

| Variable  | Example   | Type of<br>Regression                   | R function / R function for mixed models                |
|---|---|---|---|
| Continuous  | Quality of Life, linear scales  | linear                                  | lm()  |
|   |   |   | <pre>lmer(), glmmTMB()</pre>                            |
| Binary  | Success yes/no  | binary logistic                         | glm(family=binomial)                                    |
|   |   |   | <pre>glmer(*), glmmTMB(*)</pre>                         |
| Trials (or proportion of counts)                                      | 20 successes out of 30 trials   | logistic <sup>1</sup>                   | <pre>glm(cbind(trial, success), family=binomial)</pre>  |
|   |   |   | <pre>glmer(*), glmmTMB(*)</pre>                         |
| Count data  | Number of usage, counts of events   | Poisson                                 | glm(family=poisson)                                     |
|   |   |   | <pre>glmer(*), glmmTMB(*)</pre>                         |
| Count data, with excess zeros or overdispersion                       | Number of usage, counts of events (with higher variance than mean of response)  | negative<br>binomial                    | glm.nb()  |
|   |   |   | <pre>glmer.nb(), glmmTMB(family=nbinom)</pre>           |
| Count data with very many zeros (inflation)                           | see count data, but response is<br>modelled as mixture of Bernoulli &<br>Poisson distribution (two sources<br>of zeros) | zero-inflated                           | zeroinfl()  |
|   |   |   | glmmTMB(ziformula, family=poisson)                      |
| Count data, with very<br>many zeros (inflation)<br>and overdispersion | Number of usage, counts of events (with higher variance than mean of response)  | zero-inflated<br>negative<br>binomial   | zeroinfl(dist="negbin")                                 |
|   |   |   | glmmTMB(ziformula, family=nbinom)                       |
| ·   | see count data, but only for  |   | hurdle()  |
| Count data, zero-<br>truncated  | positive counts (hurdle component models zero-counts)   | hurdle (Poisson)                        | glmmTMB(family=truncated_poisson)                       |
| Count data, zero-<br>truncated and<br>overdispersion                  | see "Count data, zero-truncated",<br>but with higher variance than mean<br>of response                                  | hurdle (neg.<br>binomial)               | vglm(family=posnegbinomial)                             |
|   |   |   | glmmTMB(family=truncated_nbinom)                        |
| Proportion / Ratio  | Percentages, proportions of   |   | betareg()   |
| (without zero and one)  | continuous data   | Beta <sup>1</sup>                       | glmmTMB(family=beta)                                    |
| Proportion / Ratio<br>(including zero and one)                        | Percentages, proportions of   | Beta-Binomial,<br>zero-inflated<br>Beta | BBreg(), betabin(), vglm(family=betabinomial)           |
|   |   |   | glmmTMB(ziformula,<br>family=beta_family/ betabinomial) |
| Ordinal   | Likert scale, worse/ok/better   | ordinal,<br>proportional<br>odds        | polr(), clm()   |
|   |   |   | <pre>clmm(), mixor(), MCMCglmm()</pre>                  |
| Cumulative, multinomial   | 9 '   | cumulative link,<br>multinomial         | <pre>multinom(), clm(),bracl(), brmultinom()</pre>      |
|   |   |   | clmm(), mixor(), MCMCglmm()                             |
| Continuous, right-<br>skewed  | Financial data, reaction times  | Gamma                                   | glm(family=Gamma)                                       |
|   |   |   | <pre>glmer(*), glmmTMB(*)</pre>                         |
| Continuous, (right)<br>skewed, probably with                          | Financial data, probably exponential dispersion of variance   | Tweedie                                 | <pre>glm(family=tweedie), cpglm()</pre>                 |
| spike at zero   |   |   | <pre>cpglmm(), glmmTMB(family=tweedie)</pre>            |
| Continuous, but truncated or outliers                                 |   | truncated                               | <pre>censReg(), tobit(), vglm(family=tobit)</pre>       |
| Proportion / Ratio with > 2 categories                                | Biomass partitioning in plants (ratio of leaf, stem and root mass)  | Dirichlet                               | DirichReg()   |

<sup>\*</sup> Indicates same family-option for mixed models as for their non-multilevel counterparts.

<sup>&</sup>lt;sup>1</sup> Note that ratios or proportions from count data, like **cbind(trials, success)**, are modelled as logistic regression with **glm(cbind(trials, success), family=binomial())**, while ratios from continuous data where the response ranges from 0 to 1 are modelled using beta-regression.

