

# Introduction to Generalised Linear Models for Ecologists

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[https://github.com/niamhmimmnagh/GLME01---  
Introduction-to-Generalised-Linear-Models-for-  
Ecologists](https://github.com/niamhmimmnagh/GLME01---Introduction-to-Generalised-Linear-Models-for-Ecologists)

# Binary Data

Many real-world problems involve binary (yes/no, success/failure) outcomes:

- Did a patient survive? (yes/no)
- Was the animal infected? (yes/no)
- Did the student pass the course? (yes/no)

$$Y_i = \begin{cases} 1, & \text{if success} \\ 0, & \text{if failure} \end{cases}$$

What is 'success'?

It's whatever you want it to be! It's not necessarily the 'best' outcome.

# Binary Data

When data is binary, there are only two possible outcomes, and so when we talk about the probability of each outcome, we have:

$$\begin{aligned}P(\text{success}) &= P(Y_{ij} = 1) = \pi_{ij} \\P(\text{failure}) &= P(Y_{ij} = 0) = 1 - \pi_{ij}\end{aligned}$$

$Y_i$  has a Bernoulli distribution:

$$Y_i \sim \text{Bernoulli}(\pi_i)$$

# Why Not Use a Linear Model?

Linear regression assumes:

- The response variable is continuous and unbounded.
- The relationship between predictors and the response is linear.

Problems with using it on binary data:

- Predictions can fall outside  $[0,1]$
- Error terms are heteroscedastic (non-constant variance)
- Residuals are not normally distributed

This leads to poor model performance and invalid inference.

# The Bernoulli GLM

$$Y_i \sim \text{Bernoulli}(\pi_i)$$

We want to model the success probabilities  $\pi_i$  as a function of predictors.

Can we simply write  $\pi_i = \beta_0 + \beta_1 x_{1i} + \dots + \beta_p x_{pi}$ ?

No!

Since the  $\beta$  coefficients are unbounded (they can take any real value from  $-\infty$  to  $\infty$ ), this would result in unbounded  $\pi_i$  values

But  $\pi_i$  are probabilities, and so have to be bounded in the  $(0,1)$  interval

So we need a link function that maps the  $(0,1)$  interval to the real line.

# The Bernoulli GLM

$$Y_i \sim \text{Bernoulli}(\pi_i)$$
$$\log\left(\frac{\pi_i}{1 - \pi_i}\right) = \beta_0 + \beta_1 x_{1i} + \dots + \beta_p x_{pi}$$

The function  $\log\left(\frac{\pi_i}{1 - \pi_i}\right)$  can also be written as  $\text{logit}(\pi_i)$

‘logit’ stems from the words **logistic unit**, since its based on the cumulative distribution function of the logistic distribution

It is simply the natural logarithm of the odds

This is how we ensure predicted probabilities stay between 0 and 1.

# Probability vs. Odds

- Probability ( $\pi$ ) is the chance of an event occurring (range: 0 to 1)  
e.g.,  $\pi = 0.8$  means an 80% chance of success
- Odds compare the probability of success to the probability of failure.

$$Odds = \frac{\pi}{1 - \pi}$$

- For example, if  $\pi = 0.8$  then  $odds = \frac{0.8}{0.2} = 4$   
(4 to 1 odds of success)

$$\pi = \frac{Odds}{1 + Odds}$$

- If odds = 4, then  $\pi = \frac{4}{1+4} = \frac{4}{5} = 0.8$



# Interpreting Fixed-Effect Coefficients

$$\text{logit}(\pi) = \beta_0 + \beta_1 x = -0.5 + 1.2x$$

- $\beta_0 = -0.5$ : when  $x = 0$ , the log-odds of success are  $-0.5$ .
- Odds =  $e^{-0.5} \approx 0.607$ : when  $x = 0$ , the odds of success are 0.61 to 1 (success is less likely than failure).
- $\pi = \frac{0.607}{1+0.607} \approx 0.38$ : when  $x = 0$ , the probability of success is about 38%.
- $\beta_1 = 1.2$ : for each 1-unit increase in  $x$ , the log-odds increase by 1.2.
- Odds ratio =  $e^{1.2} \approx 3.32$ : each 1-unit increase in  $x$  multiplies the odds of success by about 3.3.
- At  $x = 1$ :  $\text{logit}(\pi) = -0.5 + 1.2(1) = 0.7 \rightarrow \text{Odds} = e^{0.7} \approx 2.01$
- $\pi = \frac{2.01}{1+2.01} \approx 0.67$ : increasing  $x$  by one unit raises probability from 38% to 67%.



