Partial residuals with the package LMMstar

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This vignette details how partial residuals can be used to illustrate model fit in a linear regression and a linear mixed model when using the package LMMstar. We thus start by loading the necessary packages:

```
library(LMMstar)
library(ggplot2)
```

1 Univariate linear regression

To illustrate the use of partial residuals we will use the state.x77 dataset:

```
lifeExp illiteracy income murder
                                                 edu
           69.05
                         2.1 3.624
                                       15.1
                                              (0,50]
Alabama
Alaska
           69.31
                         1.5 6.315
                                       11.3 (60,100]
                         1.8 4.530
           70.55
                                       7.8
                                             (50,60]
Arizona
Arkansas
           70.66
                         1.9
                              3.378
                                       10.1
                                              (0,50]
```

which contains information about life expectancy (lifeExp), income (income), illeteracy (illiteracy), murder rate (murder), and the percentage of high-school graduates (as categorical variable) in various states in the USA. For later use we display a few descriptive for each covariate value:

```
outcome observed missing
                                   mean
                                           min
                                                  max
1
       lifeExp
                     50
                              0 70.8786 67.960 73.600 1.34239
2
    illiteracy
                     50
                                1.1700 0.500 2.800 0.60953
3
        income
                     50
                                 4.4358 3.098 6.315 0.61447
4
                              0 7.3780 1.400 15.100 3.69154
        murder
                     50
  edu: (50,60]
                     50
                              0 0.5600 0.000 1.000 0.50143
6 edu: (60,100]
                     50
                              0 0.1600 0.000 1.000 0.37033
```

and check there are no missing values. Here for the categorical covariates the mean indicates the relative frequency of occurrence (56% and 16%) and other columns like sd should be ignored.

1.1 No interaction

Suppose we are interested in relating life expectancy (Y) to income (X). We cannot directly illustrate this relationship, as it could be confounded by other variables such as illiteracy (Z_1) , murder rate (Z_2) . and education (Z_3) . We will therefore use a linear model to control for those variables $(\mathbf{Z} = (Z_1, Z_2, Z_3))$ where, for simplicity, we assume a linear effect for all variables:

$$Y = \alpha + \beta X + \gamma_1 Z_1 + \gamma_2 Z_2 + \gamma_2 Z_3 + \varepsilon$$

```
e.lm <- lmm(lifeExp \sim income + illiteracy + murder + edu, data = df1) model.tables(e.lm)
```

```
estimate
                           se
                                  df
                                        lower
                                                 upper
                                                          p.value
(Intercept) 71.57828 1.138488 44.009 69.28382 73.87274 0.0000e+00
income
            0.19270 0.252649 44.009 -0.31648 0.70188 4.4971e-01
             0.17590 0.320967 44.009 -0.47096 0.82276 5.8644e-01
illiteracy
            -0.27822 0.047855 44.009 -0.37467 -0.18178 6.3280e-07
murder
             0.30141 0.414015 44.009 -0.53298 1.13580 4.7046e-01
edu(50,60]
edu(60,100]
            0.77306 0.513118 44.009 -0.26105 1.80718 1.3906e-01
```

Note that the estimates are nearly identical to the ones of the lm function:

```
coef(e.lm) - coef(lm(lifeExp \sim income + illiteracy + murder + edu, data = df1))
```

```
(Intercept) income illiteracy murder edu(50,60] edu(60,100] -2.7001e-13 2.0345e-13 -2.9035e-13 6.3283e-15 -4.2277e-13 -5.3091e-13
```

A graphical display can now be obtained by modifying the original outcome, life expectancy, had every state had the same illiteracy and murder rate:

```
df1$pres <- residuals(e.lm, type = "partial", variable = c("(Intercept)", "income"))
head(df1)</pre>
```

```
lifeExp illiteracy income murder
                                                  edu
                                                        pres
Alabama
             69.05
                          2.1
                               3.624
                                        15.1
                                               (0,50] 72.882
             69.31
                                        11.3 (60,100] 71.417
Alaska
                          1.5
                               6.315
                                         7.8
                                              (50,60] 72.102
Arizona
             70.55
                          1.8 4.530
Arkansas
             70.66
                          1.9
                               3.378
                                        10.1
                                               (0,50] 73.136
California
             71.71
                          1.1 5.114
                                        10.3 (60,100] 73.609
Colorado
             72.06
                          0.7
                               4.884
                                        6.8 (60,100] 73.056
```

By default, the partial residuals are computed substracting the effect of the covariates, i.e. had each state got no illiteracy, no murder, and the lowest education level:

```
c(69.05 - 0.17590 * 2.1 - (-0.27822) * 15.1,
69.31 - 0.17590 * 1.5 - (-0.27822) * 11.3 - 0.77306)
```

[1] 72.882 71.417

The element "(Intercept)" was specified in the variable argument to avoid to substract the mean value from the outcome and keep a plausible range of values for the outcome.

