Review of linear models

18 Oct 2023

```
## it's nice to include packages at the top
## (and NOT automatically install them)
## try not to carry over packages you don't use
library(faux)
library(brglm2) ## for lizards data
library(ggplot2); theme_set(theme_bw())
## diagnostics
library(performance)
library(DHARMa)
## downstream model evaluation
library(broom)
library(dotwhisker)
library(emmeans)
library(effects)
library(marginaleffects)
library(parameters)
## library(qqeffects)
```

Basics

- assume $\mathbf{y} \sim \text{Normal}(\mathbf{X}\beta, \sigma)^1$
- **X** is the *model matrix*, can be anything we want it to be
- the *Gauss-Markov theorem* (Wikipedia) makes weaker assumptions: $\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\epsilon}$; as long as $\boldsymbol{\epsilon}$ is mean-zero, homoscedastic with finite variance, and uncorrelated ... then the OLS solution

$$\hat{\boldsymbol{\beta}} = (\mathbf{X}^{\top} \mathbf{X})^{-1} \mathbf{X}^{\top} \mathbf{y}$$

is the BLUE (or MVUE).

¹Notation-abuse warning ...

• we'll embrace the assumptions (which are needed for inference!)

Computation

- matrix decompositions (QR with pivoting; see here)
- big problems: biglm, speedglm, RcppEigen::fastLm
 - optimized BLAS, kernel trick, etc.
 - memory vs speed vs robustness ...
 - -p vs. n vs. many-small-regressions vs. ...

Inference

- σ^2 (residual variance) is RSS/(n-p)
- The covariance matrix is $\Sigma = \sigma^2(\mathbf{X}^{\top}\mathbf{X})^{-1}$.
- Individual coefficients are *t*-distributed
- Linear combinations of coefficients (contrasts or predictions) are t-distributed with covariance matrix $\mathbf{C}^{\top}\Sigma^{-1}\mathbf{C}$
- \bullet Joint hypotheses on coefficients are F-distributed
- Wald and likelihood ratio test comparisons are equivalent (but need to be careful about marginality)

Model matrices

- model definition converted to **X** before we start
- input variables vs predictor variables (Schielzeth (2010), Gelman and Hill (2006), CV)
 - transformations
 - encoding of categorical variables: contrasts
 - interactions
 - basis expansions (e.g. polynomials)

Wilkinson-Rogers formulas

- Wilkinson and Rogers (1973), updated by Chambers and Hastie (1991, ch. 2)
- operators: +, *, :, /, -, ^
- I()

Contrasts

treatment contrasts

- intercept = baseline, subsequent values are differences
- $\begin{array}{l} \bullet \ \left\{ \beta_0 = \mu_0, \beta_i = \mu_i \mu_0 \text{ for } i > 0 \right\} \\ \bullet \ \text{equivalently: } \left\{ \mu_0 = \beta_0, \mu_i = \beta_0 + \beta_i \text{ for } i > 0 \right\} \end{array}$

$$\mathbf{C}\beta = \begin{pmatrix} 1 & 0 & 0 & \dots \\ 1 & 1 & 0 & \dots \\ 1 & 0 & 1 & \dots \\ \vdots & \vdots & \vdots & \ddots \end{pmatrix} \begin{pmatrix} \beta_0 \\ \beta_1 \\ \beta_2 \\ \dots \end{pmatrix} = \begin{pmatrix} \mu_1 \\ \mu_2 \\ \mu_3 \\ \dots \end{pmatrix} \quad .$$

Maybe easier to start from the **inverse** contrast matrix: $\beta = \mathbf{C}^{-1}\mu$.

```
C <- cbind(1, contr.treatment(3)) ## R omits the intercept by default
```

1 2 3

1 0 0

2 - 1 1 0

3 -1 0 1

We have to specify the baseline level (contr.treatment uses first level of a factor; contr.SAS() uses the last level).

It's nice when contrasts are *orthogonal*, i.e. all rows are independent $\to \mathbf{C}^\top \mathbf{C}$ is diagonal.