# Assumptions:

## Independence of Observations

- Each  $Y_i$  is an independent Bernoulli trial given the predictors (no residual correlation across observations).
- If this assumption is violated, it will lead to underestimated SEs, too-small p-values, overconfident CIs, misleading inference.

### Common violations:

- Clusters/groups: students within classes, animals within herds, patients within hospitals
- Repeated measures: multiple rows per subject over time
- Spatial/temporal autocorrelation: nearby in space/time more similar than distant

# Assumptions:

## No Perfect Multicollinearity

- Predictors should not be perfectly correlated with each other.
- High collinearity leads to unstable coefficients and inflated standard errors.
- Check Variance Inflation Factor (VIF), and drop redundant predictors if needed. The VIF measures how much the variance of a regression coefficient is increased due to correlation with other predictors. A higher VIF means the predictor is more redundant with others. A rule of thumb is:
  - $VIF > 5 \rightarrow$  moderate collinearity
  - $VIF > 10 \rightarrow$  high collinearity
- Collinearity doesn't bias the model, but makes it hard to interpret.



# Assumptions: Sample Size

- Logistic regression needs enough events for stable estimates.
- The rule of thumb is  $\geq 10$  events per variable.
- For example, if we have 5 predictors, we need at least 50 events.
- Why?
- Small sample sizes can lead to large standard errors (wide confidence intervals), unstable odds ratios (sensitive to small data changes), and convergence issues (failure to estimate coefficients or infinite estimates).

# The Logistic Function

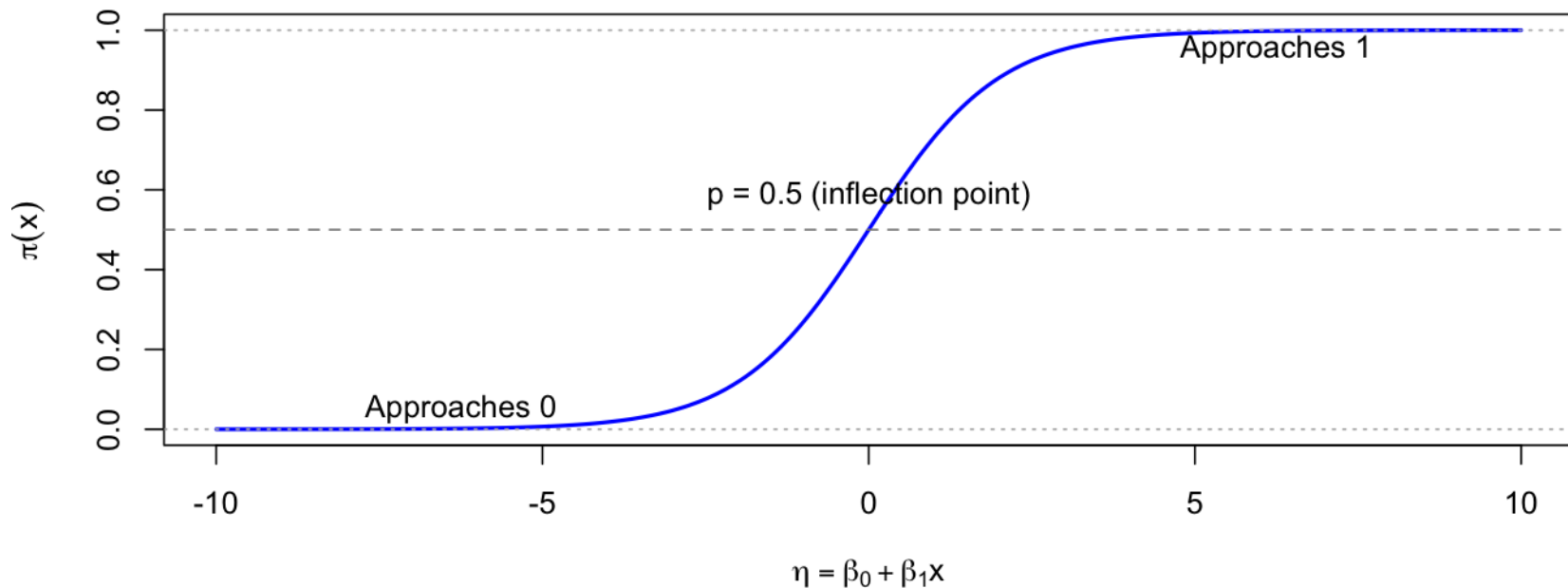
- Rewriting the logit model:

$$\pi = \frac{e^{\beta_0 + \beta_1 x}}{1 + e^{\beta_0 + \beta_1 x}}$$

- This is a sigmoid-shaped curve. The logistic curve:
  - Approaches 0 and 1 asymptotically.
  - Has an inflection point at  $p = 0.5$ .
  - Is nonlinear in probability space but linear in log-odds space.

