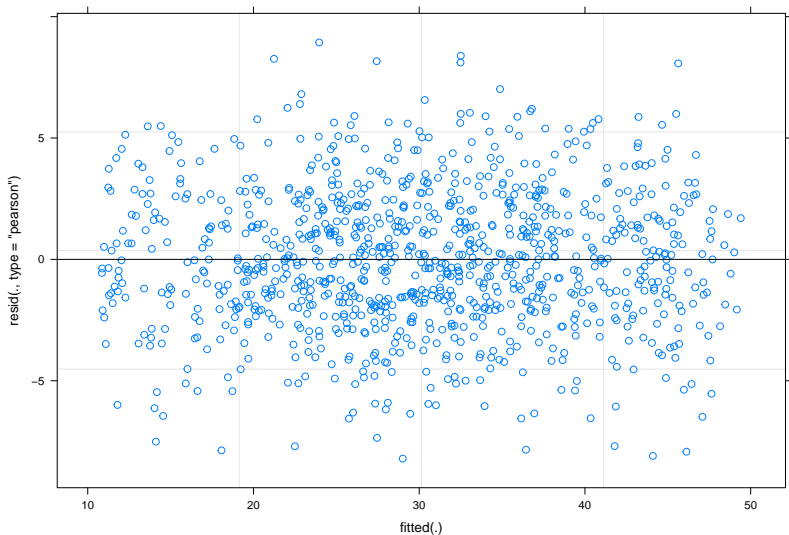
Plotting regression for individual forest plots

```
nplot <- 2  
plot(trees$dbh[trees$plot==nplot], trees$height[trees$plot==nplot],  
abline(a=coef(mixed)$plot[nplot, 1], b=coef(mixed)$plot[nplot, 2])
```



Checking residuals

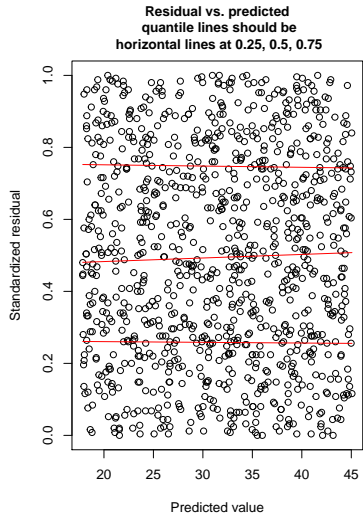
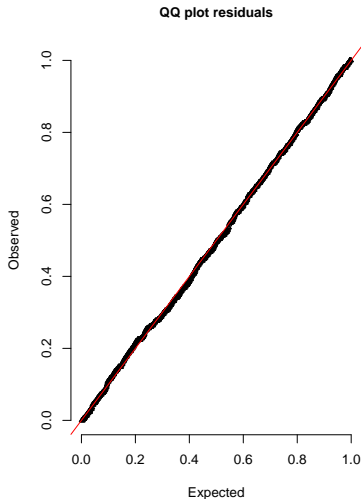
```
plot(mixed)
```



Checking residuals (DHARMA)

```
simulateResiduals(mixed, plot = TRUE, use.u = TRUE)
```

DHARMA scaled residual plots



Growing the hierarchy: adding plot-level
predictors

Model with group-level predictors

We had:

$$y_i = a_j + bx_i + \varepsilon_i$$

$$a_j \sim N(0, \tau^2)$$

$$\varepsilon_i \sim N(0, \sigma^2)$$

Now

$$y_i = a_j + bx_i + \varepsilon_i$$

$$a_j \sim N(\mu_j, \tau^2)$$

$$\mu_j = \gamma + \delta \cdot \text{predictor}_j$$

$$\varepsilon_i \sim N(0, \sigma^2)$$

Merging trees and plot data

```
plotdata <- read.csv("data-raw/plotdata.csv")
trees.full <- merge(trees, plotdata, by = "plot")
head(trees.full)
```

	plot	dbh	height	sex	dead	dbh.c	temp
1	1	28.63	22.1	female	0	3.63	15.1
2	1	44.71	39.0	female	0	19.71	15.1
3	1	28.31	29.0	female	0	3.31	15.1
4	1	19.33	19.1	male	0	-5.67	15.1
5	1	9.25	12.2	female	0	-15.75	15.1
6	1	30.02	23.1	female	0	5.02	15.1

Centre continuous variables

Plot temperatures referred as deviations from 15°C

```
trees.full$temp.c <- trees.full$temp - 15
```

