Bayesian exposure-response modeling for binary data

Outline

- ▶ Brief introduction to Bayesian analysis
- Fitting models
- ▶ Model checking

Approach to Bayesian modeling in this course

- ► For this series of classes we are going to use Stan to do Bayesian modeling
 - ► Stan is a probabilistic programming language for fitting Bayesian models.
 - ▶ By default it uses Hamiltonian Monte Carlo (HMC), specifically a variation called the no U-turn sampler (NUTS).
 - We will go into more details about HMC/NUTS later in the course
- Start with using brms as our gateway to Stan
 - brms is a package that enables simple fitting of many types models that you can fit using glm, survreg, lme, nlme, etc.
 - Allows quick access to Bayesian inference
- ► Later we will program our own Stan models and run them with rstan
 - Learn more about the Stan language
 - Fit models that are not supported in brms

A brief review of Bayesian inference

Bayes Rule is the basis for inference about model parameters θ given data y and prior knowledge about model parameters $p(\theta)$:

$$p(\theta | y) = \frac{p(\theta)p(y | \theta)}{p(y)} = \frac{p(\theta)p(y | \theta)}{\int p(\theta)p(y | \theta)d\theta}$$
$$\propto p(\theta) p(y | \theta)$$

Goals:

- ▶ Inference about θ or a function of θ
- Predictions of future observations
- The posterior summarizes what we know about θ , but typically we can't express $p(\theta | y)$ in closed-form.
 - We'll use Markov Chain Monte Carlo to obtain samples from $p(\theta | y)$.

Bayesian modeling/inference process using MCMC

- 1. Construct a model for the data, conditional on parameters θ , $p\left(y\mid\theta\right)$
- 2. Construct a prior distribution for θ , $p(\theta)$
 - Ideally based on all available evidence/knowledge (or belief)
 - Or deliberately select a non-informative (or weakly informative) prior
- 3. Sample from the posterior distribution for θ , $p(\theta | y)$.
 - Look at convergence and sampler diagnostics
 - Use for inferences regarding parameter values
- 4. Sample from the posterior predictive distribution for y_{new} : $p(y_{\text{new}} | y) = \int p(y_{\text{new}} | \theta) p(\theta | y) d\theta$.
 - ▶ Use for inferences regarding future observations
 - ▶ Sample from $p(y | \theta)$ for values of θ from step 3.

Bayesian ingredients for MCMC sampling from a posterior

- Data
- ► Model for the outcome(s) the likelihood
- ▶ Models for the parameters the prior distribution
- ► MCMC tool Stan (via brms or 'rstan")

Ingredients for HMC/NUTS

- A starting point in the parameter space (initial value, one per chain)
- Number of MCMC samples used to tune the HMC/NUTS algorithm (warmup)
 - ► This is not exactly the same as the burn-in in other MCMC algorithms
 - ▶ NUTS uses these samples to adaptively tune the sampler
- ► Total number of samples to take, including the warm-up (iter)
- ▶ Parameters which inform how the sampler should adapt
 - ▶ Defaults are usually good for 'simple' models
 - ▶ Often need to modify for more complex, hierarchical models
 - Return to these later in the course

Let's re-fit our AE model using brms

What about the prior distributions?

- By default, brms uses flat, non-informative prior distributions for regression coefficients
- We can specify priors directly through the prior argument.
 - ► More to come in a few slides

Model summary

summary(mod01_stan)

```
Family: bernoulli
   Links: mu = logit
. Formula: AEO1 ~ CAVGSS + BWT + PTTYPE + SEXTXT
    Data: aedat (Number of observations: 180)
   Draws: 4 chains, each with iter = 1000; warmup = 500; thin = 1;
         total post-warmup draws = 2000
. Population-Level Effects:
           Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
. Intercept
          -7.32
                      4.47
                             -16.50 1.00 1.00
                                                     995
                                                            1236
            0.86 0.19 0.52 1.26 1.00 1359
. CAVGSS
                                                           1324
             0.04 0.06 -0.08 0.16 1.01
BWT
                                                  925
                                                           1009
. PTTYPEPT1 1.52 0.75 0.17 3.05 1.00 1467
                                                           1227
          0.88 0.97 -1.05 2.74 1.00 1557
. PTTYPEPT2
                                                           1302
. SEXTXTMALE 0.07 0.89 -1.70 1.84 1.01
                                                    1141
                                                            1256
. Draws were sampled using sampling (NUTS). For each parameter, Bulk ESS
. and Tail ESS are effective sample size measures, and Rhat is the potential
. scale reduction factor on split chains (at convergence, Rhat = 1).
```

