Post-model-fitting procedures with **glmmTMB** models: diagnostics, inference, and model output

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The purpose of this vignette is to describe (and test) the functions in various downstream packages that are available for summarizing and otherwise interpreting glmmTMB fits. Some of the packages/functions discussed below may not be suitable for inference on parameters of the zero-inflation or dispersion models, but will be restricted to the conditional-mean model.

```
library(glmmTMB)
library(car)
library(emmeans)
library(effects)
library(multcomp)
library(MuMIn)
require(DHARMa, quietly = TRUE) ## may be missing ...
library(broom)
library(broom.mixed)
require(dotwhisker, quietly = TRUE)
library(ggplot2); theme_set(theme_bw())
library(texreg)
library(xtable)
if (huxtable_OK) library(huxtable)
## retrieve slow stuff
L <- gt_load("vignette_data/model_evaluation.rda")</pre>
```

A couple of example models:

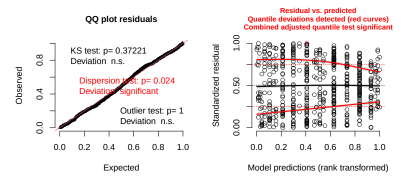
1 model checking and diagnostics

1.1 DHARMa

The DHARMa package provides diagnostics for hierarchical models. After running

```
owls_nb1_simres <- simulateResiduals(owls_nb1)
  you can plot the results:
plot(owls_nb1_simres)</pre>
```

DHARMa residual diagnostics



DHARMa provides lots of other methods based on the simulated residuals: see vignette("DHARMa", package="DHARMa")

1.1.1 issues

 DHARMa will only work for models using families for which a simulate method has been implemented (in TMB, and appropriately reflected in glmmTMB)

2 Inference

2.1 car::Anova

We can use car::Anova() to get traditional ANOVA-style tables from glmmTMB fits. A few limitations/reminders:

- these tables use Wald χ^2 statistics for comparisons (neither likelihood ratio tests nor F tests)
- they apply to the fixed effects of the conditional component of the model only (other components *might* work, but haven't been tested at all)
- as always, if you want to do type 3 tests, you should probably set sum-to-zero contrasts on factors and center numerical covariates (see contrasts argument above)

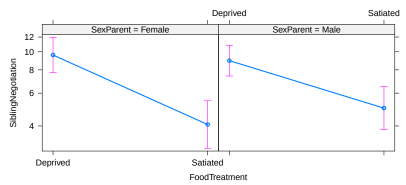
```
if (requireNamespace("car") && getRversion() >= "3.6.0") {
   Anova(owls_nb1) ## default type II
   Anova(owls_nb1, type="III")
}
```

Chisq	Df	Pr(>Chisq)
21.4	1	3.66e-06
46.1	1	1.1e-11
0.512	1	0.474
2.29	1	0.13

2.2 effects

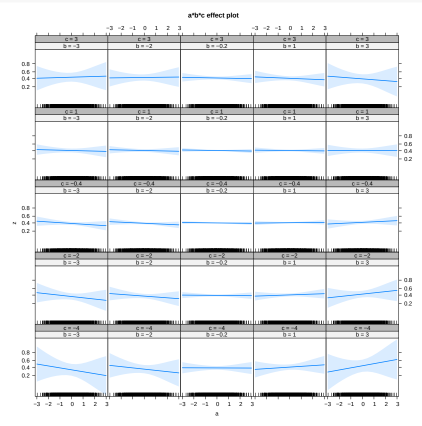
```
effects_ok <- (requireNamespace("effects") && getRversion() >= "3.6.0")
if (effects_ok) {
        (ae <- allEffects(owls_nb1))
        plot(ae)
}
## Warning in Effect.glmmTMB(predictors, mod, vcov. = vcov., ...):
overriding variance function for effects: computed variances may
be incorrect</pre>
```

FoodTreatment*SexParent effect plot



(the error can probably be ignored)

```
if (effects_ok) {
  plot(allEffects(simex_b1))
}
```



2.3 emmeans

```
emmeans(owls_nb1, poly ~ FoodTreatment | SexParent)
## $emmeans
## SexParent = Female:
  FoodTreatment emmean
                             SE df lower.CL upper.CL
                   2.30 0.1104 592
                                        2.09
                                                 2.52
   Deprived
##
   Satiated
                   1.44 0.1493 592
                                        1.15
                                                 1.74
##
## SexParent = Male:
##
  FoodTreatment emmean
                             SE
                               df lower.CL upper.CL
   Deprived
                   2.23 0.0964 592
                                        2.04
                                                 2.42
##
   Satiated
                   1.65 0.1357 592
                                        1.38
                                                 1.91
##
## Results are given on the log (not the response) scale.
## Confidence level used: 0.95
##
## $contrasts
## SexParent = Female:
   contrast estimate
                         SE df t.ratio p.value
##
   linear
              -0.859 0.149 592 -5.776 <.0001
##
## SexParent = Male:
  contrast estimate
                         SE df t.ratio p.value
##
   linear
            -0.586 0.129 592 -4.531 <.0001
##
## Results are given on the log (not the response) scale.
```