Fit multilevel model

```
group.pred <- lmer(height ~ dbh + (1 | plot) + temp.c, data = tr  
arm::display(group.pred)
```

```
lmer(formula = height ~ dbh + (1 | plot) + temp.c, data = trees.
```

	coef.est	coef.se
(Intercept)	11.79	1.75
dbh	0.61	0.01
temp.c	1.07	0.46

Error terms:

Groups	Name	Std.Dev.
plot	(Intercept)	3.61
Residual		2.89

number of obs: 1000, groups: plot, 10

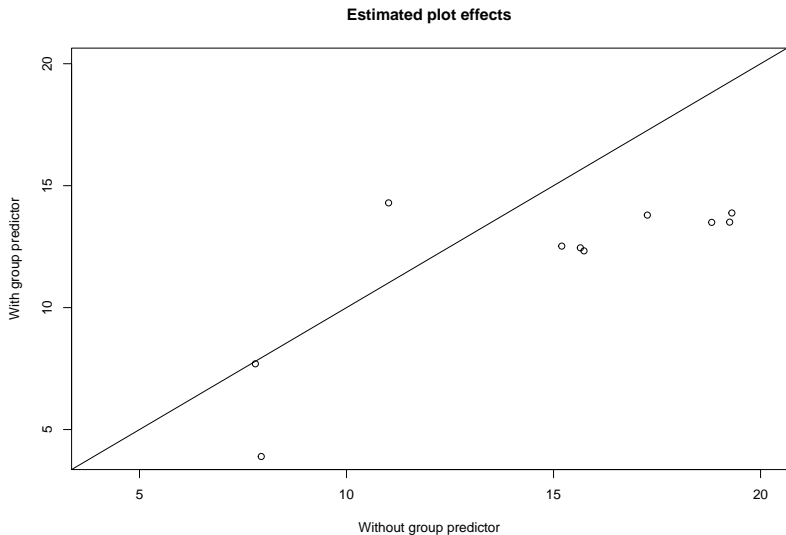
AIC = 5012.8, DIC = 4991

deviance = 4996.9

Examine model with merTools

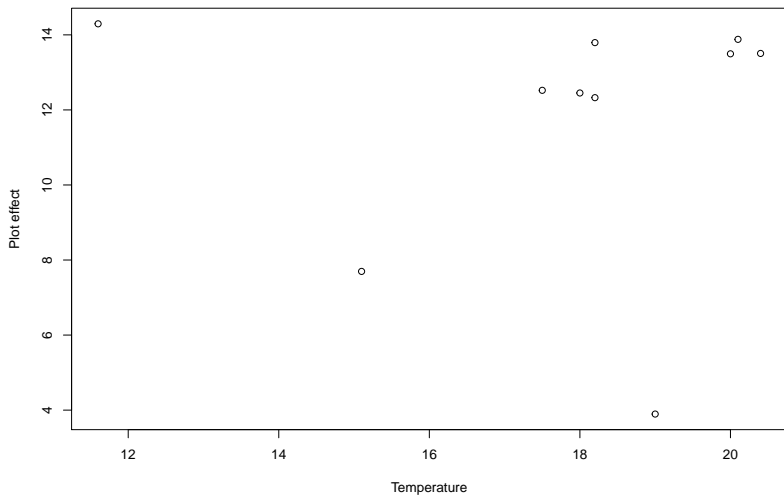
```
shinyMer(group.pred)
```

Comparing plot effects with and without group predictor



Are plot effects related to temperature?

```
plot(plotdata$temp, coef(group.pred)$plot[,1],  
      xlab = "Temperature", ylab = "Plot effect")
```



Varying intercepts and slopes

Varying intercepts and slopes

- ▶ There is overall difference in height among plots (different intercepts)

```
mixed.slopes <- lmer(height ~ dbh + (1 + dbh | plot), data=trees)
```

Varying intercepts and slopes

- ▶ There is overall difference in height among plots (different intercepts)
- ▶ AND

```
mixed.slopes <- lmer(height ~ dbh + (1 + dbh | plot), data=trees)
```


Varying intercepts and slopes

- ▶ There is overall difference in height among plots (different intercepts)
- ▶ AND
- ▶ Relationship between DBH and Height varies among plots (different slopes)

```
mixed.slopes <- lmer(height ~ dbh + (1 + dbh | plot), data=trees)
```

Varying intercepts and slopes

```
lmer(formula = height ~ dbh + (1 + dbh | plot), data = trees)
```

	coef.est	coef.se
(Intercept)	14.82	1.48
dbh	0.60	0.01

Error terms:

Groups	Name	Std.Dev.	Corr
plot	(Intercept)	4.57	
	dbh	0.01	-0.41
Residual		2.88	

number of obs: 1000, groups: plot, 10

AIC = 5018.6, DIC = 4995.9

deviance = 5001.3

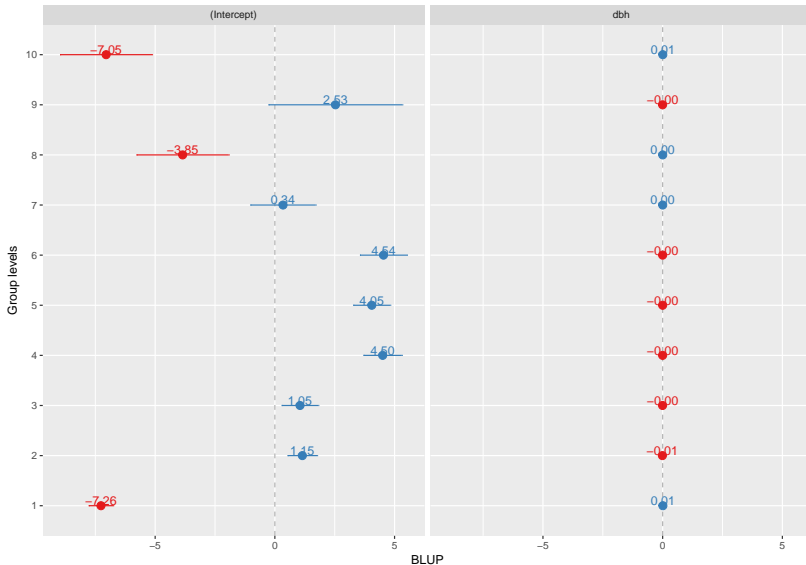
Varying intercepts and slopes

```
$plot
      (Intercept)      dbh
1      7.554578 0.6144452
2     15.966915 0.5942836
3     15.868969 0.6008673
4     19.321161 0.6031855
5     18.866370 0.6039353
6     19.355009 0.6038332
7     15.159258 0.6067449
8     10.965429 0.6080747
9     17.348840 0.6024600
10     7.769135 0.6109349
```

```
attr(,"class")
[1] "coef.mer"
```

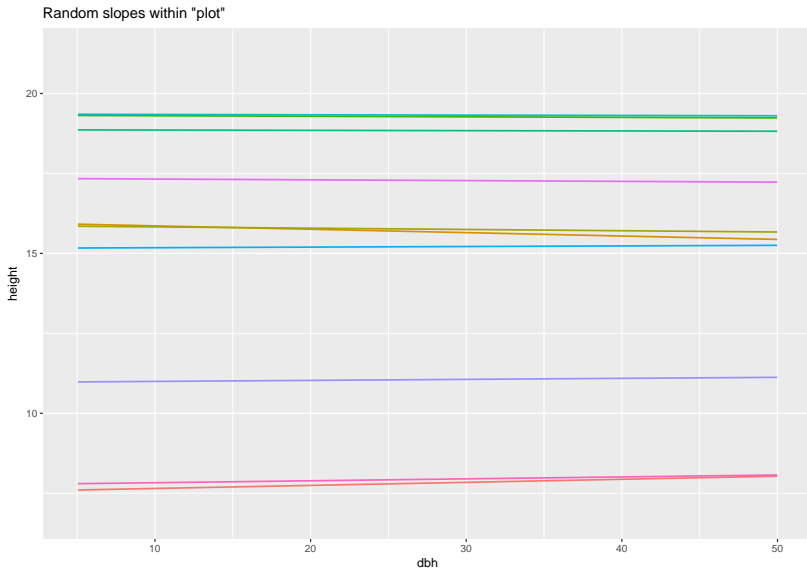
Visualising model: sjPlot

```
sjp.lmer(mixed.slopes)
```



