



# ptmixed: an R package for flexible modelling of overdispersed longitudinal counts

Mirko Signorelli<sup>1</sup>

Twitter: @signormirko

Joint work with Roula Tsonaka<sup>1</sup> and Pietro Spitali<sup>2</sup>

<sup>1</sup> Department of Biomedical Data Sciences, Leiden University Medical Center

<sup>2</sup> Department of Human Genetics, Leiden University Medical Center

June 19, 2020  
e-Rum2020



Leiden University  
Medical Center



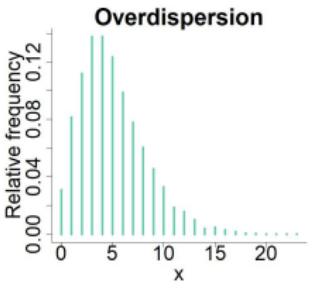
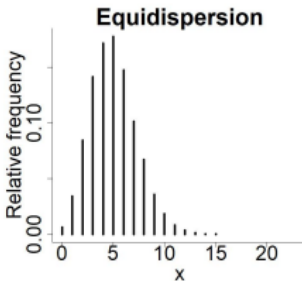
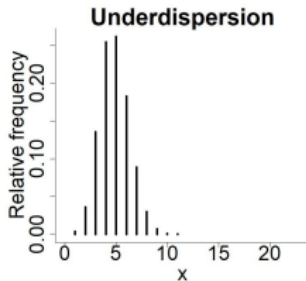
Universiteit  
Leiden

Twitter: @signormirko

Slides: [bit.ly/ptmixed](http://bit.ly/ptmixed)

# Count data come in different shapes

- ▶ Count data typically classified as
  - ▶ Underdispersed:  $\text{Var}(X) < E(X)$
  - ▶ Equidispersed:  $\text{Var}(X) = E(X)$
  - ▶ Overdispersed:  $\text{Var}(X) > E(X)$



- ▶ Most common models for overdispersed counts: **negative binomial (NB)** GLM and GLMM

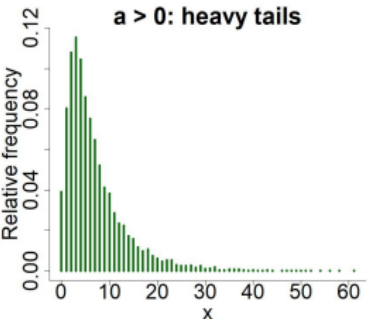
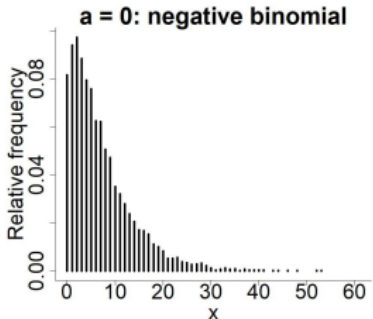
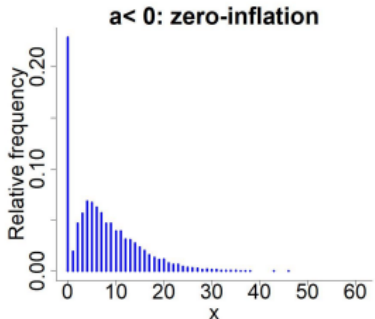
# Poisson-Tweedie: fifty shades of overdispersion



- ▶ El-Shaarawy et al. (2011) showed that NB can't capture well different levels of **zero-inflation (ZI)** and **heavy-tails (HT)** commonly observed with overdispersed counts
- ▶ Alternative: use the **Poisson-Tweedie (PT)** distribution

$$Y \sim PT(\mu, D, a)$$

- ▶ Power  $a \leq 1$  models extra ZI and HT for given dispersion  $D$ :





Applicability of Poisson-Tweedie:

- ▶ cross-sectional data → PT GLM (Esnaola et al., 2013) ✓
- ▶ longitudinal data → no GLMM extension ✗

Our proposal: ptmixed

- ▶ We propose a **Poisson-Tweedie GLMM** to flexibly model **longitudinal counts** with different levels of **zero-inflation** and **heavy-tails**
- ▶ Implementation: R package **ptmixed** (published on CRAN)



# Estimation of the PT mixed model

## Poisson-Tweedie GLMM

$$Y_{ij} \mid v_i \sim \text{PT}(\mu_{ij}, D, a)$$

$$\log(\mu_{ij}) = x_{ij}^T \beta + z_{ij}^T v_i$$

$$v_i \stackrel{\text{i.i.d.}}{\sim} N(0, \Sigma)$$

where  $\beta \rightarrow$  fixed effects,  $v_i \rightarrow$  random effects

How to estimate this model?