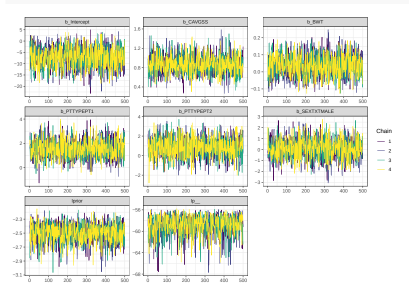
▶ Defaults: Estimate = posterior mean and Est.Err = posterior sd. Can modify these using the robust argument to brm

MCMC convergence diagnostics

- Traceplots
 - Plot of sampled values vs iteration
 - Look for stationarity and good mixing: fuzzy caterpillar
- ► Â
 - ightharpoonup Heuristically: $\frac{\text{total variance of } θ \text{ (between and within-chain)}}{\text{average within-chain variance of } θ}$
 - ▶ Target: $\hat{R} < 1.01$ (sometimes, you'll see a rule of $\hat{R} < 1.05$)
 - Output: Summary and plot (mcmc_plot(mod01_stan, type='rhat'))
- Effective sample sizes
 - bulk ESS for assessing posterior means, medians, etc
 - tail ESS for assessing tail percentiles (5th, 95th)
 - Target: Depends on your goals
 - Output: Summary

Traceplots

bayesplot::mcmc_trace(mod01_stan)



Let's see the default priors in our model

```
prior_summary(mod01_stan)
                  prior
                             class
                                         coef group resp dpar nlpar lb ub
                  (flat)
                  (flat)
                                 b
                                          BWT
                 (flat)
                                        CAVGSS
                  (flat)
                                 b PTTYPEPT1
                  (flat)
                                    PTTYPEPT2
                  (flat)
                                 b SEXTXTMALE
   student_t(3, 0, 2.5) Intercept
         source
        default
   (vectorized)
   (vectorized)
   (vectorized)
   (vectorized)
   (vectorized)
        default
```

BRMS centers all predictors in the model

Mathematically, the RHS of the model $y \sim x1 + x2$ is

$$b_Intercept + b_x1 \cdot x1 + b_x2 \cdot x2$$

Or, equivalently,

$$Intercept + b_x1 \cdot (x1 - \overline{x1}) + b_x2 \cdot (x2 - \overline{x2})$$

where

$$b_Intercept = Intercept - b_x 1 \cdot \overline{x1} - b_x 2 \cdot \overline{x2}$$

This is the parameterization that brms uses. (Why do you think that is?)

We need to specify priors for Intercept, b_x1 and b_x2

To change them use the set_priors function

A Normal(mean=0, sd=5) prior on all covariate effects:

```
priors_mod01 <- set_prior('normal(0,5)', class='b')</pre>
```

A Normal(0,5) prior on CAVGSS and a DoubleExponential prior on the other effects:

```
priors_mod01_de <- set_prior('normal(0,5)', class='b', coef='CAVGSS') +
    set_prior('double_exponential(0,1)', class='b')</pre>
```

See Stan functions reference for list of all available distributions.

Workbook Bayes01

- ► Model fitting in brms
- ► Convergence diagnostics

Model diagnostics

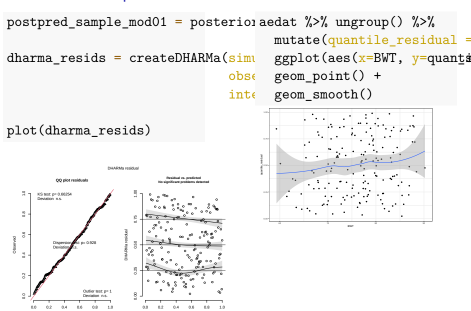
- We can use similar diagnostics as with likelihood based methods, but now using posterior predictive distributions
 - Quantile residual plots
 - Posterior predictive checks

Quantile residuals

- 1. Simulate from posterior predictive distribution
 - brms has a posterior_predict function to generate samples in a matrix
- Use createDHARMa function in the DHARMa package to format output
 - Input is a matrix of posterior samples and the observed outcome data
- 3. Make plots as before
 - Residuals vs predicted
 - Residuals vs predictors

DHARMa examples

Fxpected



Model predictions (rank transformed)

Posterior predictive checks

- 1. Simulate from posterior predictive distribution
 - tidy_bayes has an add_predicted_draws function to append the samples to a data frame
- Compute some summary statistic on each posterior draw and on the observed data
 - Summary statistic depends on what you want to diagnose
 - ► For binary models, it is almost always the expected value (mean)
- 3. Plot distribution of summary statistics and overlay observed values
 - Type of plot depends on grouping factor and summary statistic

