Statistical Modelling in Immunology

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We will cover the following topics:

- A recap on the normal model
- Generalized linear models and extensions
- Generalized additive models for location, scale, and shape
- Mixed models

GAMLSS 000

The Normal Model: A Recap

The Normal Model

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- \blacksquare Y_i is a response variable associated with experimental or observational unit i
- We assume it comes from a certain probability distribution with pmf/pdf f and vector of parameters θ
- In general, one of the parameters in θ is the mean of the distribution
- We also have predictors x_i we are interested in studying
- We may link it to a parameter of interest, typically the mean of the distribution

For the normal model, we typically write for each observation y_i :

$$y_i = \beta_0 + \beta_1 x_{1i} + \dots + \beta_p x_{pi} + \varepsilon_i$$

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

- **Each** β coefficient represents the expected mean change in y for a 1-unit increase in its associated predictor
- We can show that, from the equation above, the expected value of Y_i is

$$\mathsf{E}[Y_i] = \beta_0 + \beta_1 x_{1i} + \dots + \beta_p x_{pi}$$

and the variance is

$$Var(Y_i) = \sigma^2$$

■ Therefore, we are assuming the variance is constant

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- The error notation has its advantages. . .
- However, let's switch things up a bit

The Normal Model 0000000000000

What is Statistical Modelling?

The Normal Model 0000000000000

What is a Statistical Model?

It's all about Probability!

Building blocks

- \blacksquare Response variable (Y)
- Probability distribution
- 3 Parameters of interest

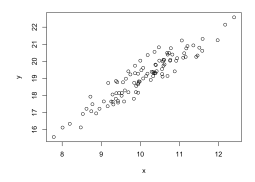
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Building blocks

- **1** Response variable (Y)
- Probability distribution
- \blacksquare Parameters of interest \leftarrow covariates / predictors

- $lackbox{ iny Y_i is a response variable associated with experimental or observational unit $i$$
- We assume it comes from a certain probability distribution with pmf/pdf f and vector of parameters θ
- Very often one of the parameters in θ is the mean of the distribution (or a function of the mean)
 - e.g.¹ for the normal distribution, $\theta = (\mu, \sigma^2)^{\top}$, where μ is the mean of the distribution
 - e.g.² for the Poisson distribution, $\theta = \mu$, where μ is the mean of the distribution
- lacktriangle We also have predictors x_i we are interested in studying
- We may link these predictors to any parameter of interest, but we typically do it for the mean

Example



Example

$$Y_i \sim \mathsf{N}(\mu_i, \sigma^2)$$

 $\mu_i = \beta_0 + \beta_1 x_i$

Example

The Normal Model

$$Y_i \sim \mathsf{N}(\mu_i, \sigma^2)$$

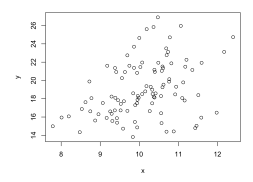
 $\mu_i = \beta_0 + \beta_1 x_i$

What about

- normality of residuals?
- homogeneity of variances?

This only makes sense for the model above!

What if you had



Maybe then you'd assume

$$Y_i \sim \mathsf{N}(\mu_i, \sigma_i^2)$$

$$\mu_i = \beta_0 + \beta_1 x_i$$

$$\log \sigma_i^2 = \gamma_0 + \gamma_1 x_i$$

No homogeneity of variances here!

The Normal Model

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Y can be assumed to have any distribution

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Y can be assumed to have any distribution Why is the normal distribution used so often then?

GLMs

- Multiple linear regression: a normal model with the identity link (Legendre, Gauss, Galton, 19th century)
- Analysis of variance (ANOVA): a normal model with the identity link (Fisher, 1918)
- Analysis of dilution assays: a binomial model with the complementary log-log link (Fisher, 1922)
- The exponential family class of distributions (Fisher, 1934)
- Probit analysis: a binomial distribution with the probit link (Bliss, 1935)
- Logistic regression: a binomial distribution with the logit link (Berkson, 1944; Dyke and Patterson, 1952)
- Item analysis: a Bernoulli distribution with the logit link (Rasch, 1960)
- Log-linear models: a Poisson distribution with the log link (Birch, 1963)
- Regression for survival data: an exponential distribution with the inverse or log links (Feigl and Zelen, 1965; Zippin and Armitage, 1966; Gasser, 1967)
- Inverse polynomials: a gamma distribution with the inverse link (Nelder, 1966)

A Brief History of GLMs

The Normal Model

J. R. Statist. Soc. A, (1972), 135, Part 3, p. 370 370

Generalized Linear Models

By J. A. Nelder and R. W. M. Wedderburn

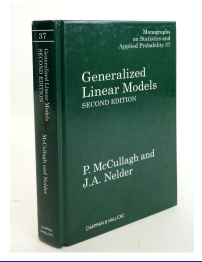
Rothamsted Experimental Station, Harpenden, Herts

SUMMARY

The technique of iterative weighted linear regression can be used to obtain maximum likelihood estimates of the parameters with observations distributed according to some exponential family and systematic effects that can be made linear by a suitable transformation. A generalization of the analysis of variance is given for these models using log-likelihoods. These generalized linear models are illustrated by examples relating to four distributions; the Normal, Binomial (probit analysis, etc.), Poisson (contingency tables) and gamma (variance components).

The implications of the approach in designing statistics courses are discussed.