# The Logistic Function

Sigmoid Shape of the Logistic Function



# Other Link Functions

- The logistic function is most commonly used for binary data, but it isn't the only choice. Other choices include:
  - Probit (normal CDF): assumes the latent propensity follows a normal distribution. It tends to give very similar results to logit but is sometimes preferred in fields like toxicology, psychometrics, or genetics where a normal latent variable is a natural assumption.
  - Complementary log-log (cloglog): asymmetric S-shape - changes faster near 0 than near 1. It's useful when the probability of an event increases rapidly and then levels off (e.g. survival models, time-to-event data, rare events).
- All are S-shaped, but with subtle differences

# Example:

## Disease Presence

- We want to estimate how an animal's age relates to the probability of disease presence and produce usable, age-specific risk estimates.

$$Y_i \sim \text{Bernoulli}(\pi_i)$$
$$\log \left( \frac{\pi_i}{1 - \pi_i} \right) = \beta_0 + \beta_1 \text{Age}_i$$

- Predictor: Age in years (we'll centre at 2 years so the intercept is interpretable)
- Baseline odds of disease presence at age 2 years:  $e^{\beta_0}$
- Age effect: each additional year multiplies odds of disease by  $e^{\beta_1}$  e.g., a 5-year difference  $\Rightarrow e^{5\beta_1}$  times the odds

# Coding Demo



# Nested Models

- Nested models are a pair of models where one is a special case of the other - i.e., the smaller model is completely contained within the larger model.
- The simpler model has fewer predictors, while the more complex model includes all the terms in the simple model plus additional predictors.
- If Model A is:  $\text{logit}(\pi) = \beta_0 + \beta_1 x$
- And Model B is:  $\text{logit}(\pi) = \beta_0 + \beta_1 x + \beta_2 z$
- Then Model A is nested inside Model B.

# Nested Models

- Comparing nested models lets us test whether the extra variables in the larger model significantly improve the model's fit.
- It helps us answer the question: “Do these new predictors help explain the outcome, beyond what we already had?”
- We can answer this question using the Likelihood Ratio Test, which compares how well each model explains the data, using the difference in log-likelihoods.

$$G^2 = -2(\log L_A - \log L_B)$$

# Coding Demo

# Analysing Model Fit

- Fitting a model is only the beginning.
- We need to verify that:
  - The model adequately describes the data.
  - The assumptions are not violated.
  - No single observation unduly influences results.

# Model Comparison: Pseudo-R<sup>2</sup> Measures

- Logistic models use pseudo-R<sup>2</sup> measures to approximate explanatory power:

## McFadden's R<sup>2</sup>:

$$R^2 = 1 - \frac{\log L_{Model}}{\log L_{null}}$$

- McFadden's R<sup>2</sup> compares the likelihood of the fitted model against the null (intercept-only) model. Values closer to 1 indicate better improvement over the null, but they are usually much lower than R<sup>2</sup> in linear regression. 0.2-0.4 already signals a well-fitting logistic model.

# Model Comparison: Deviance

- Deviance measures how well the model fits the data.
- It's based on the log-likelihood of the model.
$$\text{Deviance} = -2(\log \text{Likelihood})$$
- A lower deviance indicates a better fit.
- Deviance compares the likelihood of your model to a saturated model (a model that fits the data perfectly).
- Deviance differences between models can be used for hypothesis testing: comparing the deviance of a full model and a reduced model. A large drop in deviance means the added predictors improve model fit significantly

# Model Comparison: Deviance

In the model output, you often see:

- **Null deviance:** Fit of the intercept-only model (no predictors).
- **Residual deviance:** Fit of the model with predictors.

$$\text{Improvement} = \text{Null deviance} - \text{Residual deviance}$$

If residual deviance is much lower than null deviance, your predictors are explaining the variation in the response, and adding predictors improves the model fit compared to the intercept-only model.