2.4 drop1

stats::drop1 is a built-in R function that refits the model with various terms dropped. In its default mode it respects marginality (i.e., it will only drop the top-level interactions, not the main effects):

```
system.time(owls_nb1_d1 <- drop1(owls_nb1,test="Chisq"))
## user system elapsed
## 1.054 0.002 1.056</pre>
```

In principle, using <code>scope = .~. - (1|Nest)</code> should work to execute a "type-3-like" series of tests, dropping the main effects one at a time while leaving the interaction in (we have to use - (1|Nest) to exclude the random effects because <code>drop1</code> can't handle them). However, due to the way that R handles formulas, dropping main effects from an interaction of *factors* has no effect on the overall model. (It would work if we were testing the interaction of continuous variables.)

2.4.1 issues

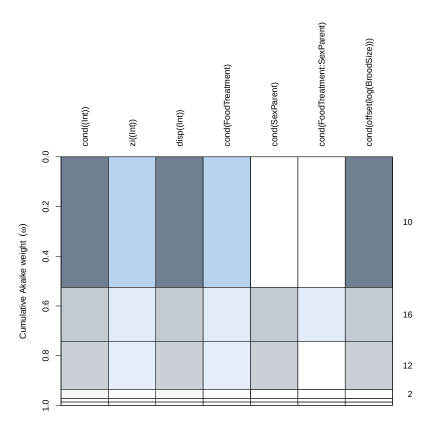
The mixed package implements a true "type-3-like" parameter-dropping mechanism for [g]lmer models. Something like that could in principle be applied here.

2.5 Model selection and averaging with MuMIn

We can run MuMIn::dredge(owls_nb1) on the model to fit all possible submodels. Since this takes a little while (45 seconds or so), we've instead loaded some previously computed results:

```
print(owls_nb1_dredge)
## Global model call: glmmTMB(formula = SiblingNegotiation ~ FoodTreatment * SexPa
       (1 | Nest) + offset(log(BroodSize)), data = Owls, family = nbinom1,
##
       ziformula = ~1, contrasts = list(FoodTreatment = "contr.sum",
##
           SexParent = "contr.sum"), na.action = na.fail, dispformula = ~1)
##
## Model selection table
##
      cnd((Int)) zi((Int)) dsp((Int)) cnd(FdT) cnd(SxP) cnd(FdT:SxP)
## 10
                    -2.094
                                    +
          0.4284
## 16
          0.4275
                    -2.055
## 12
          0.4257
                    -2.100
## 2
                    -1.990
         1.8290
## 8
         1.8280
                    -1.955
                    -1.996
## 4
         1.8260
## 9
         0.6295
                    -1.373
                    -1.232
## 1
          2.0980
## 11
                    -1.381
          0.6220
                                                       +
                    -1.236
## 3
          2.0920
                                    +
##
      cnd(off(log(BrS))) df
                               logLik
                                        AICc delta weight
                          5 -1685.978 3382.1 0.00
## 10
                                                    0.525
## 16
                       + 7 -1684.819 3383.8
                                              1.77
                                                    0.217
## 12
                       + 6 -1685.957 3384.1
                                              2.00 0.193
## 2
                          5 -1688.628 3387.4 5.30 0.037
## 8
                          7 -1687.556 3389.3
                                              7.24 0.014
                          6 -1688.610 3389.4 7.30 0.014
## 4
## 9
                       + 4 -1708.573 3425.2 43.15 0.000
                          4 -1708.672 3425.4 43.35 0.000
## 1
## 11
                       + 5 -1708.420 3426.9 44.88 0.000
## 3
                          5 -1708.509 3427.1 45.06 0.000
## Models ranked by AICc(x)
## Random terms (all models):
## 'cond(1 | Nest)'
```

```
op <- par(mar=c(2,5,14,3))
plot(owls_nb1_dredge)</pre>
```



par(op) ## restore graphics parameters

${\bf Model\ averaging:}$

```
model.avg(owls_nb1_dredge)
##
## Call:
## model.avg(object = owls_nb1_dredge)
##
## Component models:
## '14'
            1234
                     124'
                              11
                                        123
                                                 12'
                                                          ,4,
                                                                   '(Null)'
## '24'
            ,2,
##
## Coefficients:
```

2.5.1 issues

• may not work for Beta models because the family component ("beta") is not identical to the name of the family function (beta_family())? (Kamil Bartoń, pers. comm.)

