

## **D** BSSE



## Introduction to Bayesian Statistics with R

8: Exercise solutions
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First we load the tidyverse, brms and set a seed.

```
library(tidyverse); options(dplyr.summarise.inform = FALSE) # suppress summarise warnings
library(brms)
set.seed(42)
```

## Optional Exercise 8.1 - Logistic regression

To run a logistic regression, we can use the glm() function with family = "binomial" (see details in ?stats::family) and formula = cbind(cancers, noncancers) ~ ...

- Fit a logistic regression model of cancer incidence with age\_s, sex, race, and registry, as explanatory variables (no interactions). Examine the model summary and coefficients.
- Use I (age\_s 2) to add a quadratic age\_s term to the model.
- Install the visreg package, and use visreg(..., "age\_s") to visualise the fitted slope of age\_s (x-axis) with respect to the log odds (y-axis). The points are the partial residuals with respect to age\_s. Does the model fit and visreg plot change for the better when including the quadratic term?

After we load the data

```
## Call:
## glm(formula = cbind(cancers, noncancers) ~ age_s + sex + race +
       registry, family = "binomial", data = CRC_df)
##
##
## Deviance Residuals:
##
        Min
                   1Q
                         Median
                                        3Q
                                                 Max
## -12.0480
             -1.2495
                         0.3743
                                    1.5614
                                              8.4957
##
## Coefficients:
```

```
##
                Estimate Std. Error
                                      z value Pr(>|z|)
## (Intercept) -7.740603
                           0.004924 -1572.071 < 2e-16 ***
                                               < 2e-16 ***
## age s
                0.873127
                           0.001333
                                      655.139
## sexfemale
               -0.328223
                           0.003242
                                     -101.249
                                               < 2e-16 ***
## raceblack
                0.217360
                           0.005805
                                       37.441
                                               < 2e-16 ***
                                       -4.903 9.43e-07 ***
## raceother
               -0.036421
                           0.007428
## registry02
                0.151314
                           0.005807
                                       26.056 < 2e-16 ***
## registry20
                0.061408
                           0.005776
                                       10.632
                                               < 2e-16 ***
## registry21
                0.137047
                           0.009304
                                       14.729
                                               < 2e-16 ***
## registry22
                0.127811
                           0.006039
                                       21.164
                                               < 2e-16 ***
## registry23
               -0.153910
                           0.008205
                                      -18.759
                                               < 2e-16 ***
## registry25
               -0.040474
                           0.006113
                                       -6.621 3.58e-11 ***
## registry26
               -0.209244
                           0.008556
                                      -24.455 < 2e-16 ***
## registry27
                                       -6.266 3.71e-10 ***
               -0.046357
                           0.007399
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 605833
                              on 3059
                                       degrees of freedom
## Residual deviance: 19925
                              on 3047
                                       degrees of freedom
## AIC: 35111
##
## Number of Fisher Scoring iterations: 4
```

The regression coefficients are all very significant predictors of cancer incidence, with each coefficient representing the change in log-odds of cancer for a unit change in the continuous variable age\_s or a change in level (compared to reference) for the categorical variables.

To include the quadratic term in age\_s, we use the suggested syntax and obtain the following regression results:

```
glm_fit2 <- glm(formula = cbind(cancers, noncancers) ~ age_s + sex + race + registry +</pre>
                  I(age_s^2), family = "binomial", data = CRC_df)
summary(glm_fit2)
##
## Call:
   glm(formula = cbind(cancers, noncancers) ~ age_s + sex + race +
       registry + I(age_s^2), family = "binomial", data = CRC_df)
##
##
## Deviance Residuals:
##
       Min
                      Median
                                    30
                 10
                                            Max
                      0.6021
## -5.8771
           -0.5817
                                1.5625
                                         5.8589
##
## Coefficients:
##
                Estimate Std. Error
                                       z value Pr(>|z|)
## (Intercept) -7.658026
                            0.005032 -1521.894
                                               < 2e-16 ***
                            0.002598
                                       411.804
## age_s
                1.069954
                                                < 2e-16 ***
## sexfemale
               -0.319354
                           0.003242
                                       -98.517
                                                < 2e-16 ***
## raceblack
                0.205940
                           0.005800
                                        35.504 < 2e-16 ***
                            0.007401
                                        -6.029 1.65e-09 ***
## raceother
               -0.044624
## registry02
                0.148727
                            0.005805
                                        25.620
                                                < 2e-16 ***
## registry20
                0.058935
                            0.005773
                                        10.209
                                                < 2e-16 ***
```

