

Topic 12 – Nonlinear regression

ENVX1002 Statistics in Life and Environmental Sciences

Si Yang Han

The University of Sydney

Feb 2026

Module overview

- Week

- 9.

- Describing Relationships

- Correlation (calculation, interpretation)
 - Regression (model structure, model fitting)
 - What/when/why/how

- Week

- 10.

- Simple Linear Regression

- Can we use the model? (assumptions, hypothesis testing)
 - How good is the model? (interpretation, model fit)

- Week

- 11.

- Multiple Linear Regression

- Multiple Linear Regression (MLR) modelling
 - Assumptions, interpretation and the principle of parsimony

- **Week 12. Nonlinear Regression**

- Common nonlinear functions
- Transformations

Regressions

Simple linear regression

$$Y_i = \beta_0 + \beta_1 x_i + \epsilon_i$$

Ideal for predicting a continuous response variable from a single predictor variable: “*How does y change as x changes, when the relationship is linear?*”

Multiple linear regression

$$Y_i = \beta_0 + \beta_1 x_{1i} + \beta_2 x_{2i} + \dots + \beta_k x_{ki} + \epsilon_i$$

“*How does y change as x₁, x₂, ..., x_k change?*”

Nonlinear regression

$$Y_i = f(x_i, \beta) + \epsilon_i$$

where $f(x_i, \beta)$ is a nonlinear function of the parameters β : “*How do we model a change in y with x when the relationship is nonlinear?*”

Nonlinear regression



Carl Friedrich Gauss (1777-1855) and Isaac Newton (1642-1726) Gauss-Newton approach to nonlinear regression is most commonly used

Nonlinear relationships

Linear relationships are simple to interpret since the rate of change is constant.

“As one changes, the other changes at a constant rate.”

Nonlinear relationships often involve exponential, logarithmic, or power functions.

“As one changes, the other changes at a rate that is *not proportional* to the change in the other.

Dealing with nonlinearity

Transformations

Often, a nonlinear relationship may be transformed into a linear relationship by applying a transformation to the response variable or the predictor variable(s).

- **Logarithmic:** $y = \log(x)$
- **Exponential:** $y = e^x$
- **Square-root:** $y = \sqrt{x}$
- **Inverse:** $y = \frac{1}{x}$
- Usually works when y changes **monotonically** with x .
- More interpretable and easier to fit.

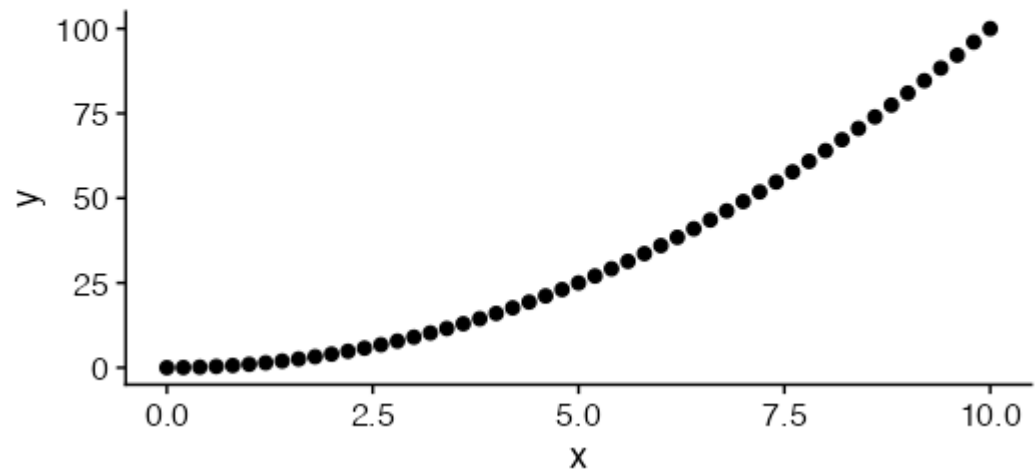
Nonlinear relationships: exponents

- x^2 is the *square* of x .
- x^3 is the *cube* of x .
- x^a is x raised to the *power* of a .

In a relationship where y is a function of x^a , as x increases, y increases nonlinearly at a rate that depends on the value of x and a ($\frac{dy}{dx} = ax^{a-1}$).

```
# Plot a simulation of above in ggplot2
set.seed(123)
tibble(x = seq(0, 10, by = 0.2), y = x^2) %>%
  ggplot(aes(x = x, y = y)) +
  geom_point() +
  labs(x = "x", y = "y") +
  ggtitle(expression(y == x^2)) +
  theme(plot.title = element_text(size = 40, face = "bold"))
```

$$y = x^2$$



Nonlinear relationships: logarithms

- $\log_e(x)$ is the *natural logarithm* of x .
- $\log_{10}(x)$ is the *common logarithm* of x .
- $\log_a(x)$ is the *logarithm* of x to the base a .

Interpretation:

- If $\log_a(y) = x$: as x increases, the value of y increases by $y = a^x$.
- If $y = \log_a(x)$: as x increases, the value of y increases by $y = \log_a(x)$. As y increases, the value of x increases by $x = a^y$.

Exponents and logarithms

	Exponents	Logarithms
Definition	If $a^n = b$, a is the base, n is the exponent, and b is the result.	If $\log_a b = n$, a is the base, b is the result, and n is the logarithm (or the exponent in the equivalent exponential form).
Example	$2^3 = 8$	$\log_2 8 = 3$
Interpretation	2 raised to the power of 3 equals 8.	The power to which you must raise 2 to get 8 is 3.
Inverse	The logarithm is the inverse operation of exponentiation.	The exponentiation is the inverse operation of logarithm.
Properties	$(a^n)^m = a^{n \cdot m}$, $a^n \cdot a^m = a^{n+m}$, $\frac{a^n}{a^m} = a^{n-m}$	$\log_a(b \cdot c) = \log_a b + \log_a c$, $\log_a\left(\frac{b}{c}\right) = \log_a b - \log_a c$, $\log_a(b^n) = n \cdot \log_a b$

i Note

For your understanding, not examinable.

Common nonlinear functions

$$f(x_i, \beta)$$

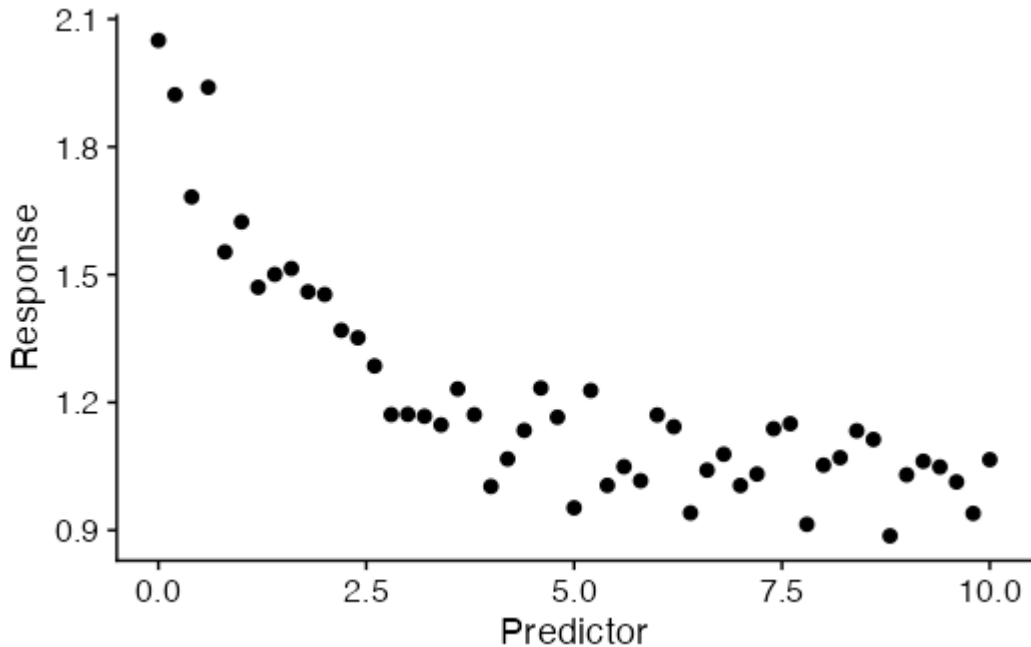
Exponential decay relationship

Response variable *decreases* and approaches limit as predictor variable increases.

$$y = a \cdot e^{-bx}$$

```
set.seed(429) # set seed
# Simulate data:
decay ← tibble(
  predictor = seq(0,10, by = 0.2),
  response = abs(exp(-0.5*predictor) + rnorm(length(predictor), mean = 1, sd = 0.1)))

ggplot(data = decay, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response")
```



Examples: radioactive decay, population decline, chemical reactions.

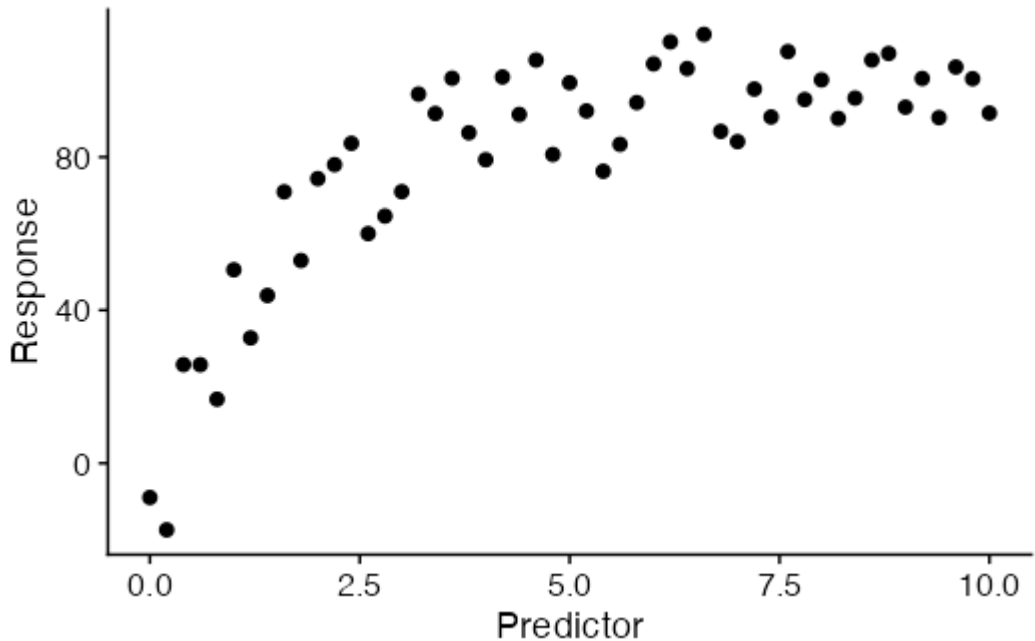
Asymptotic relationship

Response variable *increases* and approaches a limit as the predictor variable increases.

$$y = a + b(1 - e^{-cx})$$

```
set.seed(442) # set seed
# Simulate data:
asymptotic = tibble(
  predictor = seq(0, 10, by = 0.2),
  response = 100*(1-exp(-0.5*predictor)) + rnorm(length(predictor), mean = 0, sd = 10))

ggplot(data = asymptotic, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response")
```



Examples: population growth, enzyme kinetics.

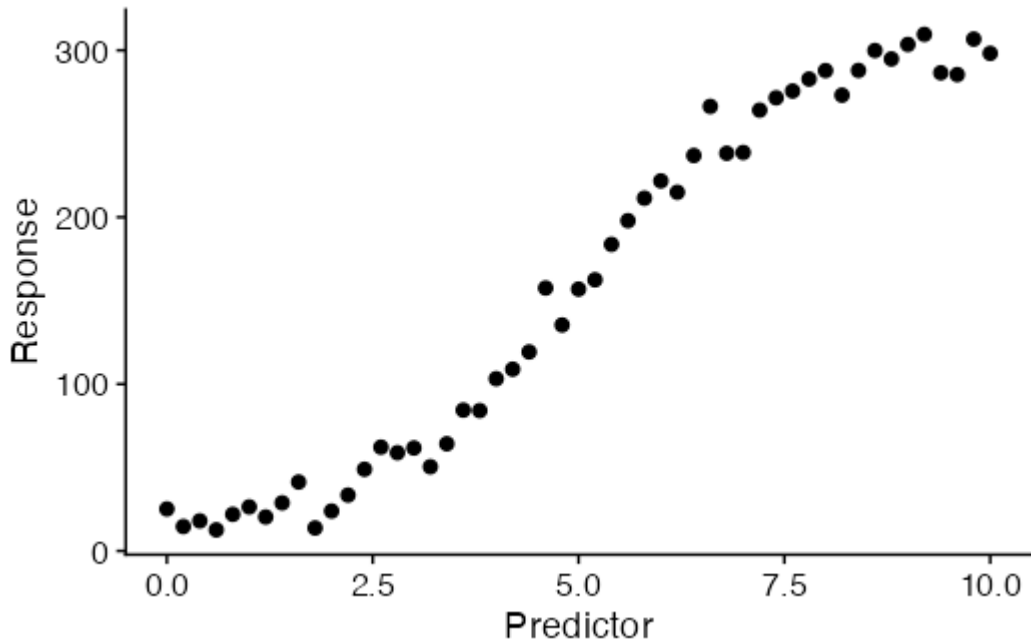
Logistic relationship

An S-shaped relationship, where the response variable is at first exponential, then asymptotic.

$$y = c + \frac{d - c}{1 + e^{-b(x-a)}}$$

```
set.seed(450)
# Simulate data:
logistic ← tibble(predictor = seq(0, 10, by = 0.2),
  response = 10 + abs(300 * (1 / (1 + exp(-0.8 * (predictor - 5))))) + rnorm(length(predictor), mean
= 0, sd = 10))

ggplot(data = logistic, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response")
```



Examples: growth of bacteria, disease spread, species growth.