- ▶ Likelihood evaluation: approximation of PT pmf (Esnaola et al., 2013) + adaptive Gauss-Hermite quadrature
- ▶ Model estimation: maximum likelihood estimation

More details → Signorelli et al. (2020, in press), [arXiv:2004.11193](https://arxiv.org/abs/2004.11193)



# Data preparation

## 1) Load package and data

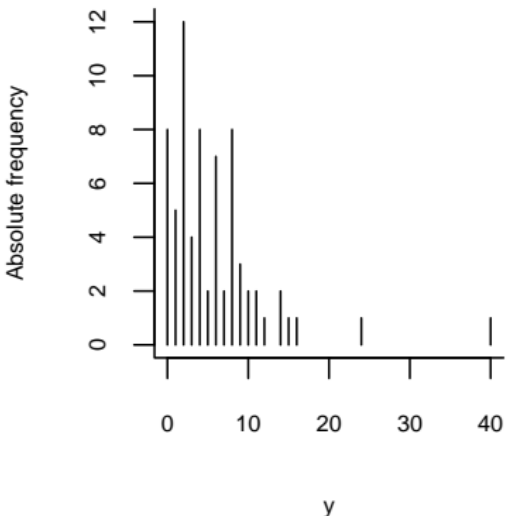
```
library(ptmixed)  
head(data.long)
```

```
##   y id  group time  
## 1 6  1 placebo  0  
## 2 0  1 placebo  1  
## 3 2  1 placebo  2  
## 4 1  1 placebo  3  
## 5 1  1 placebo  4  
## 6 12 2 placebo  0
```

- ▶  $y \rightarrow$  response variable
- ▶  $id \rightarrow$  subject id ( $1, \dots, n$ )
- ▶  $group \rightarrow$  2 groups (treated / placebo)
- ▶  $time \rightarrow$  5 time points ( $0 \rightarrow 4$ )

## 2) Visualize distribution of the response

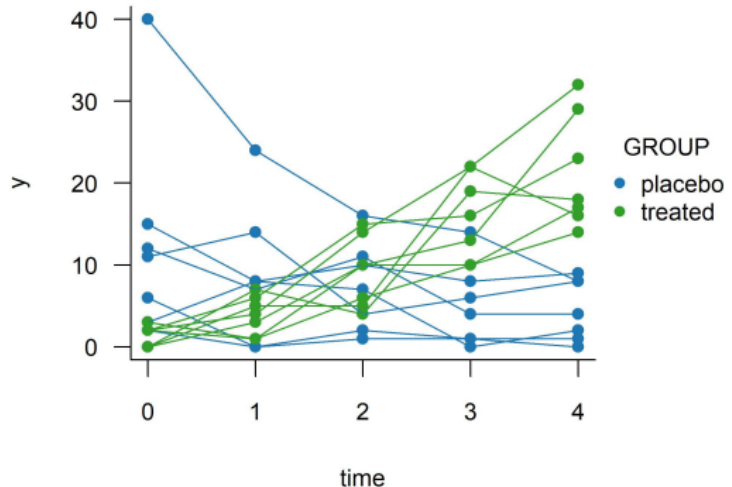
```
pmf(data.long$y)
```



# make.spaghetti()

## 3) Visualize longitudinal trajectories

```
make.spaghetti(x = time, y = y, id = id,  
                group = group, data = data.long)
```



# `make.spaghetti()`



```
make.spaghetti(x, y, id, group, data)
```

... even easier than making spaghetti!