# Model Comparison: AIC

AIC (Akaike Information Criterion) helps compare non-nested models or models with different numbers of predictors.

$$AIC = -2(\log \text{Likelihood}) + 2k$$

Where, the log likelihood measures how well the model fits the data, and  $k$  = number of parameters (complexity penalty).

Lower AIC is better.

It balances:

- **Fit** (how well the model explains the data)
- **Parsimony** (fewer predictors is better if fit is similar)





# Residuals

- Residuals measure the difference between observed and predicted outcomes. In logistic regression, residuals are calculated from the difference between the observed outcome (0/1) and the fitted probability.
- Logistic regression uses several types of residuals:
  - Deviance residuals
  - Pearson residuals
  - Standardised residuals
- Each highlights different types of model issues.



# Deviance Residuals

- Deviance residuals measure how much each observation contributes to the model's deviance. They are used to identify poorly fit observations, and plotted to detect non-random patterns.
- Large deviance residuals (positive or negative) indicate observations the model predicts poorly.
- Clusters or systematic trends in a residual plot suggest model misspecification (e.g., missing predictors, wrong link function, or non-linearity).
- In a well-fitting model, deviance residuals should look roughly random with no clear pattern.
- Plotting deviance residuals against the fitted values helps check for systematic deviations - any clear curve or structure suggests the model is mis-specified, while a random scatter around zero indicates a good fit.

# Pearson Residuals

- Pearson residuals are analogous to residuals in linear regression.

$$r_i = \frac{y_i - p_i}{\sqrt{p_i(1 - p_i)}}$$

- They measure the difference between observed and expected values. They can be used to assess overall model fit.
- Large Pearson residuals highlight observations that deviate strongly from the model's predictions. Patterns in Pearson residual plots may suggest overdispersion, missing covariates, or incorrect model structure. In a good model, Pearson residuals should be roughly standard normal, centered at zero with variance close to one.

# Standardised Residuals

- Standardised residuals are residuals divided by their estimated standard deviation.
- This rescales residuals to be approximately comparable across observations. Raw residuals aren't comparable because their variance differs across observations (due to heteroscedasticity and leverage). Standardising puts them on a common scale of “standard deviations,” making comparison meaningful.
- Useful for detecting outliers and influential data points.
- Values greater than  $|2|$  suggest potential outliers (check context).
- Values greater than  $|3|$  are often considered highly unusual.
- Always interpret in context of sample size, leverage, and model fit.

# When to Use Each Residual

- **Deviance residuals:** best for checking individual fit and influence.
  - They are derived from the model's deviance and reflect how much each observation contributes.
  - Large absolute values suggest poor fit or potential leverage points.
- **Pearson residuals:** helpful for overall goodness-of-fit statistics.
  - They are based on the difference between observed and expected counts, scaled by variance.
  - Useful for testing model adequacy in aggregate.
- **Standardised residuals:** great for detecting outliers.
  - They rescale residuals to account for differing variances across observations.
  - Values beyond  $|2|$  or  $|3|$  may signal unusual observations worth investigating.
- No single residual tells the full story – use multiple diagnostics together.



# Leverage

- Leverage measures how far an observation's predictor values are from the average predictor values.
- High-leverage points have unusual predictor values (not necessarily unusual response values).
- These points can exert strong influence on the model's fitted values.
- Examining leverage helps identify observations that disproportionately affect regression estimates.
- As a rule of thumb, leverage values greater than  $2p/n$  (where  $p$  is the number of predictors including the intercept, and  $n$  the sample size) may be considered high.

# Influence

- Influence considers both leverage (unusual predictor values) and residual size (poor fit).  
An influential point is one that, if removed, would substantially change the model.
- Influence is assessed using measures like Cook's Distance.
- We look at influential points because they can disproportionately affect parameter estimates and conclusions.
- They are not automatically “bad” - they may reflect real structure or unusual but valid cases.
- Best practice is to investigate them: check for data errors, assess context, and compare models with and without them.

# Cook's Distance

- Measures the influence of a single observation on all fitted values.
- Combines residual size (fit) and leverage (position in predictor space).
- An observation with a large residual and a high leverage will have a high Cook's Distance.
- Interprets how much model coefficients would change if that observation were removed.
- Rule of thumb: values  $> 0.5$  or  $1$  may indicate influential points.
- Should not be used mechanically - investigate influential cases for possible data issues or meaningful outliers.





# Coding Demo