Simulate from posterior predictive distribution

```
aedat_pp = add_predicted_draws(newdata = aedat, mod01_stan)
aedat_pp %>% ungroup() %>%
select(USUBJID, PTTYPE, AE01, Quartile:.prediction) %>%
slice_tail(n=4)
```

```
. # A tibble: 4 x 9
   USUBJID PTTYPE AEO1 Quartile .row .chain .iteration .draw .prediction
   <fct> <fct> <int> <chr>
                              <int> <int>
                                               <int> <int>
                                                               <int>
. 1 IITD-180 PT1
                    0 02
                               180
                                       NΑ
                                                 NA 1997
                                                                   0
. 2 UID-180 PT1 0 Q2
                               180
                                       NA
                                                 NA 1998
. 3 UTD-180 PT1
                    0 Q2
                                180
                                       NA
                                                  NA 1999
. 4 UID-180 PT1
                    0 02
                                180
                                       NA
                                                  NA 2000
```

Plot the VPC

```
# Observed data summary
obs_summary <- aedat %>%
group_by(Quartile) %>%
summarise(phat_obs = mean(AEO1))

#Simulated data summary
sim_summary <- aedat_pp %>%
group_by(.draw,Quartile) %>%
summarise(phat_sim = mean(.prediction))

# VPC
sim_summary %>%
ggplot(aes(x=Quartile, y=phat_sim)) +
geom_point(data=obs_summary, aes(y=phat_obs), labs(x='', y='Proportion with AE') +
coord_flip()
```

VPC for continuous variable: define summary statistic

- Fit generalized additive model (smoother) to each simulated dataset
- Predict at a fixed grid of values (0 to 95th percentile)

Compute summary statistics for observed data

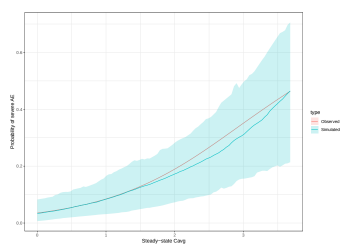
- . 1 0.0000000 0.03526032 Observed . 2 0.03718346 0.03644987 Observed . 3 0.07436693 0.03767799 Observed . 4 0.11155039 0.03894582 Observed . 5 0.14873385 0.04025454 Observed
- . 6 0.18591731 0.04160537 Observed

Compute summary on each simulated study

```
sim_summary <- aedat_pp %>%
  # Nest everying except the simulation name
 nest(cols=-.draw) %>%
  # Use 200 sims for demonstration
 sample_n(size=200) %>%
  # Compute summary stats for each simulated dataset
 mutate(predictions = map(cols, ~summary_function(.x,
                                                    .x name='CAVGSS'.
                                                    .v name='.prediction'))) %>%
 select(.draw,predictions) %>%
 unnest(cols=predictions) %>%
  # Summarise across simulated data sets
 group_by(xvar) %>%
  summarise(glo = quantile(prediction, probs = 0.05),
            qhi = quantile(prediction, probs = 0.95),
            prediction=median(prediction)
            ) %>%
 mutate(type = 'Simulated')
```

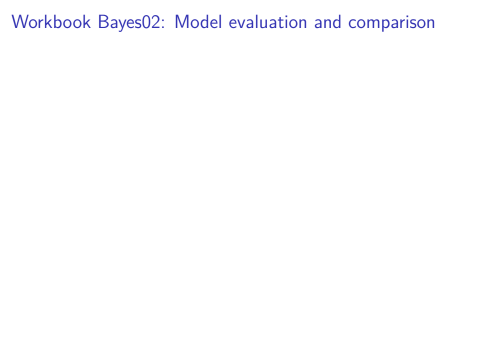
Plot VPC

```
sim_summary %>% bind_rows(obs_summary) %>%
ggplot(aes(x=xvar, y=prediction)) +
geom_line(aes(col=type, group=type)) +
geom_ribbon(aes(ymin=qlo, ymax=qdhi, fill=type), alpha=0.2) +
labs(x='Steady-state Cavg', y='Probability of severe AE')
```



Model Comparison

- Goal: maximize expected log predictive density (ELPD) for future data
 - This is a measure of out-of-sample prediction quality
- Leave-one-out cross-validation (LOO-CV) to approximate ELPD
 - Involves fitting N models
 - Can be approximated using pareto smoothed importance sampling of the posterior samples
 - ▶ loo package
- ► WAIC also approximates -2 ELPD
 - Proven to be asymptotocally equivalent to LOO-CV (modulo the -2)
 - Lower is better
 - ► LOO-CV generally preferable due to its ability to tell when the estimates are not trustworthy



References