# ptmixed(): estimate the Poisson-Tweedie GLMM!



## 4) Fit the Poisson-Tweedie GLMM

```
fit <- ptmixed(fixef.formula = y ~ group*time,  
                id = id, data = data.long)
```

The package also allows to estimate a few simpler models:

Function	Model
nbglm	Negative binomial GLM
nbmixed	Negative binomial GLMM
ptglm	Poisson-Tweedie GLM
ptmixed	Poisson-Tweedie GLMM



## summary()

- 5) View estimated  $\hat{\beta}$ ,  $\hat{D}$ ,  $\hat{\alpha}$ ,  $\hat{\sigma}_0^2$ , standard errors and univariate Wald tests

```
summary(fit)
```

```
## Loglikelihood: -140.539
## Parameter estimates:
##                               Estimate Std. error      z p.value
## (Intercept)           2.0012    0.2888  6.9286  0.0000
## grouptreated        -1.4546    0.4677 -3.1102  0.0019
## time                 -0.1360    0.0765 -1.7784  0.0753
## grouptreated:time    0.5115    0.1465  3.4908  0.0005
##
## Dispersion = 1.64
## Power = -0.14
## Variance = 0.42
```



## Predicted random effects

- 6) Compute the best linear unbiased predictor of the random effects

```
ranef(fit)
```

```
##      1      2      3      4      5      6
## -0.7467 0.3204 1.1391 -1.1709 0.4988 0.4097
##      7      8      9     10     11     12
## -0.2220 0.3224 -0.3528 -0.0221 0.1815 -0.2705
##     13     14
## -0.1814 0.6586
```



## Multivariate Wald test

7) Test more complex hypotheses, e.g.  $H_0 : \beta_1 = \beta_3 = 0$ <sup>1</sup>:

```
L = matrix(0, nrow = 2, ncol = 4)
L[1, 2] = L[2, 4] = 1
ptmixed::wald.test(fit, L = L, k = c(0, 0))
```

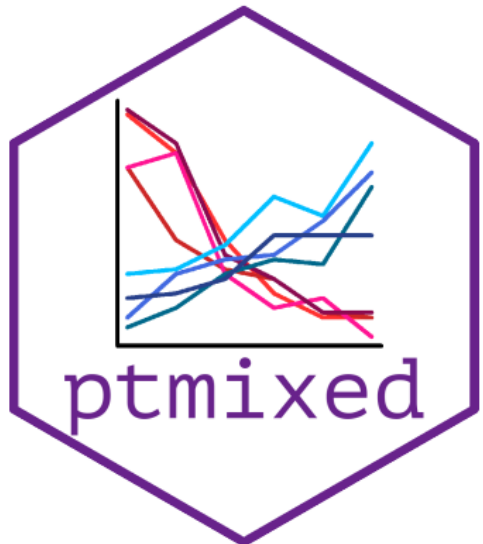
```
##          chi2 df          P
## 1 14.1051  2 0.0008651998
```

- ▶ Likelihood ratio test also possible, see sec. 2.4 of Signorelli et al. (2020)

---

<sup>1</sup>NB: the hypothesis is coded in the form  $L\beta = k$

# Useful resources



More info about ptmixed here:

- 1) arXiv:[2004.11193](https://arxiv.org/abs/2004.11193) preprint  
of Signorelli et al. (2020, in  
press)
- 2) CRAN [package page](#)
- 3) [Vignette](#) of the R package:

```
browseVignettes("ptmixed")
```



## References

- El-Shaarawi, A. H., Zhu, R., & Joe, H. (2011). Modelling species abundance using the poisson–Tweedie family. *Environmetrics*, 22(2), 152–164.
- Esnaola, M., Puig, P., Gonzalez, D., Castelo, R., & Gonzalez, J. R. (2013). A flexible count data model to fit the wide diversity of expression profiles arising from extensively replicated RNA-seq experiments. *BMC Bioinformatics*, 14(1), 254.
- Signorelli, M., Spitali, P., & Tsonaka, R. (2020). Poisson-Tweedie mixed-effects model: a flexible approach for the analysis of longitudinal RNA-seq data. *To Appear in Statistical Modelling*.  
<https://arxiv.org/abs/2004.11193>