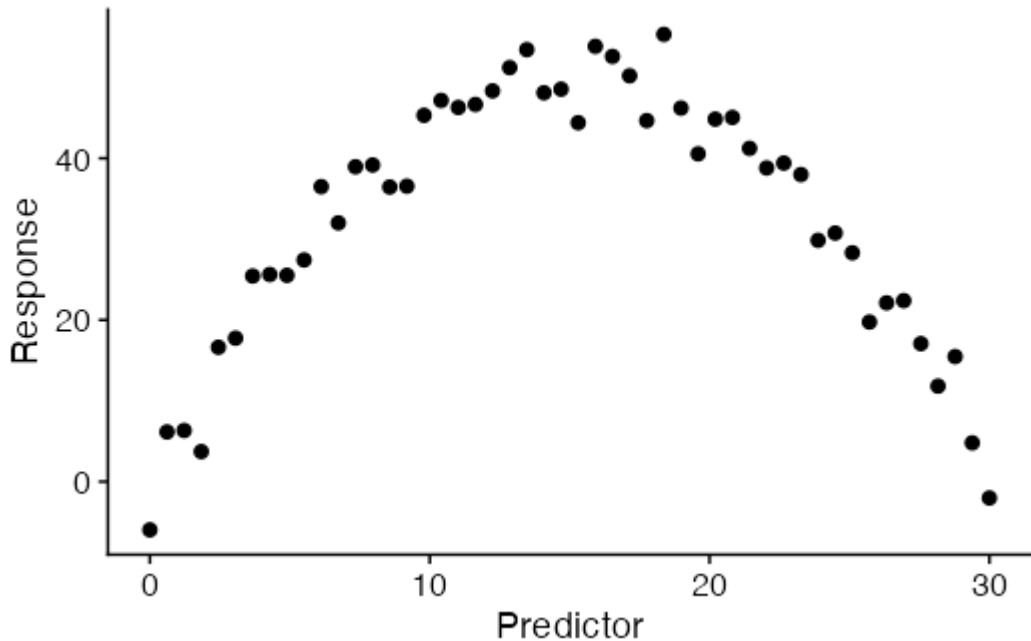
Polynomial relationship

Response variable changes in a variety of ways as the predictor variable changes. Also known as ‘curvilinear’.

$$y = a + bx + cx^2 + dx^3 + \dots$$

```
# Set seed for reproducibility
set.seed(529)
# Simulate data:
curvilinear ← tibble(predictor = seq(0, 30, length.out = 50),
  response = 50 * (1 - (predictor - 15)^2 / 225) + rnorm(length(predictor), mean = 0, sd = 5))

ggplot(data = curvilinear, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response")
```



Examples: food intake, drug dosage, exercise.

Transformations

| How far can we go?

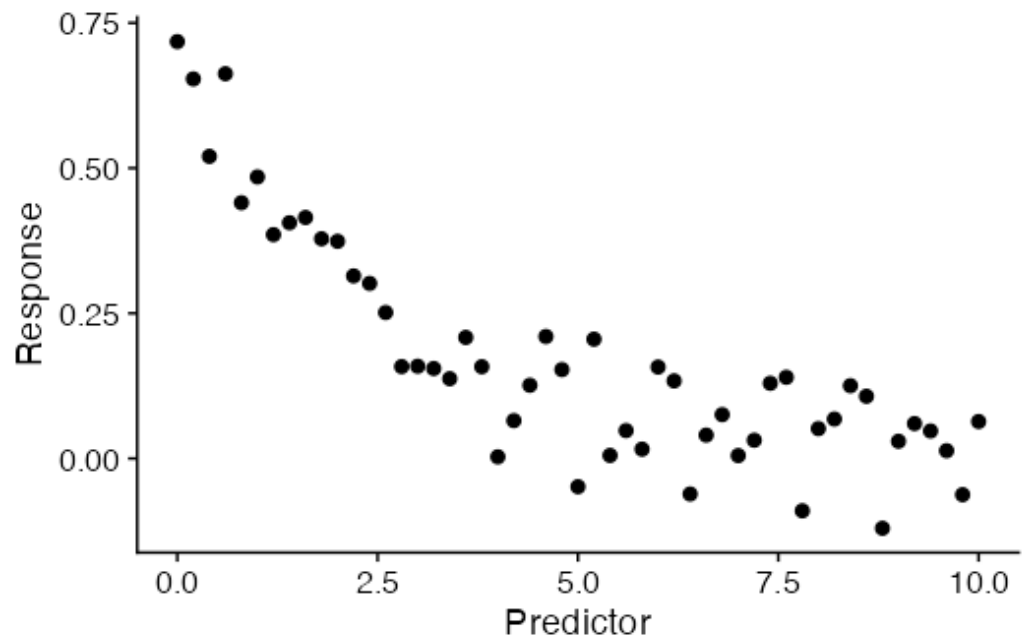
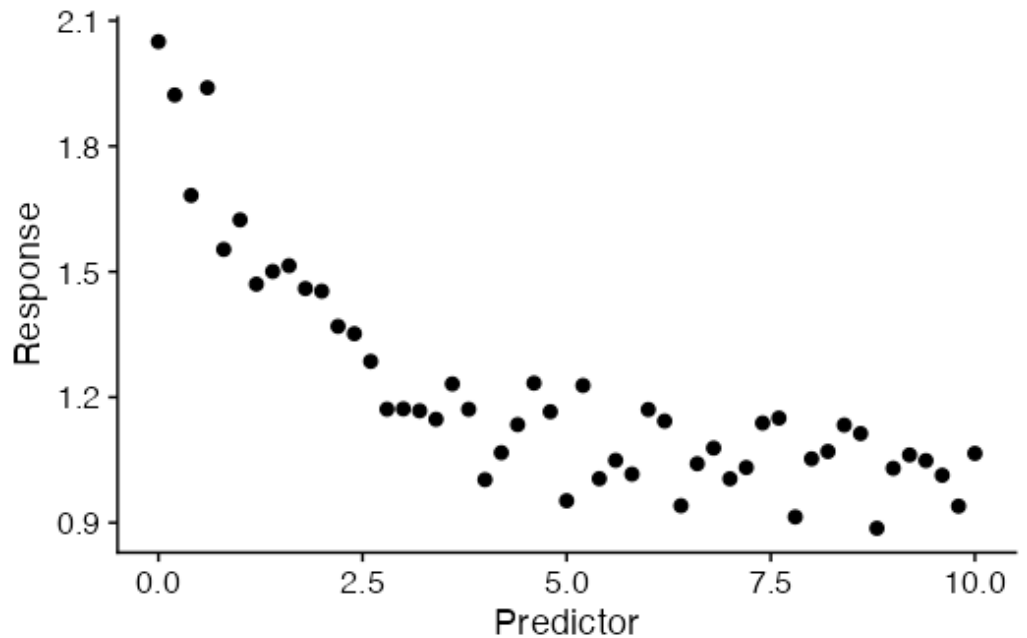
Transformations: exponential decay

Before transformation

```
ggplot(data = decay,  
       aes(x = predictor, y = response)) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```

After \log_e transform

```
ggplot(data = decay,  
       aes(x = predictor, y = log(response))) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```



Transformations: exponential decay

Before transformation

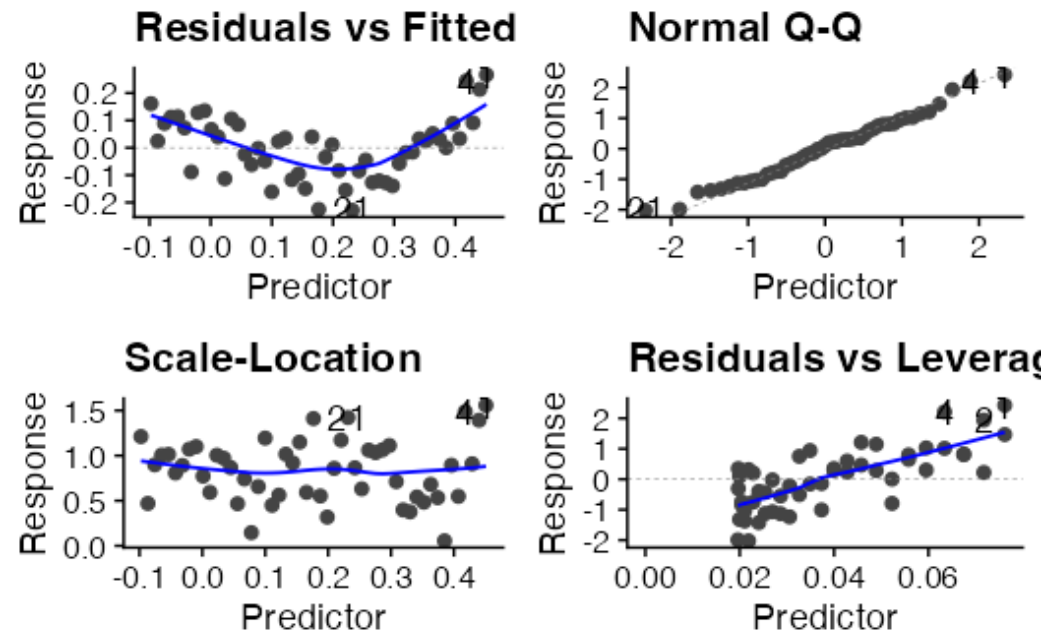
```
autoplot(lm(response ~ predictor, data = decay)) +  
  labs(x = "Predictor", y = "Response")
```

Warning: `fortify(<lm>)` was deprecated in ggplot2 4.0.0.
i Please use `broom::augment(<lm>)` instead.
i The deprecated feature was likely used in the ggfortify package.
Please report the issue at <<https://github.com/sinhrks/ggfortify/issues>>.

Warning: `aes_string()` was deprecated in ggplot2 3.0.0.
i Please use tidy evaluation idioms with `aes()`.
i See also `vignette("ggplot2-in-packages")` for more information.

After \log_e transform

```
autoplot(lm(log(response) ~ predictor, data = decay)) +  
  labs(x = "Predictor", y = "Response")
```



i The deprecated feature was likely used in the ggfortify package.

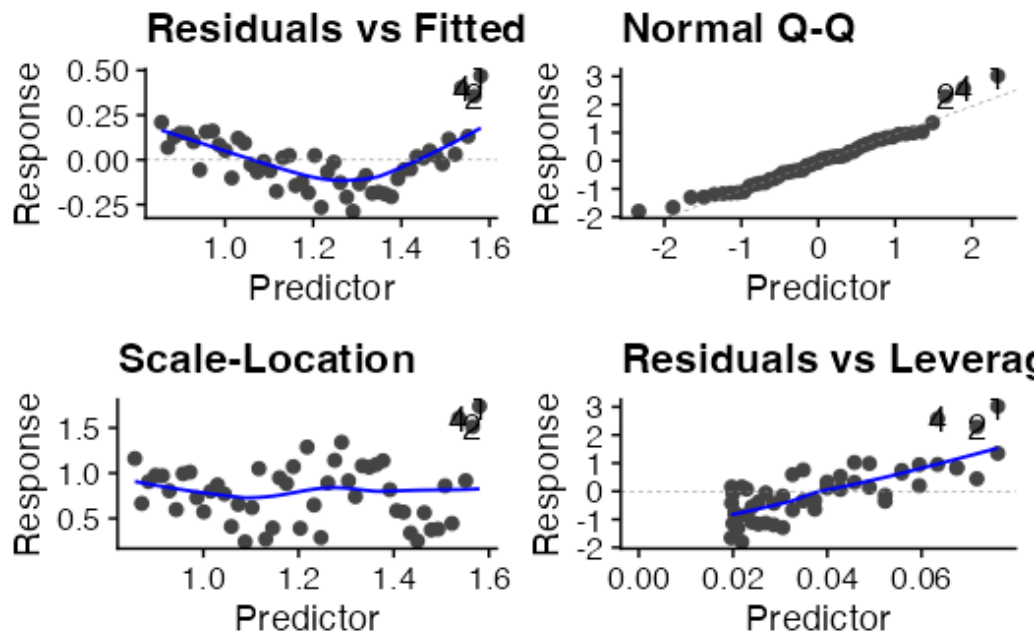
Please report the issue at <<https://github.com/sinhrks/ggfortify/issues>>.

Warning: Using `size` aesthetic for lines was deprecated in ggplot2 3.4.0.

i Please use `linewidth` instead.

i The deprecated feature was likely used in the ggfortify package.

Please report the issue at <<https://github.com/sinhrks/ggfortify/issues>>.