2.6 multcomp for multiple comparisons and post hoc tests

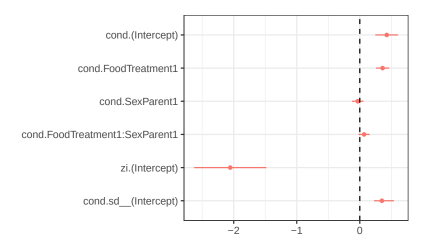
```
g1 <- glht(cbpp_b1, linfct = mcp(period = "Tukey"))
summary(g1)
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Multiple Comparisons of Means: Tukey Contrasts
##
##
## Fit: glmmTMB(formula = incidence/size ~ period + (1 | herd), data = cbpp,
      family = binomial, weights = size, ziformula = ~0, dispformula = ~1)
##
##
## Linear Hypotheses:
             Estimate Std. Error z value Pr(>|z|)
             -0.9923
                          0.3066 -3.236 0.00638 **
## 2 - 1 == 0
## 3 - 1 == 0 -1.1287
                          0.3266 -3.455 0.00283 **
## 4 - 1 == 0 -1.5803
                          0.4274 -3.697
                                          0.00111 **
## 3 - 2 == 0 -0.1363
                          0.3807
                                  -0.358
                                          0.98368
## 4 - 2 == 0 -0.5880
                          0.4703 -1.250 0.58569
## 4 - 3 == 0 -0.4516
                          0.4843 -0.933 0.78117
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
```

3 Extracting coefficients, coefficient plots and tables

3.1 broom and friends

The broom and broom.mixed packages are designed to extract information from a broad range of models in a convenient (tidy) format; the dotwhisker package builds on this platform to draw elegant coefficient plots.



3.1.1 issues

(these are more general dwplot issues)

- use black rather than color(1) when there's only a single model, i.e. only add aes(colour=model) conditionally? draw points even if std err / confint are NA (draw geom_point() as well as geom_pointrange()? need to apply all aesthetics, dodging, etc. to both ...)
- for glmmTMB models, allow labeling by component? or should this be done by manipulating the tidied frame first? (i.e.: tidy(.) \%>\% tidyr::unite(term,c(co

3.2 coefficient tables with xtable

The xtable package can output data frames as LATEX tables; this isn't quite as elegant as stargazer etc., but is not a bad start. I've sprinkled lots of hard line-breaks, spaces, and newlines in below: someone who was better at TEX could certainly do a better job. (xtable can also produce HTML output.)

```
ss <- summary(owls_nb1)
## print table; add space,
pxt <- function(x,title) {
   cat(sprintf("{\n\n\textbf{%s}\n\\ \\\\n",title))</pre>
```

```
print(xtable(x), floating=FALSE); cat("\n\n")
  cat("\\ \\\\\vspace{5pt}\\ \\\\n")
}
```

```
pxt(lme4::formatVC(ss$varcor$cond),"random effects variances")
pxt(coef(ss)$cond,"conditional fixed effects")
pxt(coef(ss)$zi,"conditional zero-inflation effects")
```

random effects variances

	Groups	Name	Std.Dev.
1	Nest	(Intercept)	0.35019

conditional fixed effects

	Estimate	Std. Error	z value	$\Pr(> z)$
(Intercept)	0.43	0.09	4.63	0.00
${\bf FoodTreatment 1}$	0.36	0.05	6.79	0.00
SexParent1	-0.03	0.05	-0.72	0.47
FoodTreatment1:SexParent1	0.07	0.05	1.51	0.13

conditional zero-inflation effects

	Estimate	Std. Error	z value	$\Pr(> z)$
(Intercept)	-2.06	0.29	-7.03	0.00

3.3 coefficient tables with texreg

For now, to avoid needing to import the texreg package, we are providing the required extract.glmmTMB in a separate R file that you can import with source(), as follows:

	Model 1
(Intercept)	0.43***
· - /	(0.09)
FoodTreatment1	0.36***
	(0.05)
SexParent1	-0.03
	(0.05)
FoodTreatment1:SexParent1	0.07
	(0.05)
$zi_{-}(Intercept)$	-2.06***
	(0.29)

^{***}p < 0.001; **p < 0.01; *p < 0.05

Table 1: Owls model

```
source(system.file("other_methods","extract.R",package="glmmTMB"))
texreg(owls_nb1,caption="Owls model", label="tab:owls")
```

See output in Table 1.