Sum-to-zero contrasts

- intercept is the (unweighted!) average rather than baseline value $(\sum \mu_i/n)$
- other parameters are differences between mean of level i and intercept $(\mu_i \sum_i \mu_j / n)$
- last level is dropped

```
mfun <- function(C) MASS::fractions(solve(C))</pre>
(C <- cbind(1,contr.sum(3)))
```

Helmert contrasts

[3,] -1/3 2/3 -1/3

- Weird but orthogonal
- intercept, diff of first two levels, diff of level 3 from 1 & 2, ...

others

[3,] -1/6 -1/6 1/3

- MASS::contr.sdif() (successive-differences)
- contr.poly() (orthogonal polynomial contrasts)
- custom (e.g., "none" vs "symbiont effect" vs "crabs vs shrimp" vs "two-symbiont effect") (McKeon et al. (2012); data here)

```
cc_{inv} \leftarrow matrix(c(1/4,1/4,1/4,1/4,
                 1,-1/3,-1/3,-1/3,
                 0,1,-1,0,
                 0,1/2,1/2,-1),
               byrow=TRUE,
               nrow=4,
               dimnames=list(c("intercept", "avg_symb", "C.vs.S", "twosymb"),
                             c("none", "C", "S", "CS")))
  ## inverse contrast matrix
  MASS::fractions(cc_inv)
         none C S
                        CS
intercept 1/4 1/4 1/4 1/4
            1 -1/3 -1/3 -1/3
avg_symb
C.vs.S
             0 1 -1
             0 1/2 1/2
twosymb
                          -1
  ## contrast matrix
  mfun(cc_inv)
     intercept avg_symb C.vs.S twosymb
               3/4
                                0
none
        1
                         0
С
              -1/4
                        1/2
                               1/3
S
        1
              -1/4
                       -1/2 1/3
CS
        1
              -1/4
                          0 -2/3
```

practical issues

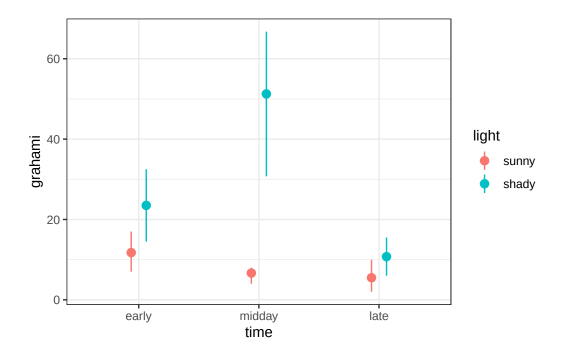
- too many ways to set contrasts (options(), contrasts(f) <-, lm(..., contrasts =
 list(...))</pre>
- terrible naming conventions: you can get used to it or use the faux package
- OK to fit models and later use emmeans to recover desired contrasts (switching linear bases)

```
mtcars$fcyl <- factor(mtcars$cyl)
lm0 <- lm(mpg ~ fcyl, mtcars)
cn <- function(x) names(coef(x))
cn(lm0)</pre>
```

Interactions

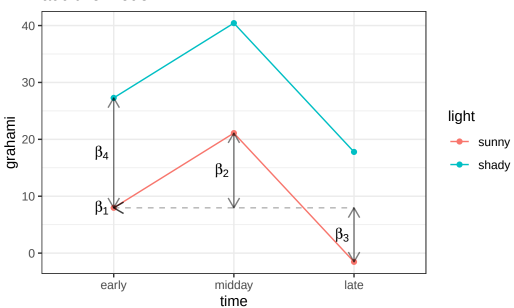
- differences in differences
- parameter values of main effects (and p values etc.) depend on contrasts/centering!
- overall model fit (R^2 , predictions, etc.) is invariant

Lizard data (Schoener (1970), from the brglm2 package):



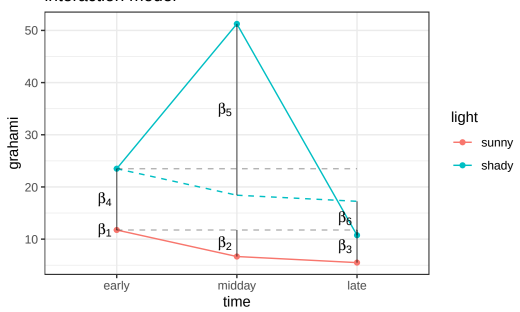
```
data("lizards", package = "brglm2")
lmTL1 <- lm(grahami~time+light,data=lizards)</pre>
pp <- with(lizards,expand.grid(time=levels(time),light=levels(light)))</pre>
pp$grahami <- predict(lmTL1,newdata=pp)</pre>
cc <- as.list(plyr::rename(coef(lmTL1),c(`(Intercept)`="int")))</pre>
labelpos <- with(cc,</pre>
  list(x=c(1,2,3,1), xend=c(1,2,3,1),
      y=c(int,int,int,int),
      yend=c(int,int+timemidday,int+timelate,int+lightshady)))
xpos < -0.1
ggplot(pp,aes(x=time,y=grahami,colour=light))+geom_point()+
  geom_line(aes(group=light))+
  annotate("segment", x=labelpos$x, xend=labelpos$xend, y=labelpos$y,
           yend=labelpos$yend,alpha=0.5,
           arrow=arrow(length = unit(0.3, "cm"), ends="both"))+
  annotate("text", x=with(labelpos, (x+xend)/2)+xpos, y=with(labelpos, (y+yend)/2),
label=paste0("beta[",1:4,"]"),parse=TRUE)+
  annotate("segment", x=labelpos$x[1], xend=labelpos$x[3], y=labelpos$y[1],
           yend=labelpos$y[1],alpha=0.3,lty=2) +
    labs(title = "additive model")
```