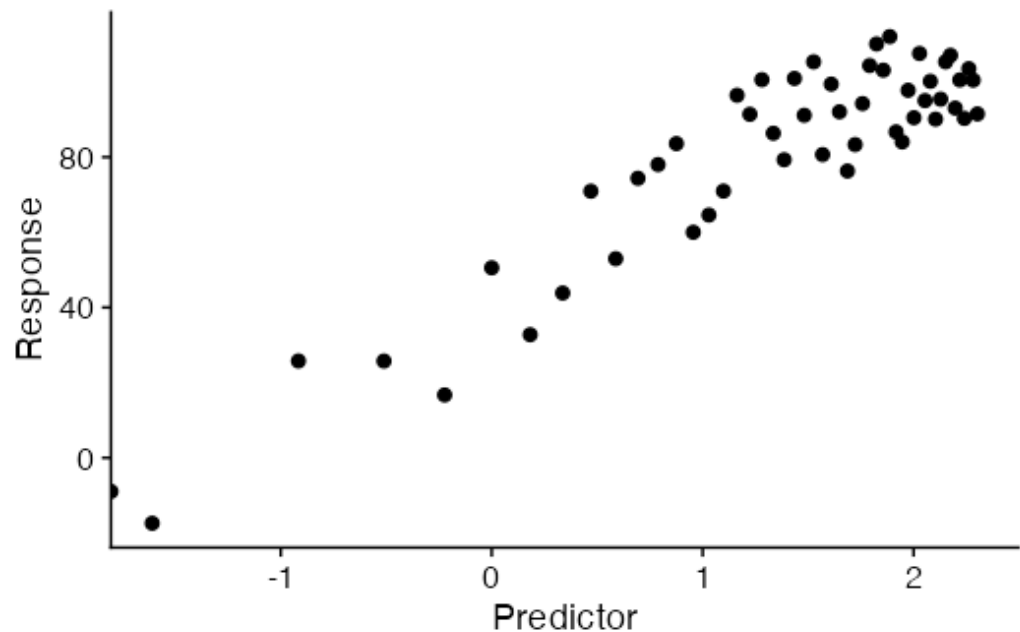
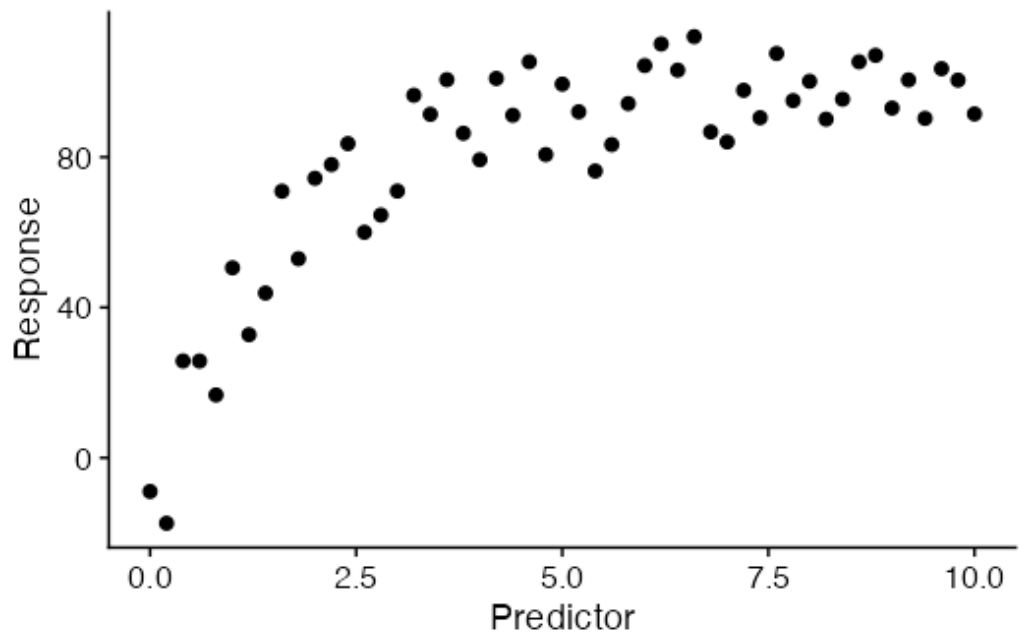
Transformations: asymptotic relationship

Before transformation

```
ggplot(data = asymptotic,  
       aes(x = predictor, y = response)) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```

After \log_e transform

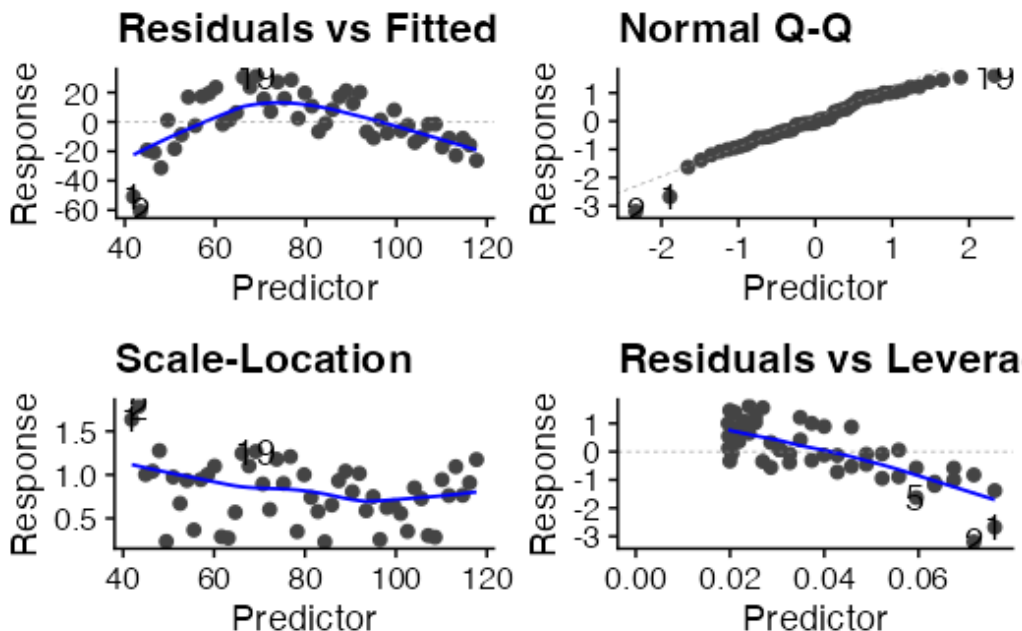
```
ggplot(data = asymptotic,  
       aes(x = log(predictor), y = response)) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```



Transformations: asymptotic relationship

Before transformation

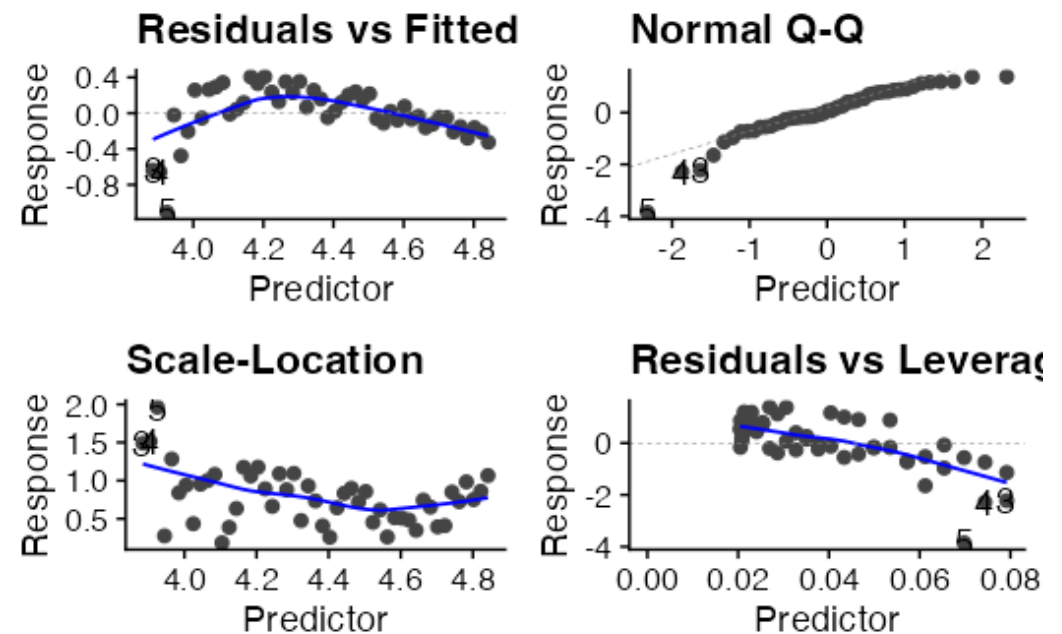
```
autoplplot(lm(response ~ predictor, data =  
asymptotic)) +  
  labs(x = "Predictor", y = "Response")
```



After \log_e transform

```
autoplplot(lm(log(response) ~ predictor, data =  
asymptotic)) +  
  labs(x = "Predictor", y = "Response")
```

Warning in log(response): NaNs produced



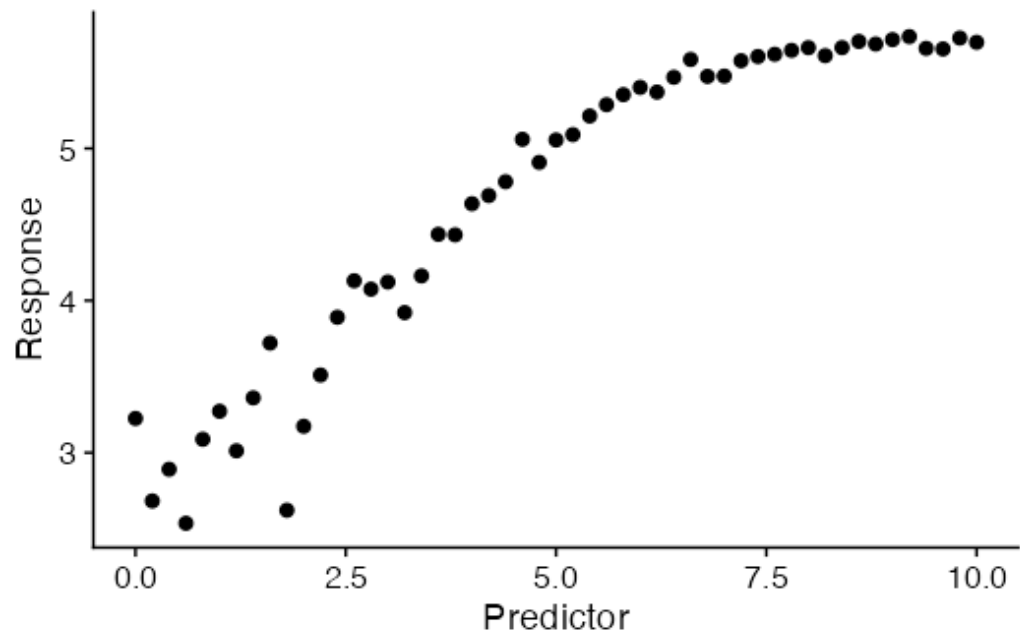
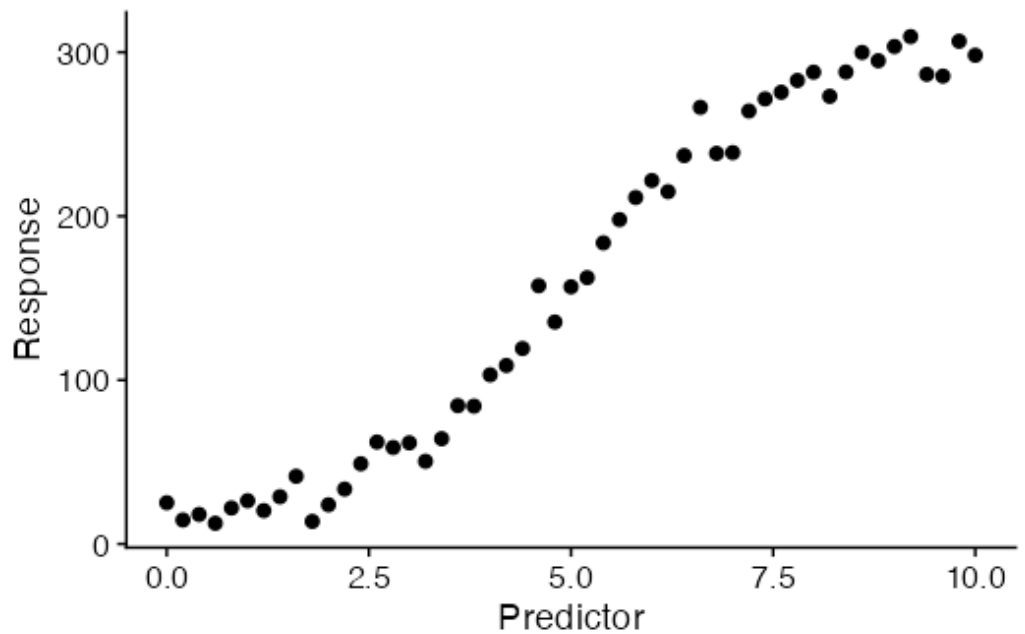
Transformations: logistic relationship