One may wish to compute the life expectancy had every state got a specific illiteracy, murder rate, and education. Say the most common in the sample: 1.17, 7.378, and (50,60] based on the descriptive statistics. This can be obtained by specifying these values in the argument at:

```
lifeExp illiteracy income murder
                                                 edu
                                                       pres pres2
            69.05
                                              (0,50] 72.882 71.336
                          2.1 3.624
                                       15.1
Alabama
             69.31
                          1.5 6.315
                                       11.3 (60,100] 71.417 69.871
Alaska
            70.55
                          1.8 4.530
                                       7.8
                                            (50,60] 72.102 70.557
Arizona
Arkansas
            70.66
                          1.9 3.378
                                       10.1
                                              (0,50] 73.136 71.590
                                       10.3 (60,100] 73.609 72.064
California
            71.71
                          1.1 5.114
                                       6.8 (60,100] 73.056 71.510
Colorado
            72.06
                          0.7 4.884
```

or doing the calcuation by hand:

```
c(69.05 - 0.17590 * (2.1-1.170) - (-0.27822) * (15.1-7.378) + 0.30141,
69.31 - 0.17590 * (1.5-1.170) - (-0.27822) * (11.3-7.378) + 0.30141 - 0.77306)
```

[1] 71.336 69.871

Note that changing in counterfactual only shifts the partial residuals by a constant, here:

```
unique(df1$pres2 - df1$pres)
```

[1] -1.5455

so does not affect the relation between the counterfactual outcome (here lifeExp) and the exposure of interest (here income). One can then get a graphical display either manually using ggplot:

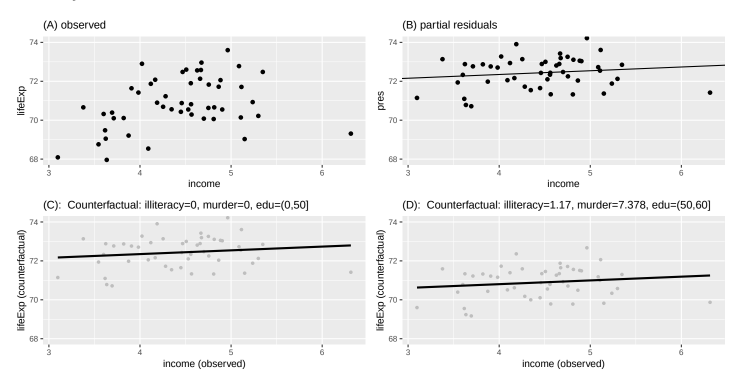
or directly via the plot function:

```
plot(e.lm, type = "partial", variable = c("(Intercept)","income")) # C
plot(e.lm, type = "partial", variable = c("(Intercept)","income"),
    at = data.frame(illiteracy = 1.17, murder = 7.378, edu = "(50,60]")) # D
```

These can be compared to displaying the observed outcome vs. income:

```
gg.obs <- ggplot(df1) + geom_point(aes(x=income, y=lifeExp))
gg.obs <- gg.obs + ggtitle("(A) observed")
gg.obs</pre>
```

where it is apparent that by using the partial residuals, the data has been normalized and exhibit less variability.