additive model



```
lmTL2 <- lm(grahami~time*light,data=lizards)</pre>
gg_color_hue <- function(n) {</pre>
  hues = seq(15, 375, length=n+1)
  hcl(h=hues, l=65, c=100)[1:n]
}
pp2 <- pp
pp2$grahami <- predict(lmTL2,newdata=pp)</pre>
cc <- as.list(plyr::rename(coef(lmTL2),c(`(Intercept)`="int",</pre>
        `timemidday:lightshady`="midshady",`timelate:lightshady`="lateshady")))
labelpos <- with(cc,</pre>
  list(x=c(1,2,3,1,2,3), xend=c(1,2,3,1,2,3),
      y=c(int,int,int,int,int+lightshady+timemidday,int+lightshady+timelate),
      yend=c(int,int+timemidday,int+timelate,int+lightshady,
             int+timemidday+lightshady+midshady,int+timelate+lightshady+lateshady)))
xpos < -0.1
ggplot(pp2,aes(x=time,y=grahami,colour=light))+geom_point()+
  geom_line(aes(group=light))+
  annotate("segment", x=1:2, xend=2:3,
           y=with(cc,c(int+lightshady,int+timemidday+lightshady)),
           yend=with(cc,c(int+timemidday+lightshady,int+timelate+lightshady)),
           colour=gg_color_hue(2)[2],lty=2)+
  annotate("segment", x=labelpos$x, xend=labelpos$xend, y=labelpos$y,
```

interaction model



Marginality

- Venables (1998)
- 'type (X) sums of squares'
- scaling and centering (Schielzeth 2010) alleviates many problems; sum-to-zero contrasts (weighted or unweighted?)

Model interpretation, visualization, testing

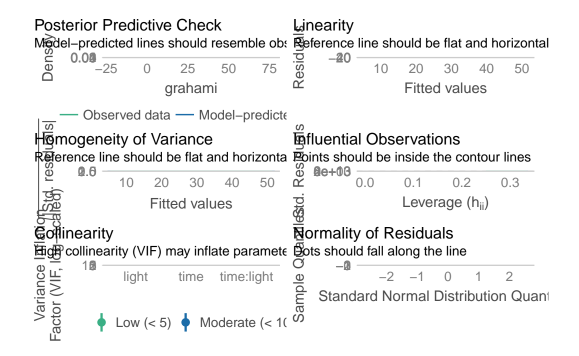
Diagnostics

- linearity > heteroscedasticity, outliers > normality
- upstream problems can induce downstream problems first
- universal plots are universal, but less interpretable than problem-specific exploration (try to identify problematic predictors/groups/etc.)

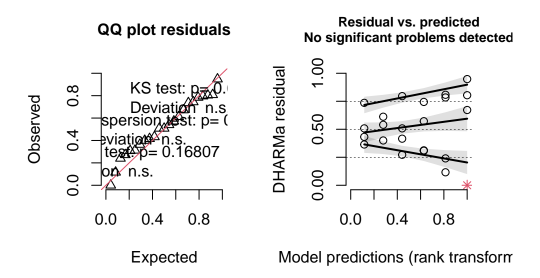
Graphical diagnostics

- base R: stats::plot.lm()
- performance::check_model()
- DHARMa (simulateResiduals(., plot = TRUE))
 - (plotResiduals(simout, form = pred_var))
- broom::augment() + plot-your-own (ggplot2)

performance::check_model(lmTL2)

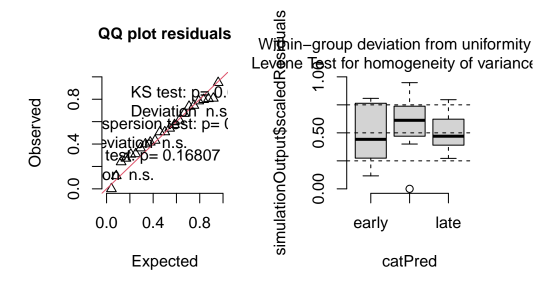


DHARMa residual



plot(ss, form = lizards\$time)