Before transformation

```
ggplot(data = logistic,  
       aes(x = predictor, y = response)) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```

After \log_e transform

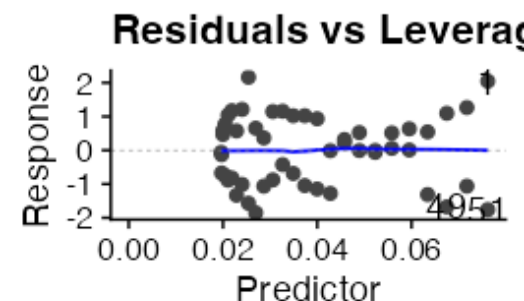
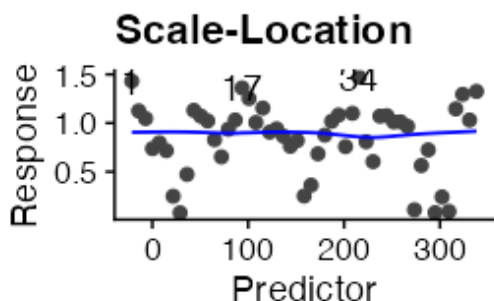
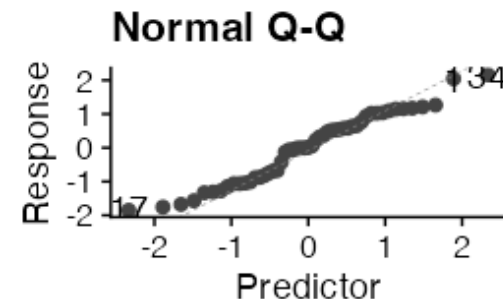
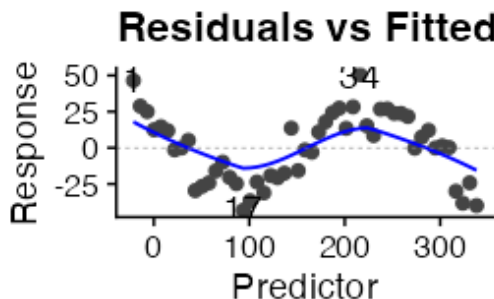
```
ggplot(data = logistic,  
       aes(x = predictor, y = log(response))) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```



Transformations: logistic relationship

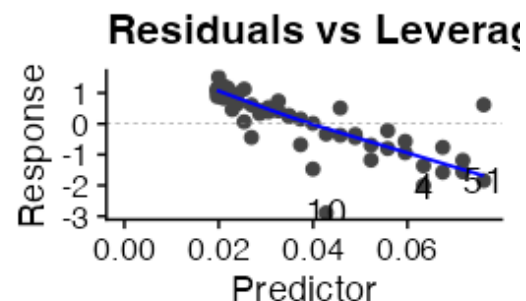
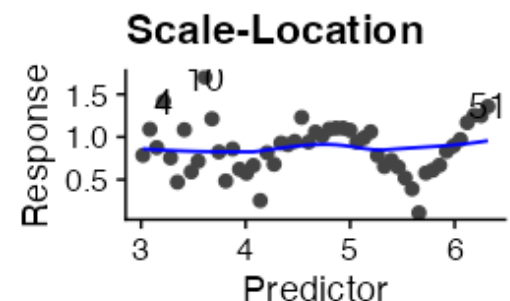
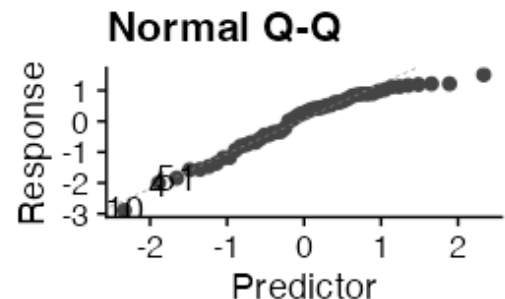
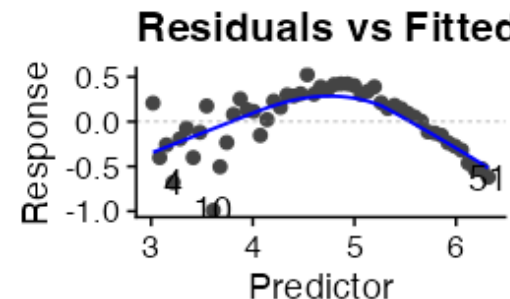
Before transformation

```
autoplplot(lm(response ~ predictor, data =  
logistic)) +  
  labs(x = "Predictor", y = "Response")
```



After \log_e transform

```
autoplplot(lm(log(response) ~ predictor, data =  
logistic)) +  
  labs(x = "Predictor", y = "Response")
```



Transformations: polynomial relationship

Before transformation

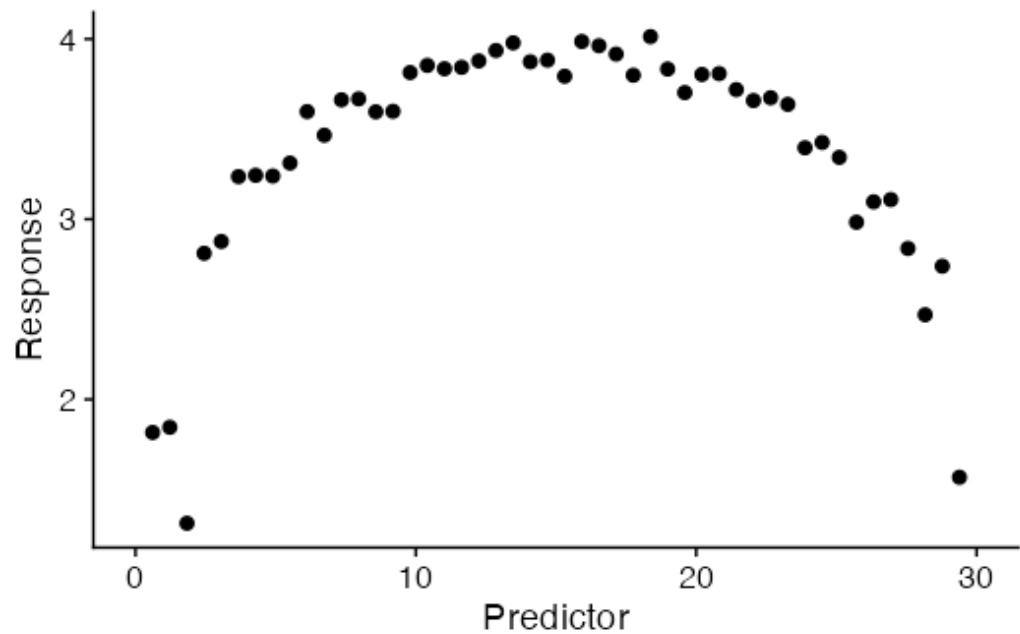
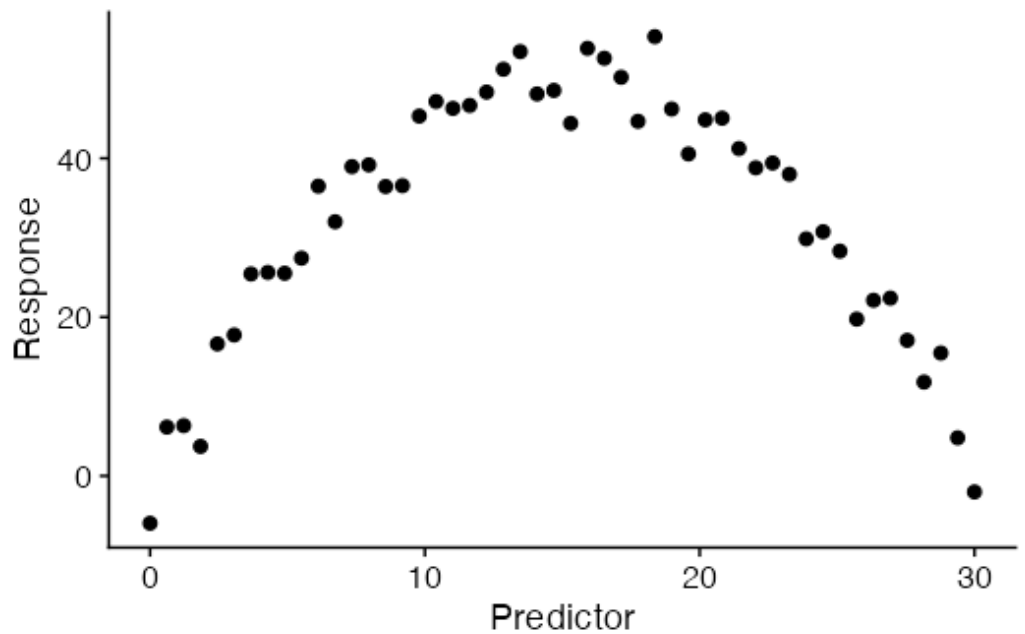
```
ggplot(data = curvilinear,  
       aes(x = predictor, y = response)) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```

After \log_e transform

```
ggplot(data = curvilinear,  
       aes(x = predictor, y = log(response))) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```

Warning in log(response): NaNs produced

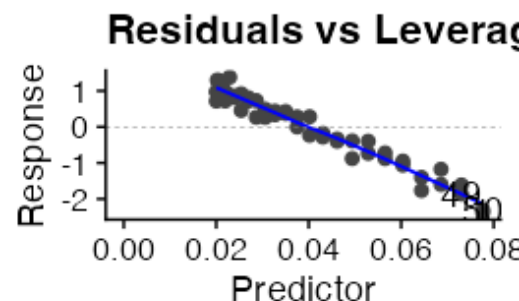
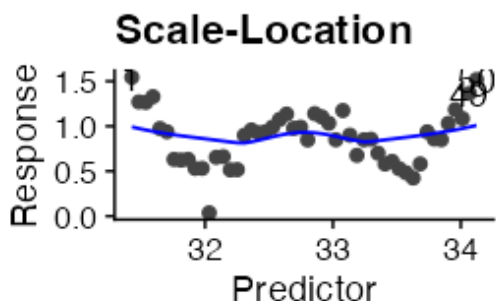
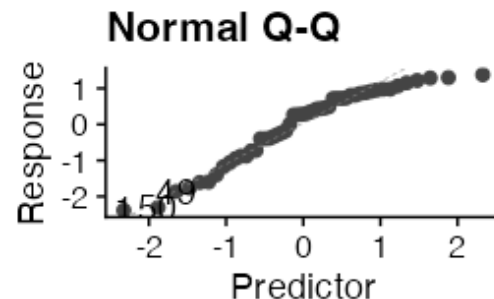
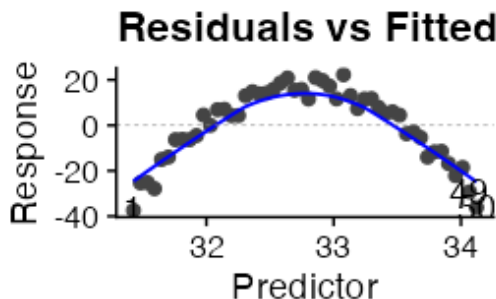
Warning: Removed 2 rows containing missing
values or values outside the scale range
(`geom_point()`).



Transformations: polynomial relationship

Before transformation

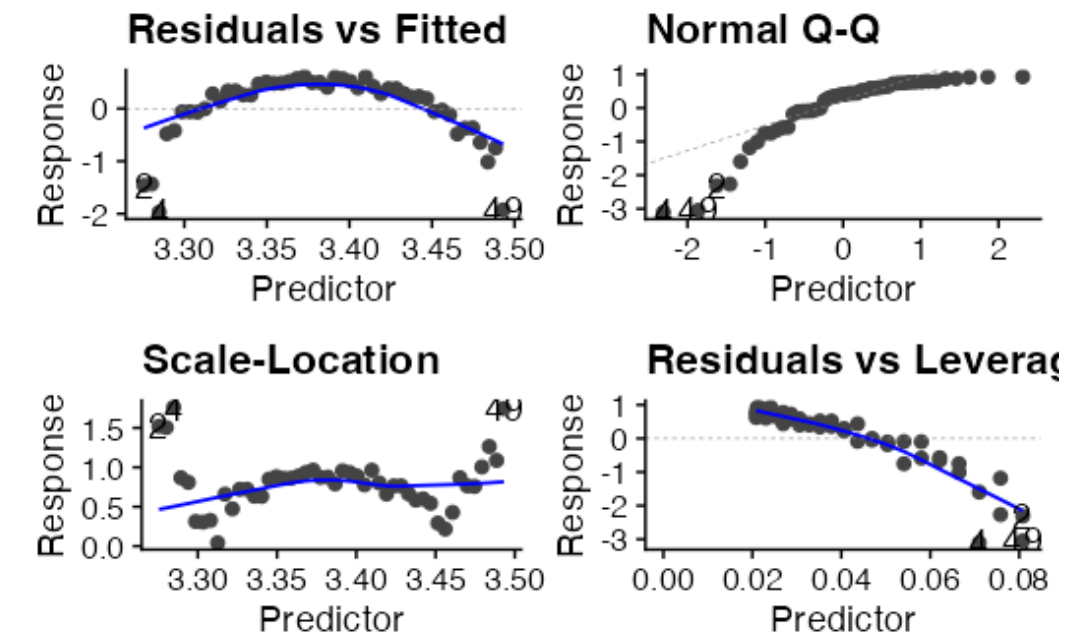
```
autoplot(lm(response ~ predictor, data =  
curvilinear)) +  
  labs(x = "Predictor", y = "Response")
```



After \log_e transform

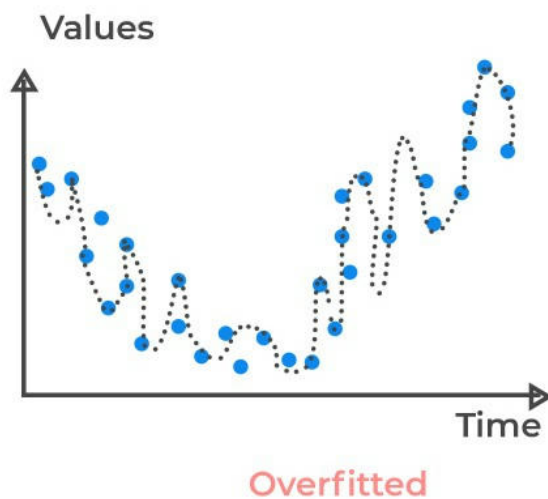
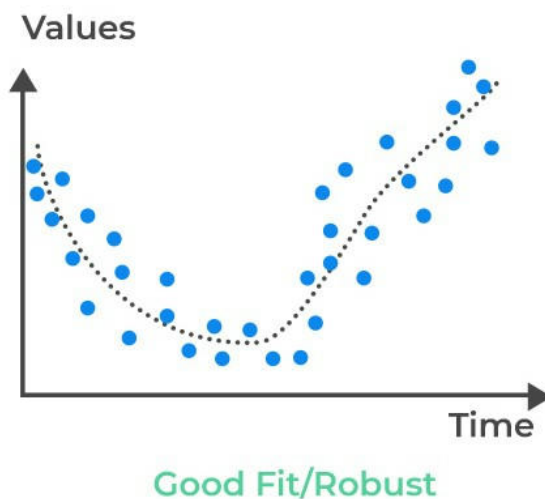
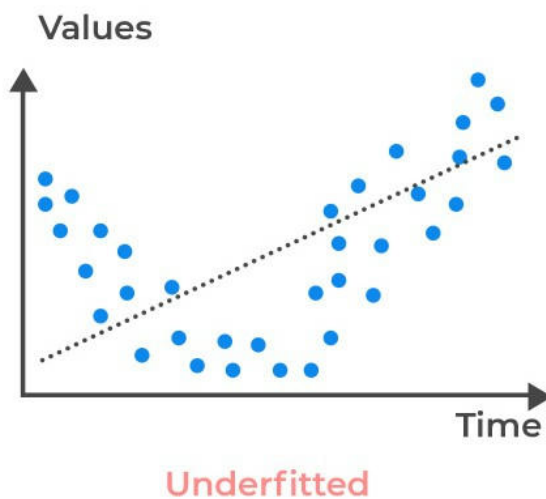
```
autoplot(lm(log(response) ~ predictor, data =  
curvilinear)) +  
  labs(x = "Predictor", y = "Response")
```

Warning in log(response): NaNs produced



Did the transformations work?