The output of the plot method is a list containing an element plot with the ggplot object and an element data with the dataset. To avoid actually displaying the graph one can use the method autoplot to only save the ggplot object:

```
ls.plot <- autoplot(e.lm, type = "partial", variable = c("(Intercept)", "income"))
lapply(ls.plot, class)</pre>
```

```
$data
```

[1] "residuals_lmm" "data.frame"

\$plot

[1] "gg" "ggplot"

One can re-create the plot based on the data argument or modify the existing plot, e.g. displaying with the y axis between 68 and 74:

```
ls.plot$plot + coord_cartesian(ylim=c(68,74))
```

1.2 What about confidence intervals?

A common question is whether one can display confidence intervals for the regression line. It is possible to add confidence intervals on the plot either via the argument ci.alpha:

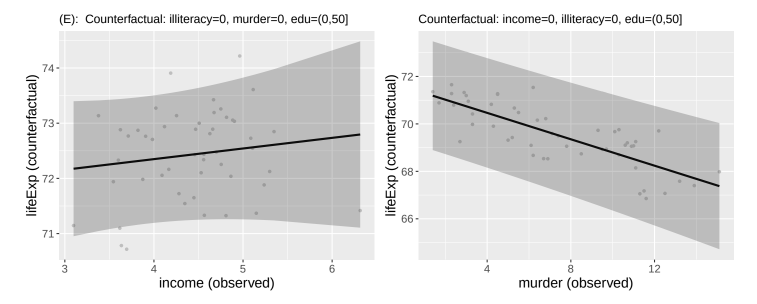
```
plot(e.lm, type = "partial", variable = c("(Intercept)","income"), ci.alpha = 0.25) ## E
```

or by requesting confidence intervals for the fitted lines via the argument pres.ci when calling residuals:

```
lifeExp illiteracy income murder
                                         edu fitted fitted.lower fitted.upper r.partial
1
    69.05
                    0
                        3.624
                                    0 (0,50] 72.277
                                                            71.115
                                                                          73.439
                                                                                     72.882
2
    69.31
                    0
                       6.315
                                      (0,50] 72.795
                                                            71.107
                                                                          74.483
                                                                                     71.417
3
    70.55
                       4.530
                                     (0,50] 72.451
                                                            71.253
                                                                          73.649
                                                                                     72.102
                    0
                                    0 (0,50] 72.229
4
    70.66
                    0
                        3.378
                                                            71.046
                                                                          73.413
                                                                                     73.136
5
    71.71
                        5.114
                                     (0,50] 72.564
                                                            71.254
                                                                          73.874
                                                                                     73.609
6
    72.06
                        4.884
                                    0 (0,50] 72.519
                                                            71.261
                                                                          73.778
                                                                                     73.056
```

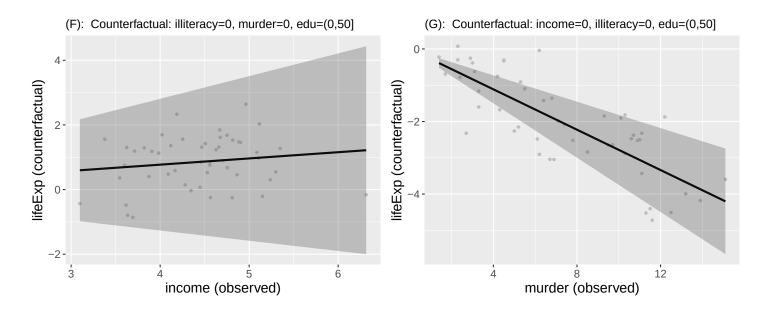
which can be added to the previous graphical display, e.g.:

The first plot is displayed in the left panel of the figure below. A similar partial residual plot but now for the murder variable is displayed in the right panel.



In many case the uncertainty represented here is of little interest, since it is the uncertainty of the intercept plus the exposure effect. This is why even though the murder variable was highly significant (p<0.001) whereas the income variable was not significant (p=0.45) the confidence intervals looks large in both cases. To only capture the uncertainty relative to the income or murder variable one should remove the intercept value, e.g. by omitting "(Intercept)" from the var argument:

```
plot(e.lm, type = "partial", variable = "income", ci.alpha = 0.25) ## F
plot(e.lm, type = "partial", variable = "murder", ci.alpha = 0.25) ## G
```



The unpleasant side effect is that the range of values on the y-axis appears unrealistic now. The statistical uncertainty may therefore be better communicated otherwise, e.g. reporting confidence intervals or p-values related to the covariate effect and keeping the partial residual plot free of confidence intervals.