## registry21

## registry22

0.131618

0.131343

0.009281

0.006037

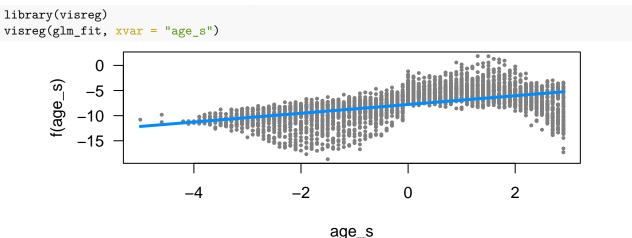
14.182 < 2e-16 \*\*\*

21.755 < 2e-16 \*\*\*

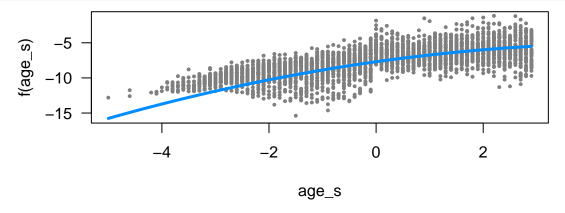
```
## registry23
               -0.163881
                           0.008204
                                       -19.977 < 2e-16 ***
## registry25
               -0.041858
                           0.006112
                                        -6.849 7.46e-12 ***
               -0.206759
## registry26
                           0.008555
                                       -24.168
                                                < 2e-16
                           0.007392
                                        -6.499 8.10e-11 ***
## registry27
               -0.048040
##
  I(age s^2)
               -0.108658
                           0.001084
                                      -100.194
                                                < 2e-16 ***
##
                           0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
##
   (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 605833.5
                                 on 3059
                                          degrees of freedom
## Residual deviance:
                                 on 3046
                                          degrees of freedom
                        8430.5
  AIC: 23619
##
##
## Number of Fisher Scoring iterations: 4
```

There are slight changes to all categorical regression coefficients, and an obvious large change for the age\_s and (Intercept) now we have the quadratic term. The quadratic term for age\_s is also a highly significant predictor, suggesting already that the quadratic dependence on age\_s is a better fit than a linear model.

When we visualise the models with visreg, without the quadratic term we get the following plot



where the partial residuals suggest a nonlinear trend in age. With the quadratic term, this looks a lot better: visreg(glm\_fit2, xvar = "age\_s")



## Exercise 8.2 - Bayesian logistic regression

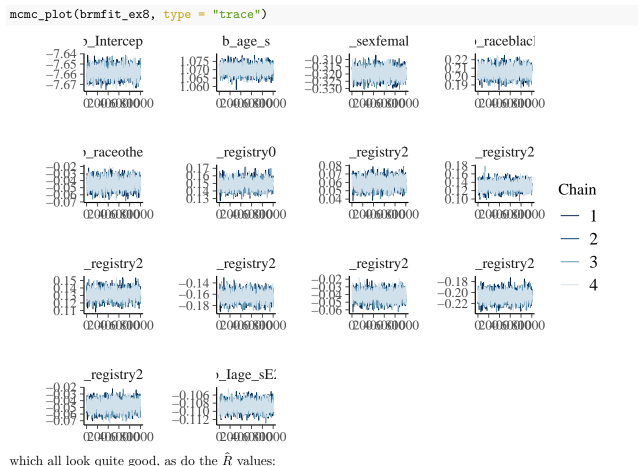
For Bayesian modelling with brms we can use the brm() function with family = binomial, but with a somewhat different syntax for the formula. We separate the number of occurrences from the number of trials (input into the trials function) with I and

formula = cancers / trials(total) ~ ...

- Fit a Bayesian logistic regression model of cancer incidence with age\_s, sex, race, and registry as explanatory variables (no interactions). Include I(age\_s^2) to add a quadratic age\_s term to the model.
- Check the model convergence.
- If you've done Exercise 8.1, compare the regression coefficients.
- What is the posterior distribution of the probability of having cancer for a 75 year-old Black female from registry 27?