- To a *certain* extent...
- **Problems:**
 - Relationships typically do not meet the linear assumption, but seem “ok” for other assumptions.
 - Poor fit to the data (over or underfitting in some areas).
 - Difficult to interpret the results.



Nonlinear regression

- A way to model complex (nonlinear) relationships.
 - ▶ i.e. phenomena that arise in the natural and physical sciences e.g. biology, chemistry, physics, engineering.
- At least *one* predictor is not linearly related to the response variable.
- Unique/specific shape - apply only if you are sure of the relationship, e.g. asymptotic, quadratic.

Performing nonlinear regression

- **Polynomial regression:** still linear in the parameters and a good place to start.
- **Nonlinear regression:** use the `nls()` function to fit the following nonlinear models:
 - ▶ Exponential growth
 - ▶ Exponential decay
 - ▶ Logistic

Polynomial regression

| A special case of multiple linear regression used to model nonlinear relationships.

Model

$$Y_i = \beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \dots + \beta_k x_i^k + \epsilon_i$$

where k is the degree of the polynomial.

- The model is still linear in the parameters β and can be fitted using least squares.
- Instead of multiple predictors, we have multiple *terms* of the same predictor (same x).
- Only the *highest-order term* is tested for significance.
- Can still be fit using `lm()`.
- The more complex, the less likely it follows a true biological relationship...

...

Adding polynomial terms

- Linear: $y = \beta_0 + \beta_1 x$
- Quadratic: $y = \beta_0 + \beta_1 x + \beta_2 x^2$
- Cubic: $y = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3$
- Each level increases the power of the predictor by 1.

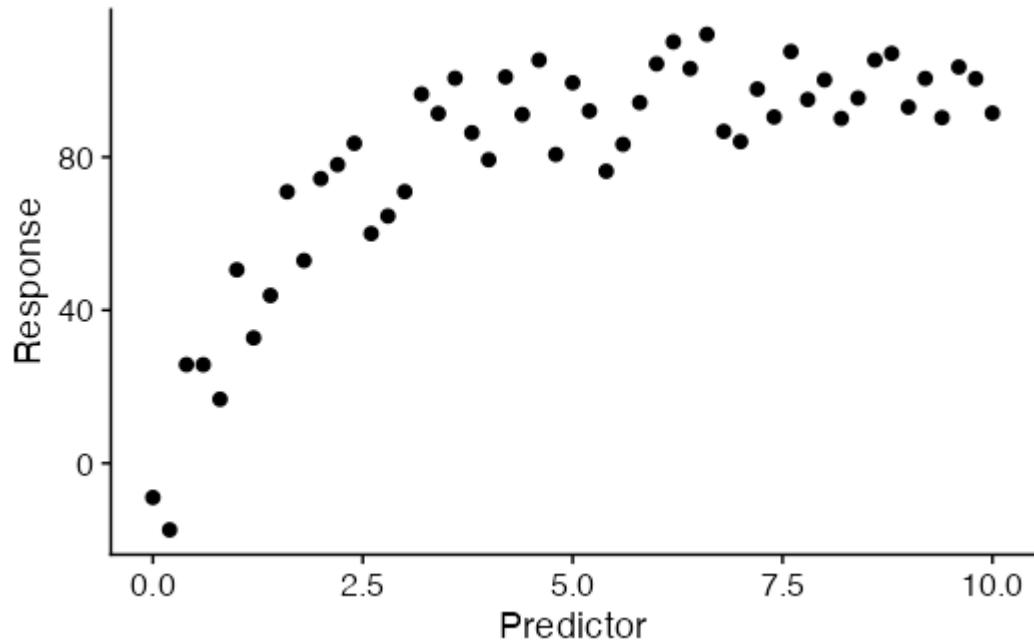
Polynomial fitting

Using the `asymptotic` data

The data

See Slide 11 for the relationship and mathematical expression.

```
ggplot(asymptotic, aes(x = predictor, y = response)) +  
  geom_point() +  
  labs(x = "Predictor", y = "Response")
```

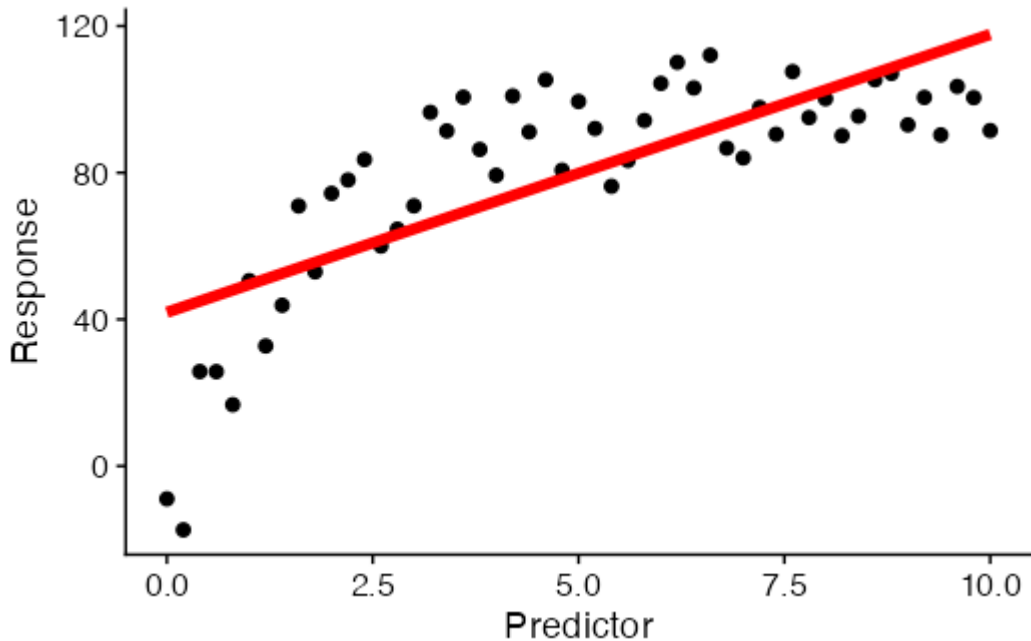


Fitting the model (linear)

$$Y_i = \beta_0 + \beta_1 x_i + \epsilon_i$$

```
lin_fit ← lm(response ~ predictor, asymptotic)
```

```
ggplot(asymptotic, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response") +
  geom_line(aes(y = predict(lin_fit)), color = "red", size = 2)
```

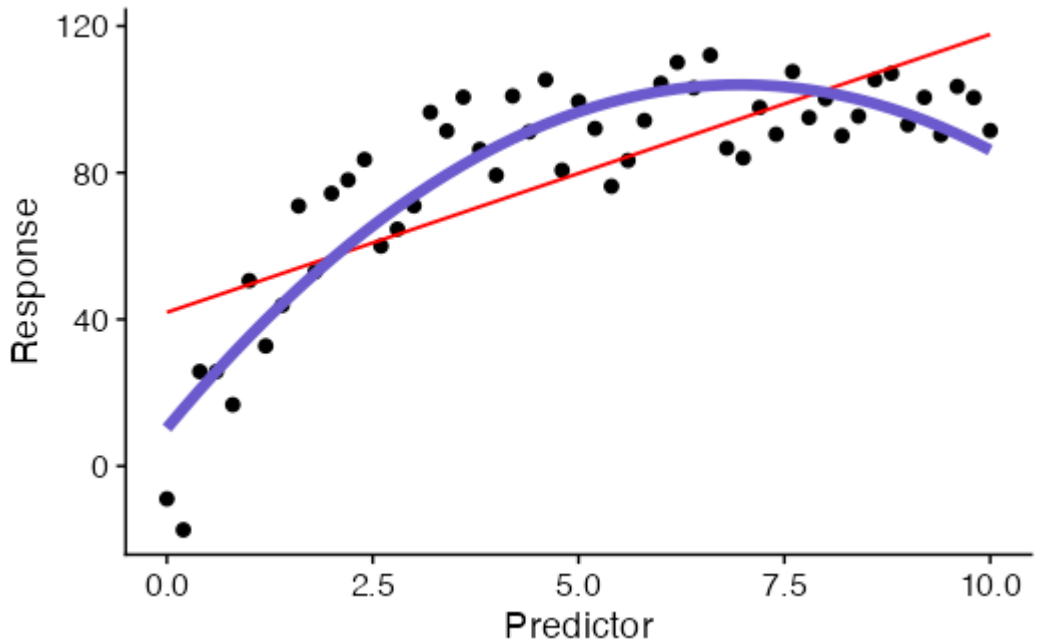


Fitting the model (`poly(degree = 2)`)

$$Y_i = \beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \epsilon_i$$

```
poly2_fit ← lm(response ~ poly(predictor, 2), asymptotic)
```

```
ggplot(asymptotic, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response") +
  geom_line(aes(y = predict(lin_fit)), color = "red") +
  geom_line(aes(y = predict(poly2_fit)), color = "slateblue", size = 2)
```

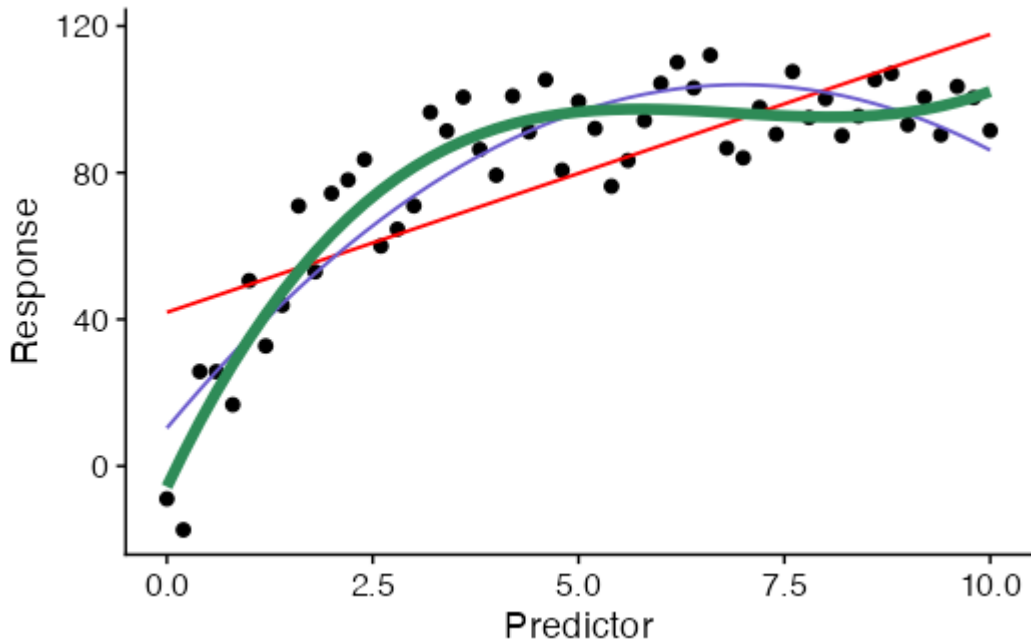


Fitting the model (`poly(degree = 3)`)

$$Y_i = \beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \beta_3 x_i^3 + \epsilon_i$$

```
poly3_fit ← lm(response ~ poly(predictor, 3), asymptotic)
```