1.3 Interaction with a categorical variable

Suppose that we are now interested in relating life expectancy (Y) to both income (X_1) for various level of education $(X_2 \in \{a, b, c\})$, adjusting for other variables such as illiteracy (Z_1) and murder rate (Z_2) . As before we assume a linear effect for all variables:

$$Y = \alpha + \beta_{1a} X_1 \mathbb{1}_{X_2 = a} + \beta_{1b} X_1 \mathbb{1}_{X_2 = b} + \beta_{1c} X_1 \mathbb{1}_{X_2 = c} + \gamma_1 Z_1 + \gamma_2 Z_2 + \varepsilon$$

where $\mathbb{1}_x$ denotes the indicator variable taking value 1 when x is true and 0 otherwise. This model can be estimated with the following R code

```
e.lmI <- lmm(lifeExp \sim income:edu + illiteracy + murder, data = df1) model.tables(e.lmI)
```

```
df
                   estimate
                                                lower
                                                                   p.value
                                   se
                                                         upper
(Intercept)
                   71.78584 1.209517 44.009 69.34823 74.22344 0.0000e+00
                    0.12870 0.319145 44.009 -0.51449
illiteracy
                                                       0.77189 6.8871e-01
                   -0.27940 0.048208 44.009 -0.37656 -0.18224 6.7276e-07
murder
income:edu(0,50]
                    0.17147 0.297725 44.009 -0.42855
                                                       0.77149 5.6760e-01
income:edu(50,60]
                    0.22526 0.252110 44.009 -0.28284
                                                       0.73335 3.7646e-01
income:edu(60,100]
                    0.30377 0.236929 44.009 -0.17373
                                                       0.78126 2.0652e-01
```

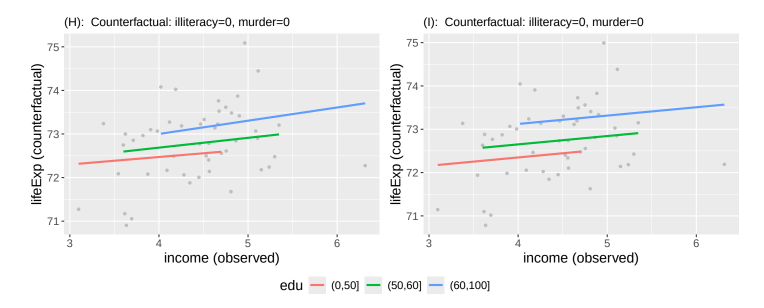
<u>Note:</u> this model is the same as $lmm(lifeExp \sim income*edu + illiteracy + murder, data = df1) but uses a different parametrisation.$

Similarly as before, we can use the plot function to display the partial residuals with respect to both income and edu:

```
plot(e.lmI, type = "partial", variable = c("(Intercept)", "income", "edu")) ## H
```

which can be compared to a plot assuming no interaction:

```
plot(e.lm, type = "partial", variable = c("(Intercept)","income","edu")) ## I
```



The partial residuals can also be output via the residuals method:

```
residuals(e.lmI, type = "partial", variable = c("(Intercept)", "income", "edu"))[1:5]
```

[1] 72.999 72.274 72.498 73.237 74.446

and one can check that they are evaluated by substracting the effect of the other variables (here illiteracy and murder), e.g.:

```
c(69.05 - 0.12870 * 2.1 - (-0.27940) * 15.1,
69.31 - 0.12870 * 1.5 - (-0.27940) * 11.3)
```

[1] 72.999 72.274

Here we computed partial residuals representing the life expectancy in the states had there be no murder nor illiteracy. We could also consider the case of average murder rate and illiteracy:

[1] 71.088 70.363 70.587 71.327 72.535

which we can also retrieve by hand:

```
c(69.05 - 0.12870 * (2.1-1.170) - (-0.27940) * (15.1-7.378),
69.31 - 0.12870 * (1.5-1.170) - (-0.27940) * (11.3-7.378))
```

[1] 71.088 70.363

2 Linear mixed model

To illustrate the use of partial residuals we will use data from a two-arm randomized trial comparing the quality of the vision over time of patients under placebo vs. active drug. We first re-shape the data:

and notice that the outcome (visual) and the covariate lesion can be missing:

```
summarizeNA(armd.long)
```

frequency	${\tt missing.pattern}$	${\tt n.missing}$	subject	lesion	line0	<pre>treat.f</pre>	miss.pat	${\tt week.num}$	visual	id	week
1106	000000000	0	0	0	0	0	0	0	0	0	0
89	00000100	1	0	0	0	0	0	0	1	0	0
1	010000000	1	0	1	0	0	0	0	0	0	0
4	010000100	2	0	1	0	0	0	0	1	0	0

This is why a warning is displayed when fitting the linear mixed model:

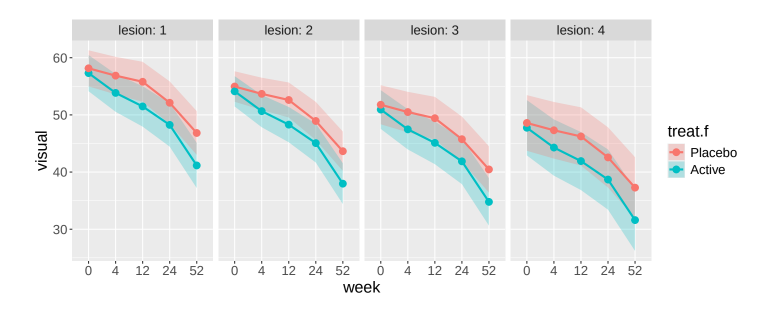
```
e.lmm <- lmm(visual \sim week*treat.f + lesion, data = armd.long, repetition =\simweek|subject)
```

Warning message:

```
In .lmmNormalizeData(as.data.frame(data)[unique(stats::na.omit(var.all))], :
   Can only handle missing values in the outcome variable visual.
   5 observations with missing values in "lesion" have been removed.
   1 cluster has been removed.
```

To visualize the model fit, we can display the fitted mean for each level of baseline lesion:

```
plot(e.lmm, facet = ~lesion, labeller = label_both, facet_ncol = 4)
```



We can retrive the fitted values from the estimated coefficients:

round(coef(e.lmm),2)

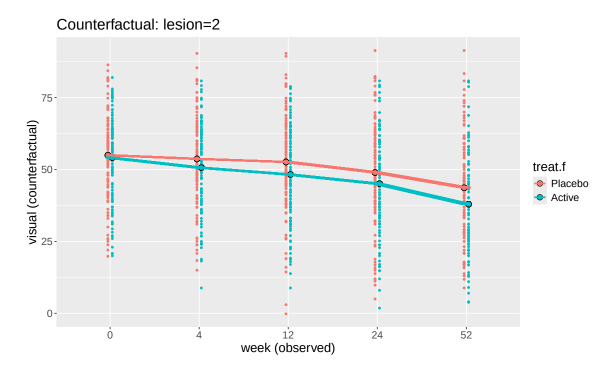
```
(Intercept)
                                                           week12
                                      week4
                                                                                 week24
               61.33
                                      -1.28
                                                            -2.35
                                                                                  -6.03
              week52
                             treat.fActive
                                                           lesion week4:treat.fActive
              -11.31
                                      -0.84
                                                            -3.19
                                                                                  -2.19
week12:treat.fActive week24:treat.fActive week52:treat.fActive
               -3.47
                                                            -4.84
                                     -3.03
```

- in the Placebo group with lesion=1, the estimated average baseline mean is (Intercept)+1*lesion, i.e. 61.33-3.19=58.14. When lesion=4, the estimated average baseline mean is (Intercept)+4*lesion, i.e. 61.33-4*3.19=48.57.
- the estimated average baseline mean in the Active group is shifted by treat.fActive i.e. -0.84 from the Placebo group.
- in the Placebo group with lesion=1, the estimated average week 52 mean is (Intercept)+week52+1*lesion, i.e. 61.33-11.31-3.19=46.83.
- the estimated average week 52 mean in the Active group is shifted by treat.fActive+week52:treat.fActive i.e. -0.84-4.84=-5.68 from the Placebo group.

Unfortunately, the display of the fitted value becomes overwhelming with considering more covariates or more covariate levels. Instead can visualize the partial residuals, e.g. here the outcome and fitted values had there be no lesion:

• on the same panel with a difference color for each treatment group:

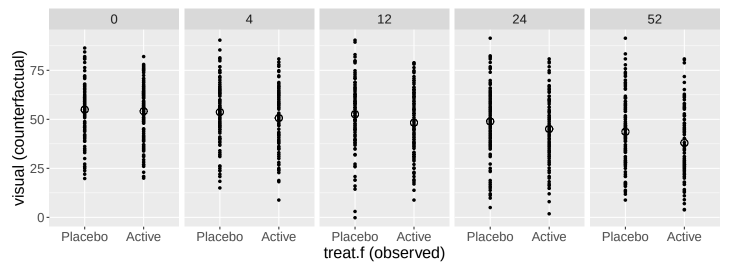
```
plot(e.lmm, type = "partial", variable = c("(Intercept)", "week", "treat.f"),
    at = data.frame(lesion = 2))
```