Running the model we use the following syntax (with our helper function in the background)

With so much data, we didn't worry too much about the default prior (especially as we use the rescaled age), and first we check the trace plots



rhat(brmfit\_ex8)

```
## b_Intercept b_age_s b_sexfemale b_raceblack b_raceother b_registry02
```

```
##
      1.0015216
                   1.0003160
                                0.9992567
                                              1.0002368
                                                           1.0005167
                                                                         1.0007795
## b_registry20 b_registry21 b_registry22 b_registry23 b_registry25 b_registry26
                                                                         1.0010904
      1.0009280
                   0.9998673
                                1.0008285
                                              1.0006513
                                                           1.0002542
                                    lprior
## b_registry27
                  b_Iage_sE2
                                                   lp__
      1.0002933
                   1.0002461
                                1.0015002
                                              1.0014880
```

and the effective sample sizes:

```
summary(brmfit_ex8)$fixed$Bulk_ESS
```

```
## [1] 1522.641 2942.224 5624.633 3140.737 2030.632 1646.479 1742.972 2103.487
## [9] 1797.273 2096.491 1789.214 2129.869 1990.103 3181.412
summary(brmfit_ex8)$fixed$Tail_ESS
```

```
## [1] 2477.065 3061.518 2881.652 2791.567 2433.148 2310.909 2531.736 2415.921 ## [9] 2629.364 2317.694 2652.390 2671.945 2591.459 2991.736
```

If we look at the regression coefficients

```
fixef(brmfit_ex8)
```

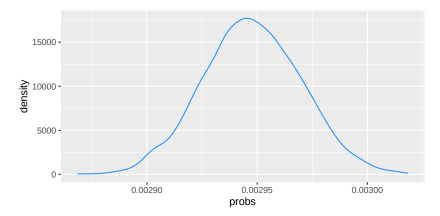
```
##
                Estimate
                           Est.Error
                                            Q2.5
                                                       Q97.5
## Intercept -7.65818507 0.005119319 -7.66820613 -7.64814300
              1.07008870 0.002562819 1.06503310 1.07499561
## age s
## sexfemale -0.31939029 0.003313728 -0.32606753 -0.31313105
## raceblack 0.20584732 0.005775137 0.19447440 0.21703037
## raceother -0.04458297 0.007262838 -0.05879977 -0.02998851
## registry02 0.14878567 0.006030355 0.13701165 0.16035380
## registry20 0.05915472 0.006004347 0.04756327 0.07069752
## registry21 0.13163374 0.009296847 0.11326611 0.15011583
## registry22 0.13142994 0.006189252 0.11953835 0.14352611
## registry23 -0.16373909 0.008493930 -0.18013640 -0.14688746
## registry25 -0.04166973 0.006344099 -0.05402193 -0.02927285
## registry26 -0.20680949 0.008591501 -0.22358324 -0.18985897
## registry27 -0.04785530 0.007448580 -0.06228802 -0.03342212
             -0.10871154 0.001083747 -0.11087805 -0.10655925
## Iage sE2
```

these are very similar to the logistic regression before, but now from these we can extract posterior estimates. For example, for a 75 year old Black female from registry 27, from the posterior samples of the regression coefficients we would have the following mapping to the sampled log-odds

```
b Intercept +2.5*b age s+b sexfemale+b raceblack +b registry 27+2.5^2*b Iage sE2
```

From the posterior samples we can therefore create a new column of the log-odds, which we transform back to probabilities with the inverse-logit, or expit, function

```
expit <- function(x) { # inverse logit
  exp(x)/(1 + exp(x))
}
as_draws_df(brmfit_ex8) %>% mutate(log_odds = b_Intercept + 2.5*b_age_s + b_sexfemale +
  b_raceblack + b_registry27 + 2.5^2*b_Iage_sE2,
  probs = expit(log_odds)) %>%
  ggplot(aes(probs)) + geom_density(color = "dodgerblue")
```



The resulting probabilities are mostly between 0.0029 and 0.003, which is closely aligned with the observed frequency of cancer in that stratum:

```
CRC_df %>% filter(age == 75, sex == "female", race == "black", registry == 27) %>%
mutate(prob = cancers/total)
```

```
## # A tibble: 1 x 9
## age sex race registry cancers noncancers total age_s prob
## <dbl> <fct> <fct> <chr> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <2000 2.5 0.00283</pre>
```

Rather than doing this by hand, we can use the posterior\_predict function (or rather the posterior\_linpred function for the linear modelling part) on new data. For this we make a small data frame of our individual

```
newdata <- data.frame(
   age_s = c(2.5), #transformed age
   sex = factor("female"),
   race = factor("black"),
   registry = c("27"),
   total = 1e5 # dummy value needed to make the functions work
)</pre>
```

and pass it into posterior\_linpred with the argument transform = TRUE to output the results in the probability space (rather than the logit-space)

```
posterior_linpred(brmfit_ex8, newdata, transform = TRUE) %>% data.frame(probs = .) %>%
    ggplot(aes(probs)) + geom_density(color = "dodgerblue")
```