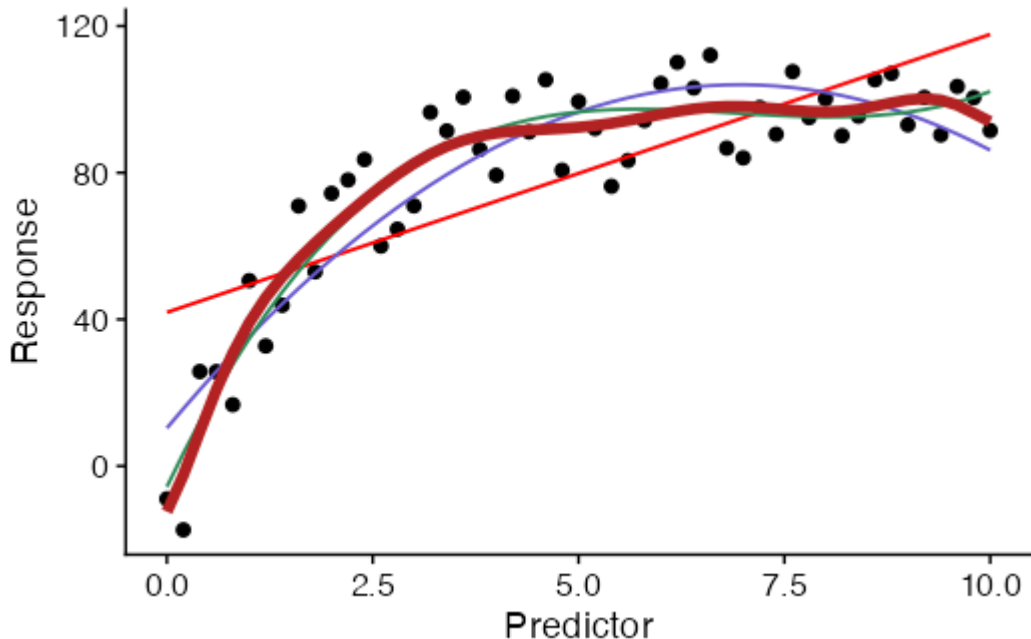
```
ggplot(asymptotic, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response") +
  geom_line(aes(y = predict(lin_fit)), color = "red") +
  geom_line(aes(y = predict(poly2_fit)), color = "slateblue") +
  geom_line(aes(y = predict(poly3_fit)), color = "seagreen", size = 2)
```



Fitting the model (`poly(degree = 10)`)

$$Y_i = \beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \dots + \beta_{10} x_i^{10} + \epsilon_i$$

```
poly10_fit ← lm(response ~ poly(predictor, 10), asymptotic)
```



Model	R2
Linear	0.570
Poly2	0.820
Poly3	0.872
Poly10	0.862

i Note

We use adjusted R² for polynomials - extra terms, extra complexity, so extra penalty.

Limitations

- Meaning of the coefficients is not always clear.
- Extrapolation can be *dangerous*.
- Extra terms can lead to overfitting and are difficult to interpret:
- Parsimony: is the most complex term (highest power) significant? If not, use a lower power.

```
summary(poly10_fit)
```

Call:

```
lm(formula = response ~ poly(predictor, 10), data = asymptotic)
```

Residuals:

Min	1Q	Median	3Q	Max
-17.1659	-8.6908	-0.0494	8.8003	16.4012

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	79.818	1.552	51.426	< 2e-16 ***

```
poly(predictor, 10)1  159.368    11.084   14.378 < 2e-16 ***
poly(predictor, 10)2 -106.939    11.084  -9.648 5.37e-12 ***
poly(predictor, 10)3  48.570    11.084   4.382 8.28e-05 ***
poly(predictor, 10)4 -19.411    11.084  -1.751  0.0876 .
poly(predictor, 10)5  1.193     11.084   0.108  0.9148
poly(predictor, 10)6 -2.769     11.084  -0.250  0.8040
poly(predictor, 10)7 -1.343     11.084  -0.121  0.9042
poly(predictor, 10)8 -4.009     11.084  -0.362  0.7195
poly(predictor, 10)9 -2.851     11.084  -0.257  0.7984
poly(predictor, 10)10 5.769     11.084   0.520  0.6056
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
Residual standard error: 11.08 on 40 degrees of freedom
Multiple R-squared:  0.8897,    Adjusted R-squared:  0.8621
F-statistic: 32.26 on 10 and 40 DF,  p-value: 4.846e-16
```

Still:

- Easy to fit: just add polynomial terms to the model.
- Simple to perform: use `lm()`.

Nonlinear fitting

Fitting a nonlinear model

If you have some understanding of the underlying relationship (e.g. mechanistic process) between the variables, you can fit a nonlinear model.

Mathematical expression

$$Y_i = f(x_i, \beta) + \epsilon_i$$

where $f(x_i, \beta)$ is a nonlinear function of the parameters β .

- Y_i is the continuous response variable.
- x_i is the vector of predictor variables.
- β is the vector of unknown parameters.
- ϵ_i is the random error term (residual error).

Assumptions

Like the linear model, the nonlinear model assumes *INE*:

- Error terms are independent (**Independence**).
- Error terms are normally distributed (**Normality**).
- Error terms have equal/constant variance (**Homoscedasticity**).

Basically:

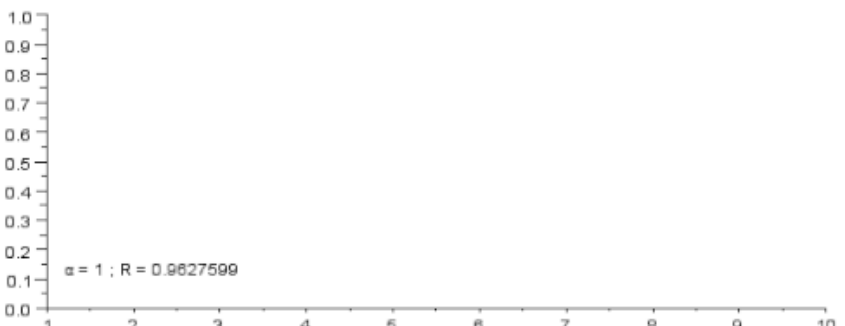
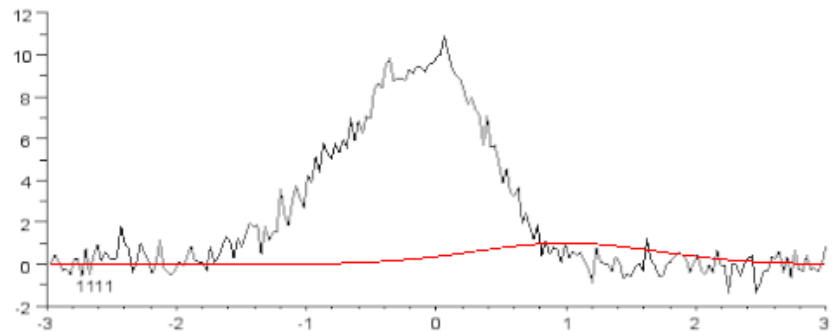
$$\epsilon_i \sim N(0, \sigma^2)$$

...

Like all other models we have seen, we focus on the residuals to assess the model fit, since the residuals are the only part of the model that is random.

Estimating the model parameters

- The parameters are estimated using the **method of least squares**.
- For nonlinear models, a nonlinear optimization algorithm is used to find the best fit, rather than ordinary least squares:
 - ▶ **Gauss-Newton algorithm**
 - ▶ **Levenberg-Marquardt algorithm**
- This can only be performed iteratively and depends on a “best guess” of the parameters *as a start*.
 - ▶ **i.e. we need to provide a starting point for a nonlinear least squares algorithm to begin.**



Source: [Wikipedia](#)

Two methods in R

Use `nls()` function in R.

```
nls(formula, data, start)
```

- `formula`: a formula object, response variable ~ predictor variable(s).
- `data`: a data frame containing the variables in the model (response, predictor).
- `start`: a named list of starting values for the parameters in the model.

Self-starting functions: `SSexpf()`, `SSasymp()`, `SSlogis()`, etc.

- Self-starting functions estimate the starting values for you.
- Named after the models they fit.
- Existing functions have pre-set formulas.
- Can define own functions but more complex than `nls()`.

Example: Fitting an exponential model

With `nls()`

$$y = y_0 e^{kx}$$

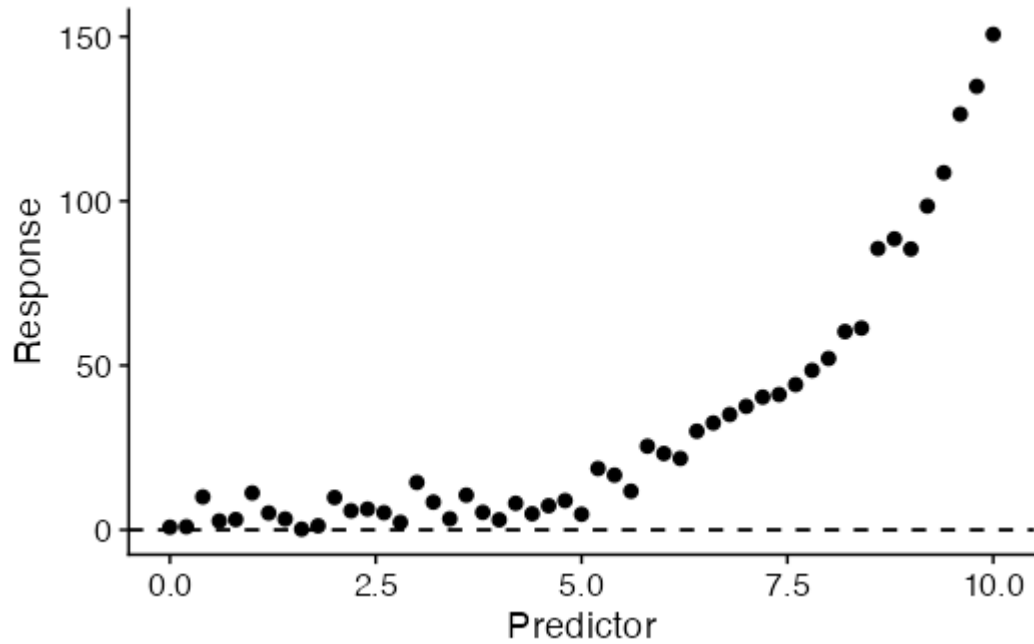
where

- y is the response and x is the predictor
- y_0 is the value of y when $x = 0$
- k is the rate of change

k can be estimated with the equation $slope = k = \frac{\log_e y_{max} - \log_e y_{min}}{x_{max} - x_{min}}$, but usually a value of 1 is a good starting point.

```
set.seed(123)
growth ← tibble(
  predictor = seq(0, 10, by = 0.2),
  response = abs(exp(0.5*predictor) + rnorm(length(predictor), mean = 1, sd = 5)))

ggplot(data = growth, aes(x = predictor, y = response)) +
  geom_point() +
  geom_hline(yintercept = 0, linetype = "dashed") +
  labs(x = "Predictor", y = "Response")
```

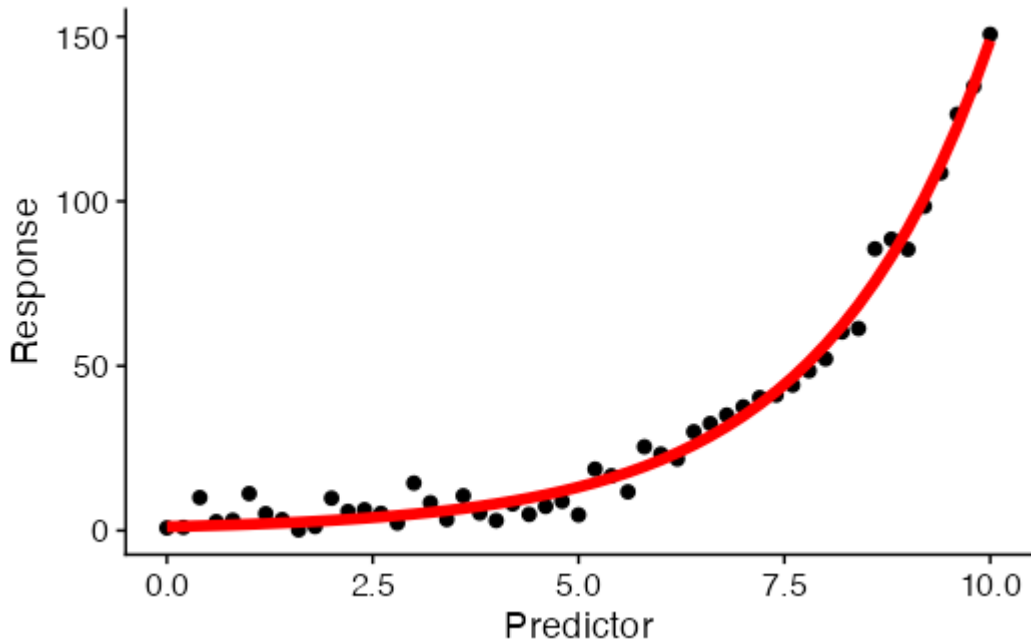


First guess

Based on the plot, we can estimate $y_0 \neq 0$ and $k = 1$. Because of the equation, $y = y_0 e^{kx}$, y_0 cannot be 0!