Keywords: ANALYSIS OF VARIANCE; CONTINGENCY TABLES; EXPONENTIAL FAMILIES; INVERSE POLYNOMIALS; LINEAR MODELS; MAXIMUM LIKELIHOOD; QUANTAL RESPONSE; REGRESSION; VARIANCE COMPONENTS; WEIGHTED LEAST SOUARES



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- The generalized linear model can be defined using three components:
- **1** The random component: a distribution belonging to the exponential family
- The systematic component: a linear predictor
- The link function: a function that links the mean to the linear predictor

The Generalized Linear Model

GI Ms

- The generalized linear model can be defined using three components:
- The random component: a distribution belonging to the exponential family
- The systematic component: a linear predictor
- The link function: a function that links the mean to the linear predictor
- More specifically, we have independent random variables $Y_i, i = 1, ..., n$
- The linear predictor can be written as $\eta = X\beta$ where X is the $n \times (p+1)$ design (or model) matrix and $\beta = (\beta_0, \beta_1, \dots, \beta_n)^{\top}$ is the vector of model coefficients
- The link function $g(\cdot)$ relates the mean μ_i to η_i , i.e. $q(\mu_i) = \eta_i = \mathbf{x}_i^{\top} \boldsymbol{\beta}$, where \mathbf{x}_i is the *i*-th row of \mathbf{X}

The Simple Linear Regression Model as a GLM

- Random component: $Y_i \sim N(\mu_i, \sigma^2)$
- 2 Systematic component: $\eta_i = \beta_0 + \beta_1 x_{1i}$
- 3 Link function: $g(\mu_i) = \mu_i$, the *identity* link

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 - More simply put:

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 - More simply put:

The Normal Model

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■ Example: IFNG gene expression data

Half-Normal Plots with a Simulated Envelope

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- One way of empirically checking whether a GLM (or any¹ statistical model, in fact), is to use a graphical technique called half-normal plot with a simulated envelope
- It consists in plotting ordered residuals in absolute value versus expected order statistics of the half-normal distribution, and adding a simulated envelope based on the fitted model
- The envelope serves as a guide to what expect if the observed data are a plausible realisation of the fitted model
- In R it is implemented as the hnp package
- If the majority of points lie within the simulated envelope, the model can be considered to be well-fitted to the data

¹as long as response variables can be simulated from it

GAMs

GAMs

The Normal Model

■ The polynomial and spline regression models can be regarded as special cases of a more general type of regression model known as a generalized additive model (GAM)

GAMs

Example of a normal GAM:

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

 $\mu_i = f_1(x_{1i}) + f_2(x_{2i}) + f_3(x_{3i}, x_{4i})$

 $f_n(\cdot)$ are smooth functions

GAMLSS

GAMLSS

The Normal Model

 Generalized Additive Models for Location, Scale, and Shape (GAMLSS: Rigby and Stasinopoulos, 2005) are a very flexible semi-parametric modelling framework

$$Y \sim f(\mu, \sigma, \nu, \tau)$$

- Includes many distributions with up to 4 parameters
- Allows for distributional regression, i.e. all parameters can be modelled with covariates
- Spline and loess smoothing, as well as random effects allowed
- Unified framework for model diagnostics

GAMLSS

- The gamlss package is the main implementation
- Extra features available through companion packages
- The package gamlss.add includes extra features, such as the inclusion of regression trees in the linear predictor for any parameter of interest

Mixed Models

Mixed Models

- Mixed models are a broad class of models that are applied to data that consist of sub-groups, or clusters
- Mixed models have many other names in the literature and in practice
- You may have heard/seen: multilevel models, hierarchical models, conditional models, random effects models
- In practice, these are all synonyms, but each name highlights a different part of the construction of a mixed model
- The defining feature of multilevel models is that they are models of models²

²Andrews, M. (2021). Doing Data Science in R: An Introduction for Social Scientists. SAGE Publishing, London, UK.

Fixed vs Random effects

- It is important to understand the conceptual and practical differences between fixed versus random effects.
- We typically use fixed effects when we assume there is a systematic difference between levels of a factor, or a trend over time or space.
- We typically use random effects when we assume that what we observe is a random sample from a population.
 - reflect design
 - accommodate extra-variability

- Expression data from 559 genes $(g_i, i=1,\ldots,559)$, 805 subjects $(j=1,\ldots,805)$, and 7 stimuli $(s_k, k=1,\ldots,7)$
- Random effects model:

$$Y_{ijk}|g_i \sim \mathsf{N}(\mu_{ijk}, \sigma^2)$$

 $g_i \sim \mathsf{N}(0, \sigma_g^2)$
 $\mu_{ijk} \sim \beta_0 + g_i$

- Expression data from 559 genes $(g_i, i = 1, ..., 559)$, 805 subjects (j = 1, ..., 805), and 7 stimuli $(s_k, k = 1, ..., 7)$
- Mixed effects model:

$$Y_{ijk}|g_i \sim \mathsf{N}(\mu_{ijk}, \sigma^2)$$

 $g_i \sim \mathsf{N}(0, \sigma_g^2)$
 $\mu_{ijk} \sim s_k + g_i$

Example: Gene expression data

- **Expression** data from 559 genes $(g_i, i = 1, \dots, 559)$, 805 subjects (j = 1, ..., 805), and 7 stimuli $(s_k, k = 1, ..., 7)$
- Random effects per stimulus:

$$\begin{array}{ccc} Y_{ijk}|g_i & \sim & \mathsf{N}(\mu_{ijk},\sigma^2) \\ g_{ik} & \sim & \mathsf{N}(0,\sigma_k^2) \\ \mu_{ijk} & \sim & s_k + g_{ik} \\ \mathsf{Corr}(g_{ik},g_{ik'}) & = & 0 \end{array}$$