Rstanarm - easy implementation of Bayesian analysis

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Why consider a Baysian analysis?

- Jarad (Lunchinators, 8 Feb 2019): for the interpretation
 - Credible intervals are what we all want confidence intervals to be
- Other reasons
 - Account for all sources of uncertainty (more below)
 - Inference on derived quantities (see below)
 - Small sample inference for many problems (get rid of the z score)
 - Can easily fit more realistic biological models (hierarchical modeling, not discussed here)

The Bayesian paradigm:

• prior + likelihood + data -> posterior

$$f(\theta \mid data) = \frac{f(data \mid \theta) \ g(\theta)}{\int f(data \mid \theta) \ g(\theta)}$$

The integral in the denominator can be very difficult! Could be very high dimensional.

Long ago (pre 1990): Bayes restricted to combinations of prior and model (likelihood) with analytic integrals (conjugate priors)

1990's: MCMC revolution. Can sample from the posterior distribution without integration

* Gibbs sampler, Metropolis-Hasting sampler

How do I do a Bayesian analysis?

- 1. Think about how the data relate to my question(s)? i.e., what are the relevant parameters?
- 2. Write out an appropriate model. specifies how data relate to parameters
- 3. Think about what I believe before seeing the data (prior distribution(s) for parameters)

Then

- 1. Do some math, write your own samplers
- 2. BUGS / WinBUGS: code the model and priors, requires loops
 - RWinBUGS: R interface to WinBUGS. Very inefficient.
- 3. JAGS: better implementation, still code the model and priors
 - rjags: R interface to JAGS much smoother than RwinBUGS
- 4. STAN: new samplers (Hamiltonian MC), much faster!
 - RStan: R interface to STAN I haven't used
 - rstanarm: uses R modeling language to write models

rstanarm: Implements many R models, including

- 1. Linear models
- 2. Generalized linear models
- 3. Linear mixed effect models
- 4. many others

Simple example: Regress Y on X and X²

```
nobs <- 20
test <- data.frame(x=runif(nobs, 2, 10))</pre>
testy <- 10 - 0.2*(test$x-5)^2 + rnorm(nobs,0,1)
test$x2 <- test$x^2
y.lm \leftarrow lm(y \sim x + x2, data=test)
summary(y.lm)$coefficients
##
                Estimate Std. Error
                                       t value
                                                    Pr(>|t|)
## (Intercept) 3.431303 1.70192725 2.016128 5.987138e-02
                2.853089 0.63641828 4.483041 3.274161e-04
## x
## x2
               -0.289506 0.05391124 -5.370049 5.090503e-05
confint(y.lm)
                     2.5 %
##
                               97.5 %
## (Intercept) -0.1594500 7.0220553
                1.5103640 4.1958144
## x
## x2
               -0.4032488 -0.1757632
y.stan <- stan_glm(
    y \sim x + x2,
    family=gaussian,
    data=test,
    chains = 4,
    cores = 4
 )
```

Explanation of code:

- * stan modeling functions are stan_(R function), e.g., stan_lm, stan_glm, stan_lmer
- * write model as you would write the R model, including data=
- * glm with family=gaussian is an lm
- * stan lm requires priors (specify r^2 for each variable)
- * stan_glm/gaussian allows different priors and has a reasonable default
- * chains= specifies how many independent chains to sample, 3 or 4 are common choices
- * cores= specifies how many cores to use for parallel processing. Do not exceed 1/2 to 2/3 number on your machine

How many cores does my machine have?

```
library(parallel)
detectCores()
```

```
## [1] 8
```

What are the default prior distributions?

```
prior_summary(y.stan)
```

```
## Priors for model 'y.stan'
## ----
## Intercept (after predictors centered)
##
   ~ normal(location = 0, scale = 10)
##
        **adjusted scale = 20.02
##
## Coefficients
   ~ normal(location = [0,0], scale = [2.5,2.5])
##
##
        **adjusted scale = [2.26,0.19]
##
## Auxiliary (sigma)
   ~ exponential(rate = 1)
        **adjusted scale = 2.00 (adjusted rate = 1/adjusted scale)
##
## ----
## See help('prior_summary.stanreg') for more details
```

Change by adding prior = (for regression coeff.) prior_intercept = (for intercept) and/or prior_aux = (for sigma) to the stan_glm call

What can I do once I fit the model?

Diagnostics:

- biggest concern is whether the sampler has converged to the posterior distribution
- default is 1000 samples "warmup" (discarded), keep next 1000 samples (both per chain)
- want chains to look similar
- Rhat measures discrepancy between chains. Want close to 1.
 - Over 1.1 is usually considered bad unless model really hard to sample
- how many samples: want small MC standard error
- Graphical exploration rstanarm shiny app

sampling

see help('prior_summary')

4000 (posterior sample size)

```
launch_shinystan(y.stan)
```

algorithm:

priors:
sample:

There are also posterior predictive checks. compare data to predictions from the posterior distribution.