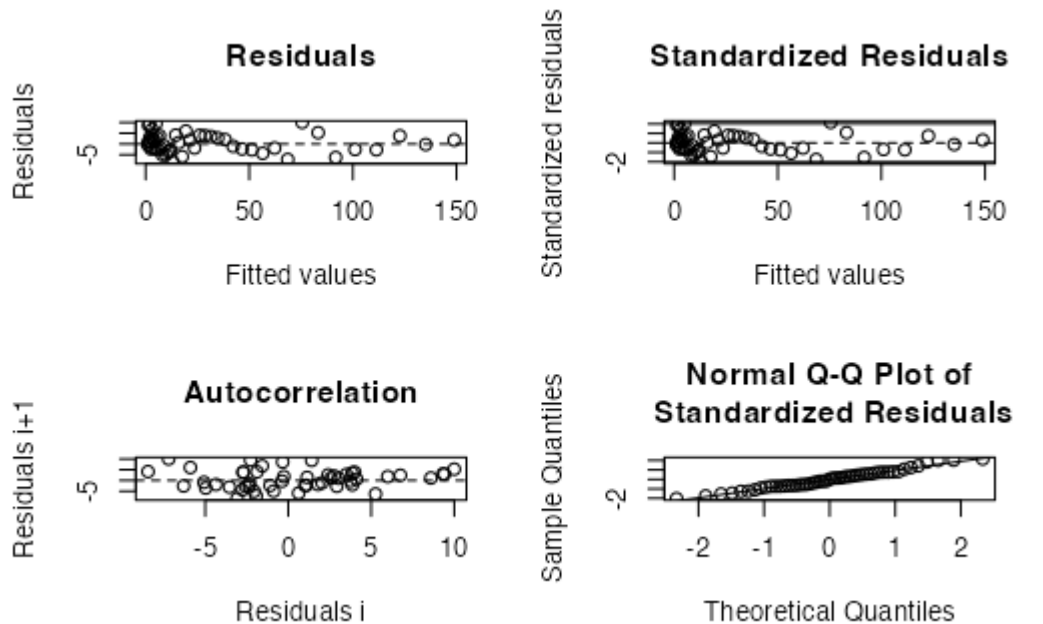
```
fit_exponential ← nls(response ~ y0*exp(k*predictor), data = growth,
  start = list(y0 = 0.1, k = 1))
```

```
ggplot(data = growth, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response") +
  geom_line(aes(y = predict(fit_exponential)), color = "red", size = 2)
```



Check assumptions

```
library(nlstools)
resids ← nlsResiduals(fit_exponential)
plot(resids)
```



- These plots determine if the residuals are normally distributed and have equal variance
- *Normal QQ* looks good
- *Residuals vs fitted* and *Standardized Residuals* even spread but slight fanning.
- With *Autocorrelation* we want random scatter around 0 – this indicates independence. Harder to meet with time-series data.
- Nonlinear models typically should meet assumptions because they are fitted specifically to the data.

Interpretation

```
summary(fit_exponential)
```

Formula: response ~ y0 * exp(k * predictor)

Parameters:

	Estimate	Std. Error	t value	Pr(> t)
y0	1.1694	0.1291	9.059	4.82e-12 ***
k	0.4847	0.0121	40.057	< 2e-16 ***

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

Residual standard error: 4.409 on 49 degrees of freedom

Number of iterations to convergence: 8

Achieved convergence tolerance: 1.192e-06

- The model is significant since the p-value is less than 0.05 for all parameters.
- If this were real data (e.g. population growth), the parameters themselves e.g. rate of change, are useful

- The parameterised model is:

$$y = 1.17 \cdot e^{-0.484x}$$

The R-squared value is not reported for nonlinear models as the sum of squares is not partitioned into explained and unexplained components. You can use the **residual standard error** and plots instead to compare between models.

A really bad guess

What if we don't estimate our parameters very well? R will either give an error or get there eventually.

Note the parameters and residual standard error are the same as the previous slide - but the
Number of iterations to convergence is higher.

```
fit_exponential ← nls(response ~ y0*exp(k*predictor), data = growth,  
start = list(y0 = 50, k = 1.5)) # totally bogus numbers  
  
summary(fit_exponential)
```

Formula: response ~ y0 * exp(k * predictor)

Parameters:

	Estimate	Std. Error	t value	Pr(> t)
y0	1.1694	0.1291	9.059	4.82e-12 ***
k	0.4847	0.0121	40.057	< 2e-16 ***

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

Residual standard error: 4.409 on 49 degrees of freedom

Number of iterations to convergence: 28

Achieved convergence tolerance: 2.003e-06



Tip

If an error pops up, try different starting values - the **rate of change** is most likely the problem.

Fitting the model with `SSexpf()`

- `SSexpf()` is from the `nltraa` package.
- It has the same formula as above – different names for parameters ($y_0 = a$, $k = c$) but we can re-define them to anything we want
- Reaches the same result but with less effort.

```
library(nltraa)
```

```
Attaching package: 'nltraa'
```

```
The following object is masked from 'package:lattice':
```

```
barley
```

```
fit_exponential_ss ← nls(response ~ SSexpf(predictor, y0, k), data = decay)
summary(fit_exponential_ss)
```

Formula: response ~ SSexpf(predictor, y0, k)

Parameters:

	Estimate	Std. Error	t value	Pr(> t)
y0	1.65486	0.04699	35.22	< 2e-16 ***
k	-0.06527	0.00590	-11.06	6.33e-15 ***

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

Residual standard error: 0.1471 on 49 degrees of freedom

Number of iterations to convergence: 5

Achieved convergence tolerance: 1.048e-06

Example: Fitting an asymptotic model

The equation

- There are multiple equations for asymptotic models, this is the equation that `SSasymp()` (base R) uses:

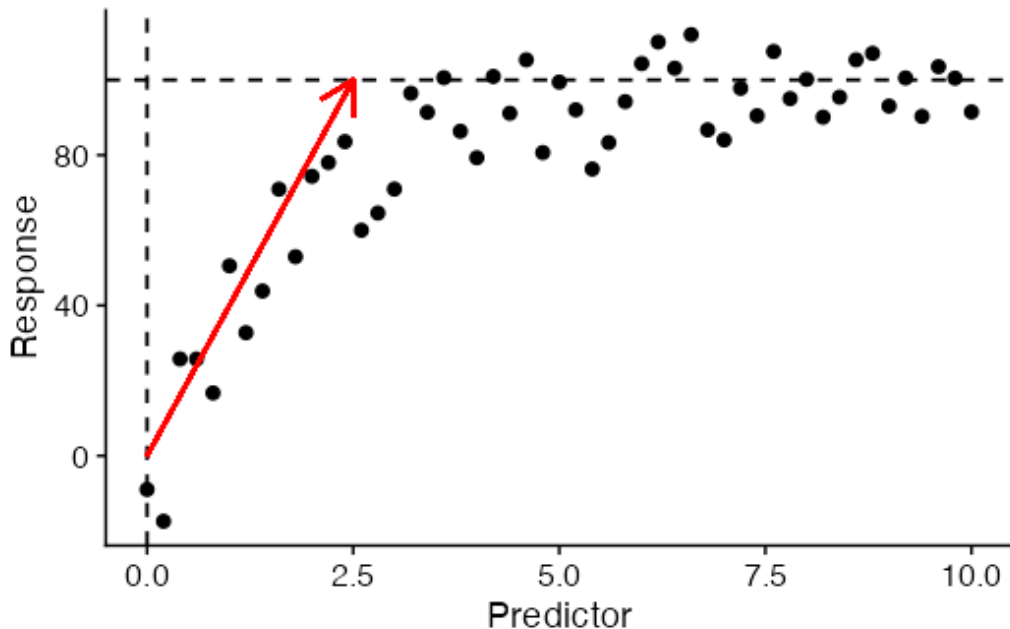
$$y = Asym + (R_0 - Asym) \cdot e^{-e^{lrc} \cdot x}$$

- R_0 is value of y when $x = 0$.
- $Asym$ is the upper limit: the maximum value of y .
- lrc is the rate of change: the rate at which y approaches the upper limit.

...

```
ggplot(data = asymptotic, aes(x = predictor, y = response)) +
  geom_point() +
  geom_hline(yintercept = 100, linetype = "dashed") +
  geom_vline(xintercept = 0, linetype = "dashed") +
  ## plot the rate
  geom_segment(aes(x = 0, y = 0, xend = 2.5, yend = 100),
               arrow = arrow(length = unit(0.5, "cm")),
               color = "red") +
  labs(x = "Predictor", y = "Response")
```

```
Warning in geom_segment(aes(x = 0, y = 0, xend = 2.5, yend = 100), arrow = arrow(length =  
unit(0.5, : All aesthetics have length 1, but the data has 51 rows.  
i Please consider using `annotate()` or provide this layer with data containing  
a single row.
```



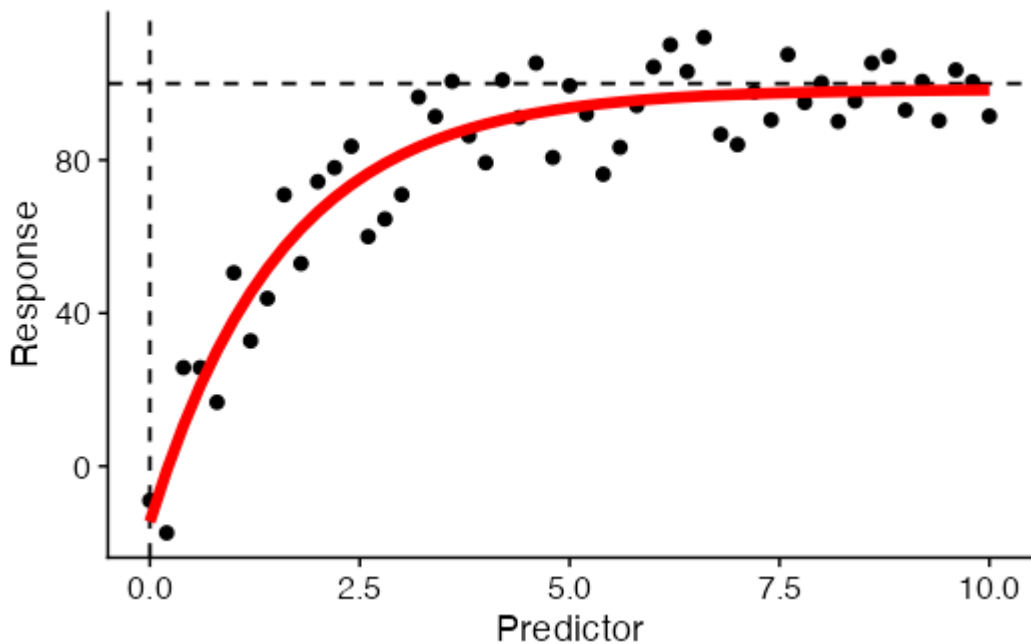
Some plausible estimates – $R_0 = 0$, $Asym = 100$, $lrc = 0.8$.

Fit model

```
# fit_asymptotic ← nls(response ~ Asym + (R0-Asym)*(exp(-exp(lrc)*predictor)), data = asymptotic,
#   start = list(R0 = 0, Asym = 100, lrc = 0.8))

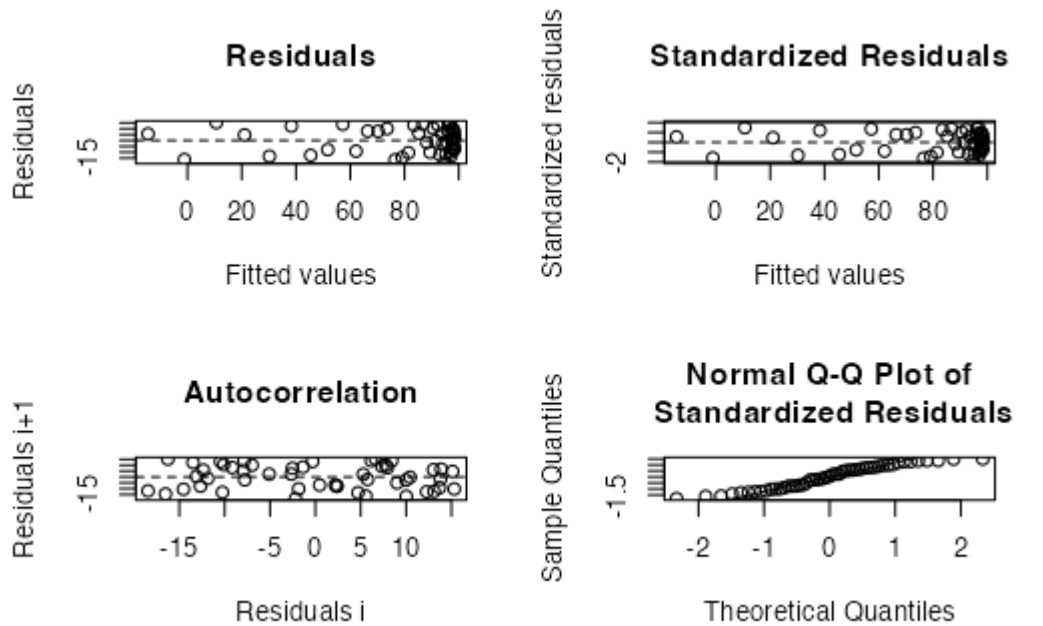
fit_asymptotic ← nls(response ~ SSasymp(predictor, Asym, R0, lrc), data = asymptotic)

ggplot(data = asymptotic, aes(x = predictor, y = response)) +
  geom_point() +
  geom_hline(yintercept = 100, linetype = "dashed") +
  geom_vline(xintercept = 0, linetype = "dashed") +
  labs(x = "Predictor", y = "Response") +
  geom_line(aes(y = predict(fit_asymptotic)), color = "red", size = 2)
```



Check assumptions

```
library(nlstools)
resids ← nlsResiduals(fit_asymptotic)
plot(resids)
```



Interpretation

```
summary(fit_asymptotic)
```

Formula: response ~ SSasymp(predictor, Asym, R0, lrc)

Parameters:

	Estimate	Std. Error	t value	Pr(> t)
Asym	98.5204	2.2852	43.113	< 2e-16 ***
R0	-14.5176	6.6416	-2.186	0.03374 *
lrc	-0.4626	0.1134	-4.079	0.00017 ***

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

Residual standard error: 10.21 on 48 degrees of freedom

Number of iterations to convergence: 0

Achieved convergence tolerance: 3.341e-07

- The model is significant since the p-value is less than 0.05 for all parameters.