DHARMa residual



Solutions to problems

- nonlinearity: transformation, add covariates (??), add interactions, add polynomial
- outliers: drop values (report both!), use robust regression
- heteroscedasticity: transformation, model dispersion explicitly, GLMs
- non-Normality: transformation, GLMs

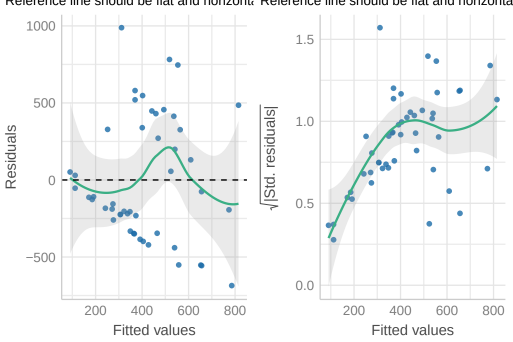
Transformation

- May do too much at once (GLMs and GAMs allow more flexibility)
- Log-transformation is often interpretable and solves problems
- ullet Transforming boundary values (e.g. $\log(0)$) is problematic
- **Box-Cox transformations**: $y \to \frac{y^{\lambda}-1}{\lambda}$ (include Jacobian term $GM^{\lambda-1}$ in denominator to keep log-likelihood comparable)
 - flexible
 - in practice people often use 'round numbers': $\lambda = 0$ (log), 1/2 (square root), etc.
 - MASS::boxcox()
 - hard to interpret!

Example

```
library(faraway)
m1 <- lm(perm ~ area, rock)
performance::check_model(m1, check = c("linearity", "homogeneity", "outliers", "qq"))
```

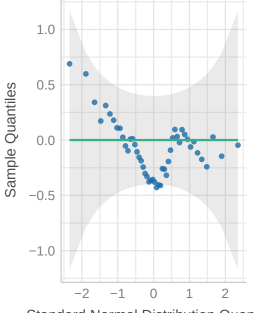
Linearity Homogeneity of Variance Reference line should be flat and horizontal Reference line should be flat and horizontal



Influential Observations Points should be inside the contour lines

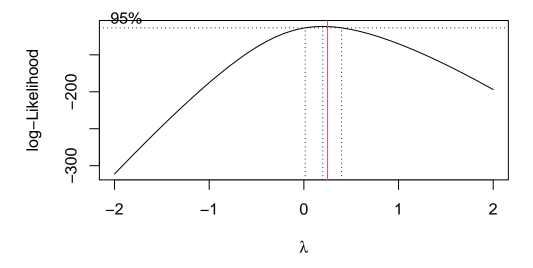
Sesional de inside the contour lines both and the serional seriona

Normality of Residuals Dots should fall along the line



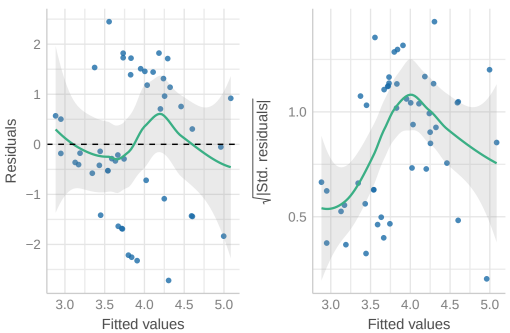
Standard Normal Distribution Quanti

```
m <- MASS::boxcox(m1, interp = FALSE)
abline(v=1/4, col = 2)</pre>
```



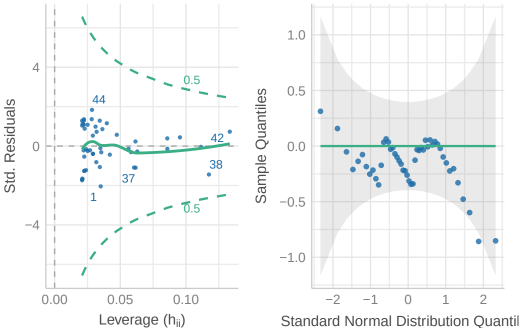
```
m2 <- update(m1, perm^(1/4) ~ .)
performance::check_model(m2, check = c("linearity", "homogeneity", "outliers", "qq"))</pre>
```

Linearity Homogeneity of Variance
Reference line should be flat and horizon Reference line should be flat and horizontal