Summarize results, once fit looks reasonable

```
##
## Model Info:
##
## function: stan_glm
## family: gaussian [identity]
## formula: y ~ x + x2
```

```
observations: 20
##
    predictors:
##
## Estimates:
##
                    mean
                            sd
                                   2.5%
                                           25%
                                                  50%
                                                          75%
                                                                 97.5%
                                                  4.67
## (Intercept)
                    4.74
                            1.74
                                   1.44
                                           3.56
                                                         5.86
                                                                 8.34
                                                  2.36
                                                          2.79
## x
                    2.34
                            0.65
                                   1.01
                                           1.92
                                                                 3.57
## x2
                   -0.25
                            0.05
                                  -0.35
                                          -0.28
                                                 -0.25 -0.21
                                                                -0.14
## sigma
                    1.14
                            0.22
                                   0.82
                                           0.99
                                                  1.11
                                                          1.26
                                                                 1.66
## mean_PPD
                    8.97
                            0.37
                                   8.24
                                           8.73
                                                  8.97
                                                          9.21
                                                                 9.70
## log-posterior -38.90
                           1.61 -42.89 -39.67 -38.53 -37.71 -36.92
##
## Diagnostics:
##
                  mcse Rhat n_eff
## (Intercept)
                  0.04 1.00 1689
## x
                  0.02 1.00 1541
## x2
                  0.00 1.00 1526
## sigma
                  0.01 1.00 1756
                  0.01 1.00 2385
## mean_PPD
## log-posterior 0.05 1.01 1238
##
## For each parameter, mcse is Monte Carlo standard error, n_eff is a crude measure of effective sample
Why are estimates so different?
* rstanarm centers all X variables - reduces correlation of estimates
+ sampling the posterior is easier
* detail: mean of X^2 is not (mean X)^2, so should center variables ourselves (same principle for interactions)
Redo with hand-centered variables
test$xc <- test$x - mean(test$x)</pre>
test$xc2 <- test$xc^2</pre>
yc.lm \leftarrow lm(y \sim xc + xc2, data=test)
summary(yc.lm)
##
## lm(formula = y ~ xc + xc2, data = test)
##
## Residuals:
        Min
                   10
                        Median
                                       30
                                               Max
## -1.99560 -0.45389 -0.03835 0.67842
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 10.31769
                             0.34618
                                      29.804 4.07e-16 ***
## xc
                             0.11134 -3.654 0.00196 **
                -0.40684
## xc2
                -0.28951
                             0.05391
                                     -5.370 5.09e-05 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.06 on 17 degrees of freedom
## Multiple R-squared: 0.7493, Adjusted R-squared: 0.7198
## F-statistic: 25.4 on 2 and 17 DF, p-value: 7.818e-06
```

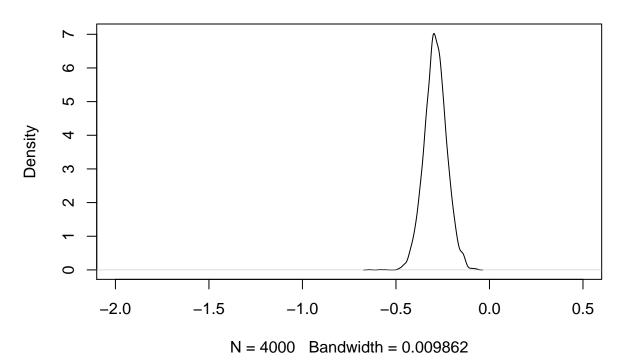
```
confint(yc.lm)
                     2.5 %
                               97.5 %
## (Intercept)
                9.5873070 11.0480647
## xc
                -0.6417553 -0.1719316
                -0.4032488 -0.1757632
## xc2
yc.stan <- stan_glm(</pre>
    y \sim xc + xc2,
    family=gaussian,
    data=test,
    chains = 4,
    cores = 4
  )
summary(yc.stan, digits=2)
##
## Model Info:
##
## function:
                   stan_glm
## family:
                   gaussian [identity]
## formula:
                   y \sim xc + xc2
## algorithm:
                   sampling
##
    priors:
                   see help('prior_summary')
##
    sample:
                   4000 (posterior sample size)
    observations: 20
##
    predictors:
##
## Estimates:
                                  2.5%
                                          25%
                                                 50%
                                                        75%
                                                                97.5%
                           sd
                   mean
## (Intercept)
                   10.31
                           0.38
                                  9.55
                                         10.07
                                                10.31 10.57
                                                               11.08
                                                -0.41 -0.33
## xc
                   -0.41
                           0.12 - 0.64
                                         -0.48
                                                               -0.17
## xc2
                   -0.29
                           0.06 - 0.41
                                        -0.33
                                                -0.29 -0.25
                                                               -0.17
## sigma
                   1.13
                           0.21
                                  0.80
                                          0.98
                                                 1.10
                                                        1.24
                                                                1.63
## mean_PPD
                   8.96
                           0.36
                                  8.26
                                          8.72
                                                 8.95
                                                        9.19
                                                                9.69
                           1.62 -41.34 -38.07 -36.95 -36.07 -35.28
## log-posterior -37.30
##
## Diagnostics:
                 mcse Rhat n_eff
## (Intercept)
                 0.01 1.00 3205
## xc
                  0.00 1.00 3545
                  0.00 1.00 2921
## xc2
## sigma
                 0.00 1.00 2497
## mean_PPD
                  0.01 1.00 3691
## log-posterior 0.04 1.00 1309
## For each parameter, mcse is Monte Carlo standard error, n_eff is a crude measure of effective sample
"
Can extract all the samples of the posterior distribution.
```