3.4 coefficient tables with huxtable

The huxtable package allows output in either LaTeX or HTML: this example is tuned for LaTeX.

```
statistics = "nobs" # don't include logLik and AIC
)
names(h0)[2:3] <- c("estimate", "std. err.")
## allow use of math notation in name
h1 <- set_cell_properties(h0,row=5,col=1,escape_contents=FALSE)
cat(to_latex(h1,tabular_only=TRUE))
}</pre>
```

intercept (mean)	0.427 ***	(0.092)
food treatment (starvation)	0.361 ***	(0.053)
parental sex (M)	-0.033	(0.047)
$\rm food \times sex$	0.068	(0.045)
nobs	599	

^{***} p < 0.001; ** p < 0.01; * p < 0.05.

3.4.1 issues

• huxtable needs quite a few additional LATEX packages: use report_latex_dependencies() to see what they are.

4 influence measures

Influence measures quantify the effects of particular observations, or groups of observations, on the results of a statistical model; leverage and Cook's distance are the two most common formats for influence measures. If a projection matrix (or "hat matrix") is available, influence measures can be computed efficiently; otherwise, the same quantities can be estimated by brute-force methods, refitting the model with each group or observation successively left out.

We've adapted the car::influence.merMod function to handle glmmTMB models; because it uses brute force, it can be slow, especially if evaluating the influence of individual observations. For now, it is included as a separate

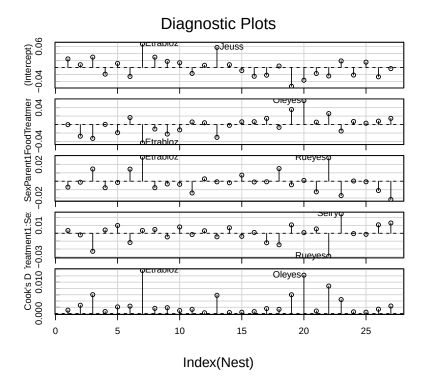
source file rather than exported as a method (see below), although it may be included in the package (or incorporated in the car package) in the future.

```
source(system.file("other_methods","influence_mixed.R", package="glmmTMB"))
```

```
owls_nb1_influence_time <- system.time(
  owls_nb1_influence <- influence_mixed(owls_nb1, groups="Nest")
)</pre>
```

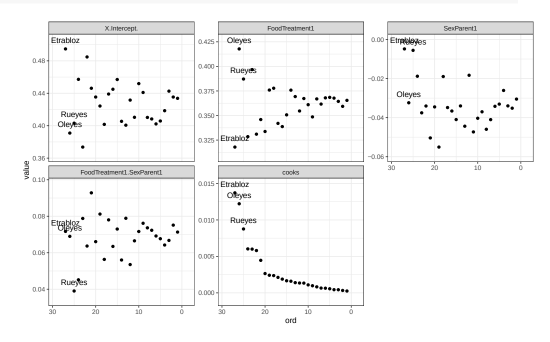
Re-fitting the model with each of the 27 nests excluded takes 16 seconds (on an old Macbook Pro). The car::infIndexPlot() function is one way of displaying the results:

```
car::infIndexPlot(owls_nb1_influence)
```



Or, you can transform the results and plot them however you like:

```
inf <- as.data.frame(owls_nb1_influence[["fixed.effects[-Nest]"]])</pre>
inf <- transform(inf,</pre>
                 nest=rownames(inf),
                  cooks=cooks.distance(owls_nb1_influence))
inf$ord <- rank(inf$cooks)</pre>
if (require(reshape2)) {
  inf_long <- melt(inf, id.vars=c("ord", "nest"))</pre>
  gg_infl <- (ggplot(inf_long,aes(ord,value))</pre>
    + geom_point()
    + facet_wrap(~variable, scale="free_y")
    ## n.b. may need expand_scale() in older ggplot versions ?
    + scale_x_reverse(expand=expansion(mult=0.15))
    + scale_y_continuous(expand=expansion(mult=0.15))
    + geom_text(data=subset(inf_long,ord>24),
                 aes(label=nest), vjust=-1.05)
  print(gg_infl)
## Loading required package:
                                reshape2
```



5 to do

- \bullet more plotting methods (\mathtt{sjplot})
- \bullet output with ${\tt memisc}$
- AUC etc. with ModelMetrics