• on a separate panel for each timepoint:

```
plot(e.lmm, type = "partial", variable = c("(Intercept)", "week", "treat.f"),
    facet =~week, facet_nrow = 1, time.var = "treat.f", color = FALSE,
    at = data.frame(lesion = 2))
```

Counterfactual: lesion=2



The calculation of the partial residuals is similar to the univariate regression:

```
subject lesion lineO treat.f miss.pat week.num visual id week
                                                                           pres
1.0
           1
                  3
                                         --XX
                                                      0
                                                                       0 62.187
                        12
                            Active
                                                             59
                                                                 1
2.0
           2
                  1
                                         ----
                                                      0
                                                                 2
                        13
                            Active
                                                             65
                                                                       0 61.813
3.0
           3
                  4
                         8 Placebo
                                        ---X
                                                      0
                                                             40
                                                                 3
                                                                       0 46.373
4.0
           4
                  2
                        13 Placebo
                                        ----
                                                      0
                                                             67
                                                                       0 67.000
5.0
           5
                  1
                        14 Active
                                        XXXX
                                                      0
                                                             70
                                                                 5
                                                                       0 66.813
6.0
                  3
           6
                        12
                           Active
                                                      0
                                                             59
                                                                 6
                                                                       0 62.187
```

here substract the estimated lesion effect from the observed outcome:

```
c(59 - (-3.19) * (3-2),
65 - (-3.19) * (1-2))
```

[1] 62.19 61.81

In particular, the partial residuals for patient with lesion equal to two is the observed outcome.

3 R session

Details of the R session used to generate this document:

sessionInfo()

R version 4.3.3 (2024-02-29)

Platform: x86_64-pc-linux-gnu (64-bit) Running under: Ubuntu 22.04.4 LTS

Matrix products: default

BLAS: /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.10.0 LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0

locale:

[1] LC_CTYPE=en_US.UTF-8	LC_NUMERIC=C	LC_TIME=en_US.UTF-8
[4] LC_COLLATE=en_US.UTF-8	LC_MONETARY=en_US.UTF-8	LC_MESSAGES=en_US.UTF-8

[7] LC_PAPER=en_US.UTF-8 LC_NAME=C LC_ADDRESS=C

[10] LC_TELEPHONE=C LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

time zone: Europe/Copenhagen
tzcode source: system (glibc)

attached base packages:

[1] stats graphics grDevices utils datasets methods base

other attached packages:

[1] lava_1.8.0 nlme_3.1-163 LMMstar_1.1.0 lme4_1.1-35.2 Matrix_1.6-5 ggpubr_0.6.0

[7] ggplot2_3.5.1

loaded via a namespace (and not attached):

[1]	utf8_1.2.4	future_1.33.2	generics_0.1.3	tidyr_1.3.1
[5]	rstatix_0.7.2	lattice_0.22-5	listenv_0.9.1	digest_0.6.35
[9]	magrittr_2.0.3	grid_4.3.3	backports_1.4.1	survival_3.5-8
[13]	<pre>gridExtra_2.3</pre>	purrr_1.0.2	fansi_1.0.6	scales_1.3.0
[17]	numDeriv_2016.8-1.1	codetools_0.2-19	abind_1.4-5	cli_3.6.2
[21]	rlang_1.1.3	parallelly_1.37.1	<pre>future.apply_1.11.2</pre>	cowplot_1.1.3
[25]	munsell_0.5.1	splines_4.3.3	withr_3.0.0	tools_4.3.3
[29]	parallel_4.3.3	nloptr_2.0.3	ggsignif_0.6.4	minqa_1.2.6
[33]	dplyr_1.1.4	colorspace_2.1-0	boot_1.3-30	globals_0.16.3
[37]	broom_1.0.5	vctrs_0.6.5	R6_2.5.1	lifecycle_1.0.4
[41]	car_3.1-2	MASS_7.3-60.0.1	pkgconfig_2.0.3	pillar_1.9.0
[45]	gtable_0.3.5	glue_1.7.0	Rcpp_1.0.12	tibble_3.2.1
[49]	tidyselect_1.2.1	farver_2.1.1	labeling_0.4.3	carData_3.0-5
[53]	compiler_4.3.3			