Influential Observations Points should be inside the contour lines Dots should fall along the line

Normality of Residuals 1.0



When we transform we have to add the logs of the <code>Jacobian</code> of the transform to the log-likelihood, or subtract $\sum \log(J_i)$ from the negative log-likelihood. e.g. if we log-transform, $\sum \log(J(y_i)) = \sum \log\left(\partial(\log(y_i))/\partial y\right) = \sum \log(1/y_i) = -\sum \log y_i$. In this case $\sum \log(J(y_i)) = \sum \log((1/4)P^(-3/4)) = -N\log 4 - 3/4 \sum \log(P)$:

Check:

[1] 23.52213

```
logJ <- -(nrow(rock)*log(4) + 3/4*sum(log(rock$perm)))
lm2 <- logLik(m2) + logJ
lm1 <- logLik(m1)
lm2-lm1

'log Lik.' 23.3844 (df=3)

## difference from Box-Cox sequence
max(m$y)-m$y[m$x==1]</pre>
```

Reminder about parameter scaling

- Schielzeth (2010), Gelman (2008)
- centering ensures no problems with marginality (analogous to sum-to-zero contrasts)
- scaling predictors gives coefficients equal units (analogous to Cohen's d)
- scaling response makes coefficients *standardized partial regression coefficients* (equal to partial correlation coefficients if predictors are independent)
- scaling by 2 SD makes coefficients equivalent to those from balanced binary predictors
- scaling inputs or scaling predictors?

What about correlated predictors?

- Can compute variance inflation factors (VIFs)
- Dropping correlated factors is dubious: Graham (2003), Dormann et al. (2012), Morrissey and Ruxton (2018), Vanhove (2021)
- perfect collinearity gets handled automatically by R's pivoting, but may want to change contrasts/model setup

```
summary(lmTL2)
```

Call:

lm(formula = grahami ~ time * light, data = lizards)

Residuals:

Min 1Q Median 3Q Max -30.250 -4.125 1.250 6.875 17.750

Coefficients:

	${\tt Estimate}$	${\tt Std.} \ {\tt Error}$	t value	Pr(> t)	
(Intercept)	11.750	5.375	2.186	0.04311	*
timemidday	-5.083	8.211	-0.619	0.54408	
timelate	-6.250	7.602	-0.822	0.42238	
lightshady	11.750	7.602	1.546	0.14061	
timemidday:lightshady	32.833	11.190	2.934	0.00927	**
timelate:lightshady	-6.500	10.751	-0.605	0.55343	

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

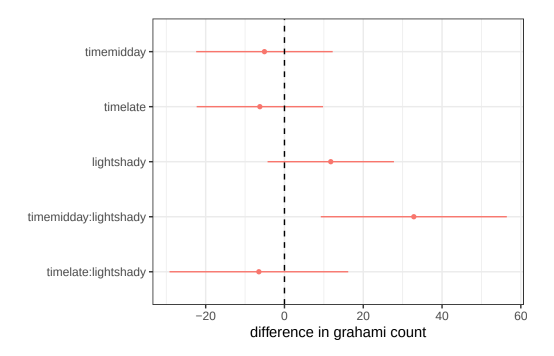
Residual standard error: 10.75 on 17 degrees of freedom Multiple R-squared: 0.7504, Adjusted R-squared: 0.677 F-statistic: 10.22 on 5 and 17 DF, p-value: 0.0001179

broom::tidy(lmTL2)

A tibble: 6 x 5

	term	${\tt estimate}$	${\tt std.error}$	${\tt statistic}$	p.value
	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
1	(Intercept)	11.8	5.38	2.19	0.0431
2	timemidday	-5.08	8.21	-0.619	0.544
3	timelate	-6.25	7.60	-0.822	0.422
4	lightshady	11.8	7.60	1.55	0.141
5	timemidday:lightshady	32.8	11.2	2.93	0.00927
6	timelate:lightshady	-6.50	10.8	-0.605	0.553

```
## automatically drop intercept; optional by_2sd argument
dotwhisker::dwplot(lmTL2) + geom_vline(xintercept = 0, lty = 2) +
    labs(x="difference in grahami count")
```



Interpretation and testing

- Look at coefficient tables: summary() or coef(summary())
- model comparison: drop1(), anova(), car::Anova()
- coefficient *plots*: broom + ggplot2, dotwhisker

Downstream methods

- plot predictions with data
- partial residuals plots (e.g. remef package)
- prediction, effects plots
- uncertainty of predictions
- emmeans, marginaleffects, effects, sjPlot ...

References

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