Visualising model: sjPlot

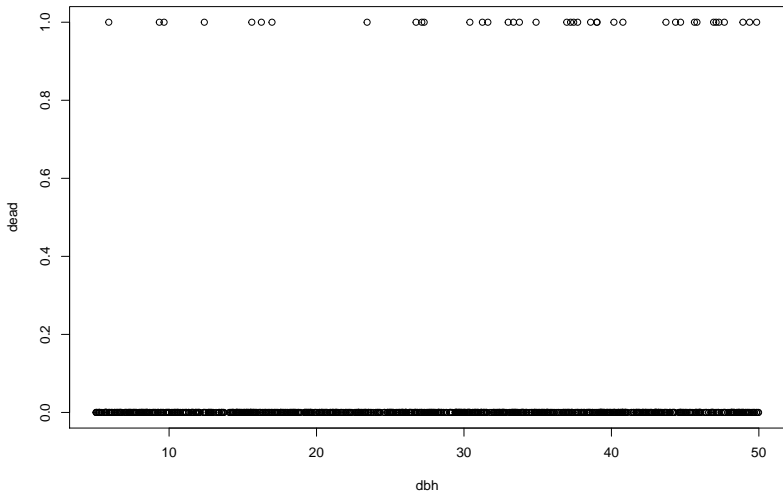
```
sjp.lmer(mixed.slopes, type = "rs.ri")
```



Multilevel logistic regression

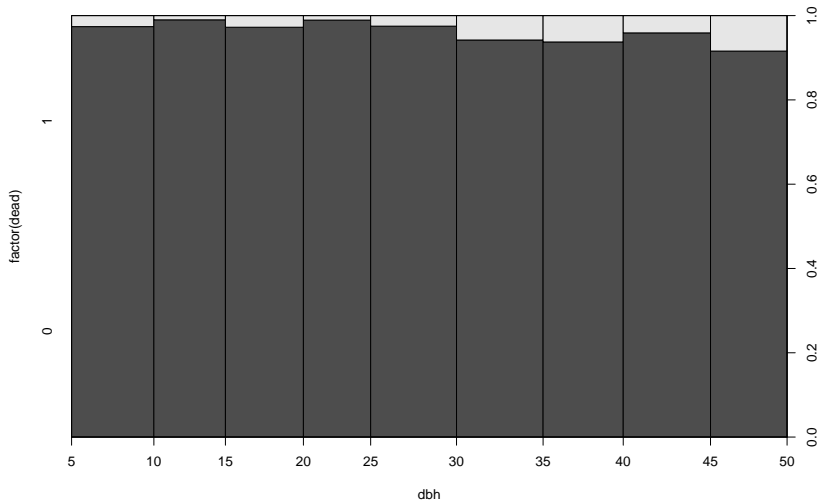
Q: Relationship between tree size and mortality

```
plot(dead ~ dbh, data = trees)
```



Q: Relationship between tree size and mortality

```
plot(factor(dead) ~ dbh, data = trees)
```



Fit simple logistic regression

```
simple.logis <- glm(dead ~ dbh, data = trees, family=binomial)
```

Call:

```
glm(formula = dead ~ dbh, family = binomial, data = trees)
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-0.4121	-0.3287	-0.2624	-0.2048	2.9127

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-4.46945	0.49445	-9.039	< 2e-16 ***
dbh	0.04094	0.01380	2.967	0.00301 **

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 329.51 on 999 degrees of freedom

Logistic regression with *independent* plot effects

```
logis2 <- glm(dead ~ dbh + factor(plot), data = trees, family=bi
```

Call:

```
glm(formula = dead ~ dbh + factor(plot), family = binomial, data
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-0.5923	-0.3198	-0.2549	-0.1940	2.8902

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-4.40106	0.52997	-8.304	<2e-16 ***
dbh	0.04060	0.01386	2.929	0.0034 **
factor(plot)2	-0.59168	0.52132	-1.135	0.2564
factor(plot)3	0.54576	0.47094	1.159	0.2465
factor(plot)4	0.05507	0.57434	0.096	0.9236
factor(plot)5	-0.38312	0.64222	-0.597	0.5508
factor(plot)6	-0.08426	0.76908	-0.110	0.9128
factor(plot)7	0.03126	1.06064	0.029	0.9765

Fit multilevel logistic regression

```
mixed.logis <- glmer(dead ~ dbh + (1|plot), data=trees, family =
```

```
glmer(formula = dead ~ dbh + (1 | plot), data = trees, family =  
      coef.est coef.se  
(Intercept) -4.47      0.49  
dbh           0.04      0.01
```

Error terms:

Groups	Name	Std.Dev.
plot	(Intercept)	0.00
Residual		1.00

number of obs: 1000, groups: plot, 10

AIC = 325.9, DIC = 319.9

deviance = 319.9

Retrieve model coefficients

```
coef(mixed.logis)
```

```
$plot
```

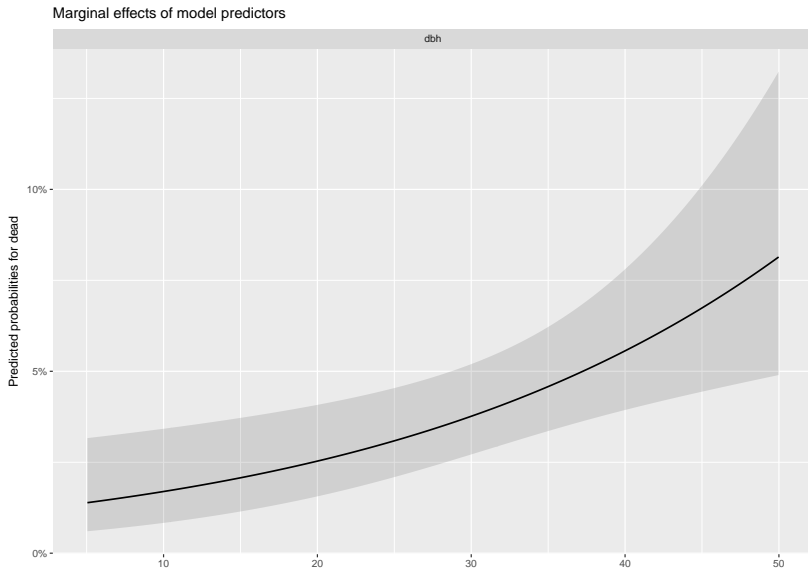
	(Intercept)	dbh
1	-4.469446	0.04093806
2	-4.469446	0.04093806
3	-4.469446	0.04093806
4	-4.469446	0.04093806
5	-4.469446	0.04093806
6	-4.469446	0.04093806
7	-4.469446	0.04093806
8	-4.469446	0.04093806
9	-4.469446	0.04093806
10	-4.469446	0.04093806

```
attr(,"class")
```

```
[1] "coef.mer"
```

Visualising model: sjPlot

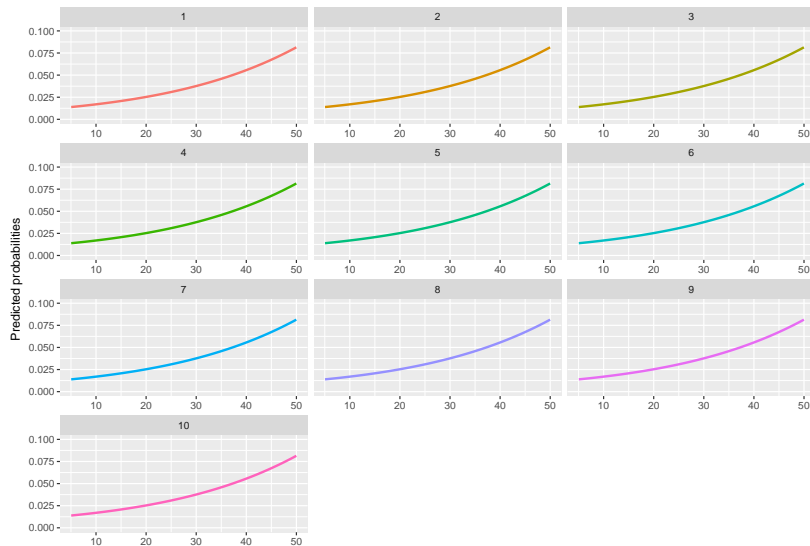
```
sjp.glmer(mixed.logis, type = "eff", show.ci = TRUE)
```



Visualising model: sjPlot

```
sjp.glmer(mixed.logis, type = "ri.slope")
```

Predicted probabilities of dbh on dead



Advantages of multilevel models

- ▶ Perfect for structured data (space-time)

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- ▶ Accommodate variation in treatment effects
- ▶ More efficient inference of regression parameters
- ▶ Using all the data to perform inferences for groups with small sample size

Formula syntax for different models

$y \sim x + (1 \mid \text{group})$ # varying intercepts

$y \sim x + (1 + x \mid \text{group})$ # varying intercepts and slopes

$y \sim x + (1 \mid \text{group/subgroup})$ # nested

$y \sim x + (1 \mid \text{group1}) + (1 \mid \text{group2})$ # varying intercepts, crossed

$y \sim x + (1 + x \mid \text{group1}) + (1 + x \mid \text{group2})$ # varying intercepts and slopes, crossed

END



Figure 9:

Source code and materials:

<https://github.com/Pakillo/LM-GLM-GLMM-intro>