- If this were real data (e.g. population growth), the parameters themselves e.g. rate of change, are useful
- The parameterised model is:

$$y = 98.5 + (-14.5 - 98.5) \cdot e^{-e^{-0.463} \cdot x}$$

Example: fitting a logistic model

The equation

There are multiple equations for logistic models, but they all have an ‘S’ or sigmoid shape. The equation that `SSlogis()` (base R) assumes y is positive and uses:

$$y = \frac{Asym}{1 + e^{\frac{xmid-x}{scal}}}$$

where

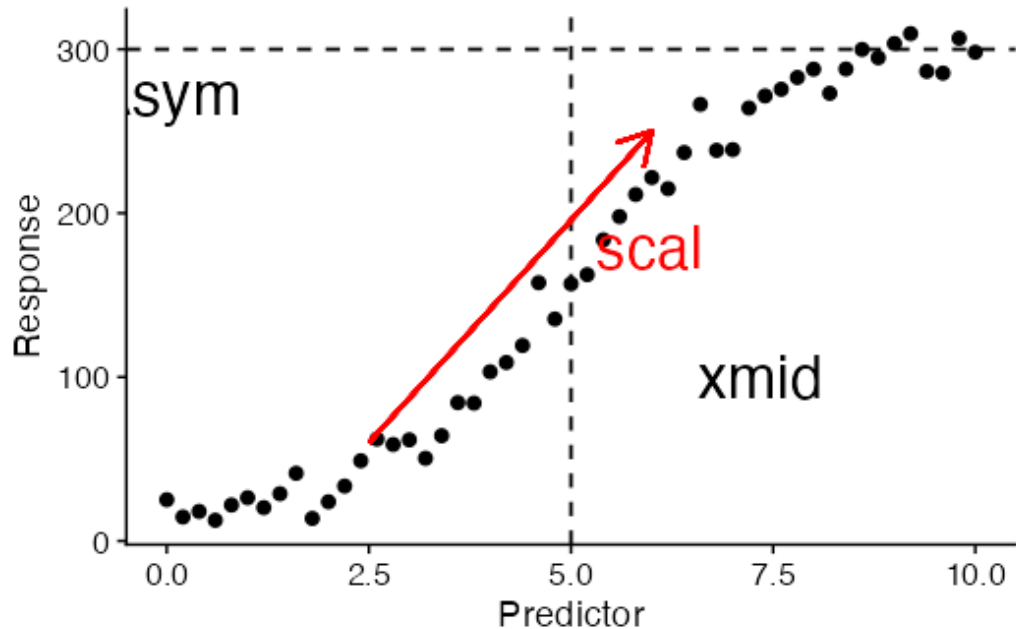
- $Asym$ is the upper limit: the maximum value of y .
- $xmid$ is the value of x when y is halfway between the lower and upper limits.
- $scal$ is the rate of change: the rate at which y approaches the upper limit.

```
ggplot(data = logistic, aes(x = predictor, y = response)) +
  geom_point() +
  labs(x = "Predictor", y = "Response") +
  geom_hline(yintercept = 300, linetype = "dashed") +
  geom_vline(xintercept = 5, linetype = "dashed") +
  # label the lines above
  annotate("text", x = 0, y = 300, label = "Asym", size = 8, vjust = 1.5) +
```

```
annotate("text", x = 5, y = 100, label = "xmid", size = 8, hjust = -1) +
## plot the rate
geom_segment(aes(x = 2.5, y = 60, xend = 6, yend = 250),
             arrow = arrow(length = unit(0.5, "cm")),
             color = "red") +
# label the rate
annotate("text", x = 4, y = 180, label = "scal", size = 8, colour = "red", hjust = -1)
```

Warning in geom_segment(aes(x = 2.5, y = 60, xend = 6, yend = 250), arrow = arrow(length =
unit(0.5, : All aesthetics have length 1, but the data has 51 rows.

i Please consider using `annotate()` or provide this layer with data containing
a single row.



Some starting values would be $Asym = 300$, $xmid = 5$, $scal = 1$.

Fit model

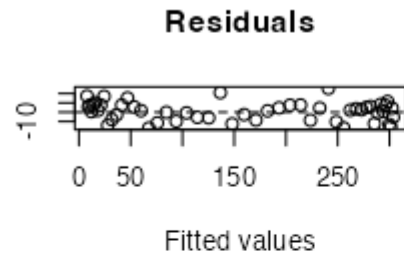
Estimating the parameters or using the self-starting function `SSlogis()` gives a near-identical result.

```
# fit_logistic ← nls(response ~ Asym/(1+exp((xmid-predictor)/scal)), data = logistic,
#   start = list(Asym = 300, xmid = 5, scal = 1))

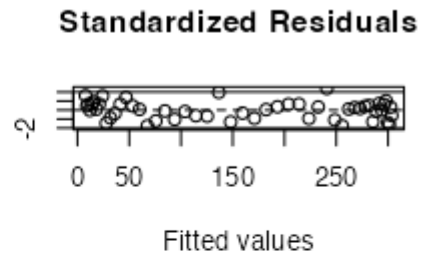
fit_logistic ← nls(response ~ SSlogis(predictor, Asym, xmid, scal), data = logistic)

resids ← nlsResiduals(fit_logistic)
plot(resids)
```

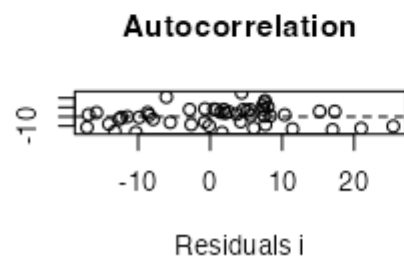
Residuals



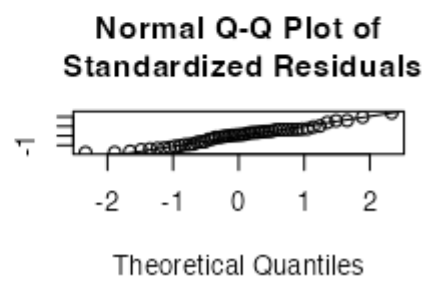
Standardized residuals



Residuals $i+1$



Sample Quantiles



Interpretation

`SSlogis()` guessed the parameters on the first try.

```
summary(fit_logistic)
```

Formula: response ~ SSlogis(predictor, Asym, xmid, scal)

Parameters:

	Estimate	Std. Error	t value	Pr(> t)
Asym	310.64727	4.62579	67.16	<2e-16 ***
xmid	4.92715	0.07142	68.99	<2e-16 ***
scal	1.34877	0.05418	24.90	<2e-16 ***

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

Residual standard error: 10.22 on 48 degrees of freedom

Number of iterations to convergence: 1

Achieved convergence tolerance: 6.632e-07

- The model is significant since the p-value is less than 0.05 for all parameters.
- If the model visually fits well and relationship has reasoning (parameter significance not always important).
- The parameterised model is:

$$y = \frac{310}{1 + e^{\frac{4.93-x}{1.35}}}$$

How do we know which model is better? (Advanced)

Note: this is non-examinable content but might be useful for your project.

Example: polynomial regression

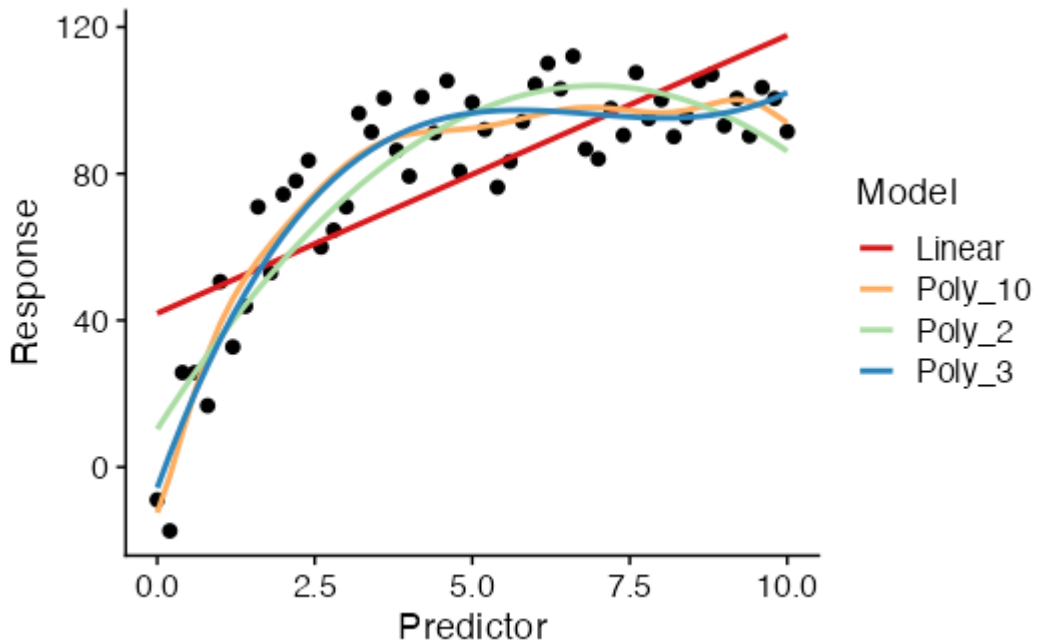
```
library(tidyr)

# Create a new data frame with predictor values and model predictions
predictions ← data.frame(
  predictor = asymptotic$predictor,
  Linear = predict(lin_fit),
  Poly_2 = predict(poly2_fit),
  Poly_3 = predict(poly3_fit),
  Poly_10 = predict(poly10_fit)
)

# Reshape the data to long format
predictions_long ← predictions %>%
  pivot_longer(cols = -predictor, names_to = "Model", values_to = "response")

# Plot the data
ggplot(predictions_long, aes(x = predictor, y = response, color = Model)) +
  geom_point(data = asymptotic, aes(x = predictor, y = response), inherit.aes = FALSE) +
  geom_line(linewidth = 1) +
```

```
labs(x = "Predictor", y = "Response") +  
  scale_color_brewer(palette = "Spectral")
```



Prediction quality

We can use prediction quality metrics to compare the fits.

- Akaike information criterion (AIC) and Bayesian information criterion (BIC).
 - ▶ Useful for comparing model fits.
 - ▶ Has a penalty for more predictors
- Residual standard error, residual sum of squares (`deviance(mod)`), root mean squared error (RMSE) and mean absolute error (MAE).
 - ▶ Essentially the difference between observed and predicted (residuals).
 - ▶ RMSE penalises larger residuals.

AIC and BIC

Use the `broom` package to extract the AIC and BIC values from the model fits.

```
library(broom)
# collect all polynomial fits into a single tibble using glance
poly_fits ← tibble(
  model = c("linear", "poly2", "poly3", "poly10"),
  fit = list(lin_fit, poly2_fit, poly3_fit, poly10_fit)) %>%
  mutate(glance = map(fit, glance)) %>%
  unnest(glance) %>%
  select(model, AIC, BIC)
poly_fits
```

```
# A tibble: 4 × 3
  model     AIC     BIC
  <chr>   <dbl>   <dbl>
1 linear   453.   459.
2 poly2    409.   416.
3 poly3    392.   402.
4 poly10   402.   425.
```

- The smaller the AIC or BIC, the better the fit compared to other models.

Calculate RMSE and MAE

```
predictions ← data.frame(  
  observed = asymptotic$response,  
  Linear = predict(lin_fit),  
  Poly_2 = predict(poly2_fit),  
  Poly_3 = predict(poly3_fit),  
  Poly_10 = predict(poly10_fit)  
)  
  
errors ← predictions %>%  
  pivot_longer(cols = -observed, names_to = "Model", values_to = "Predicted") %>%  
  group_by(Model) %>%  
  summarise(  
    RMSE = sqrt(mean((observed - Predicted)^2)),  
    MAE = mean(abs(observed - Predicted)))  
)  
  
knitr::kable(errors, digits=2, caption = "Comparison of RMSE and MAE for different models")
```

Model	RMSE	MAE
Linear	19.38	15.17
Poly_10	9.82	8.57
Poly_2	12.30	9.88
Poly_3	10.25	8.83

- From the results, the polynomial to the degree of 10 has the lowest error - but visually we know it is overfitting, and the cubic polynomial is more parsimonious.
- We can say the model has a prediction error of 10.25 units (RMSE) and 8.83 units (MAE).

i Note

Both the RMSE and MAE measure error on the same scale as the response variable. e.g. if the response variable is in kg, the error will be in kg.

Summary

- With nonlinear relationships, there are three possible approaches:
 1. **Linearise** the relationship by transforming:
 - Fit: easy
 - Interpret: difficult
 2. Add **polynomial** terms:
 - Fit: easy
 - Interpret: difficult
 3. Fit the model using a **nonlinear** algorithm:
 - Fit: difficult
 - Interpret: easy



...

- Nonlinear models:
 - Useful for modelling more complex relationships. Require some understanding of the underlying relationship and equations.

- ▶ Mainly for prediction rather than interpreting relationships.
- ▶ Self-starting functions have limited pre-defined formulas.
- ▶ Assumptions **INE**.

Thanks!

This presentation is based on the [SOLES Quarto reveal.js template](#) and is licensed under a [Creative Commons Attribution 4.0 International License](#).