Very useful if you want a transformation of parameters, e.g. X where quadratic is min or max

```
y.post <- as.matrix(yc.stan)
head(y.post)</pre>
```

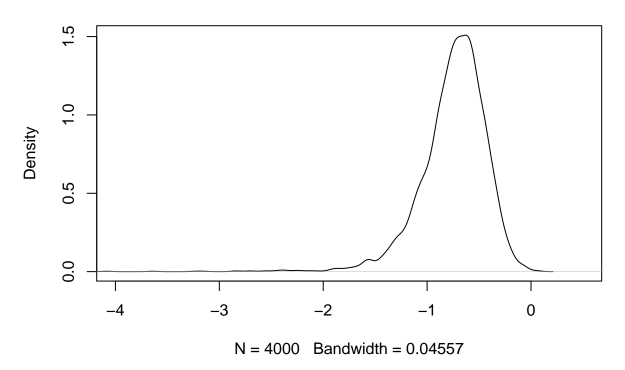
```
##
             parameters
  iterations (Intercept)
                                             xc2
                                                      sigma
                                   xc
         [1,]
                10.283755 -0.5736322 -0.2873951 1.1006859
##
##
         [2,]
                10.099205 -0.4159217 -0.2063932 0.8996655
         [3,]
                10.488654 -0.4163559 -0.2809807 1.0048600
##
##
         [4,]
                10.287575 -0.3453727 -0.3307754 1.0401509
                10.254443 -0.5048244 -0.2180562 1.1217078
##
         [5,]
         [6,]
                 9.858499 -0.4301444 -0.2240405 0.8044317
##
plot(density(y.post[,3]), xlim=c(-2,0.5))
```

density.default(x = y.post[, 3])



```
xmax <- -y.post[,2]/(2*y.post[,3])
plot(density(xmax), xlim=c(-4, 0.5))</pre>
```

density.default(x = xmax)



```
summary(xmax)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
## -5.38342 -0.90183 -0.71034 -0.75154 -0.54543 0.07354

quantile(xmax, c(0.025, 0.05, 0.5, 0.95, 0.975))

## 2.5% 5% 50% 95% 97.5%
## -1.4702041 -1.2960908 -0.7103427 -0.3299226 -0.2603170
```

Examples of other models fit with rstanarm

Generalized linear model

benefit of Bayes - appropriate inferences for small samples

```
donner <- read.csv('donner.csv')</pre>
donner.glm <- glm(survival ~ age + femc, data=donner,</pre>
                                                           family=binomial)
summary(donner.glm)
##
## glm(formula = survival ~ age + femc, family = binomial, data = donner)
##
## Deviance Residuals:
                       Median
                                     3Q
       Min
                  1Q
                                             Max
## -1.7445 -1.0441 -0.3029
                                          2.0472
                                0.8877
```

```
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) 3.23041
                           1.38686
                                     2.329
                                             0.0198 *
## age
               -0.07820
                           0.03728 -2.097
                                             0.0359 *
## femcM
              -1.59729
                           0.75547 -2.114
                                             0.0345 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 61.827 on 44 degrees of freedom
## Residual deviance: 51.256 on 42 degrees of freedom
## AIC: 57.256
##
## Number of Fisher Scoring iterations: 4
confint(donner.glm)
## Waiting for profiling to be done...
##
                    2.5 %
                               97.5 %
## (Intercept) 0.8514190 6.42669512
## age
               -0.1624377 -0.01406576
## femcM
              -3.2286705 -0.19510198
donner.stan <- stan_glm(</pre>
   survival ~ age + femc,
   family=binomial,
   data=donner,
   chains = 4,
    cores = 4
summary(donner.stan, digits=2)
##
## Model Info:
##
## function:
                  stan_glm
## family:
                  binomial [logit]
## formula:
                  survival ~ age + femc
## algorithm:
                  sampling
## priors:
                  see help('prior_summary')
##
   sample:
                  4000 (posterior sample size)
##
   observations: 45
##
   predictors:
##
## Estimates:
##
                          sd
                                 2.5%
                                        25%
                                               50%
                                                      75%
                                                             97.5%
                  mean
## (Intercept)
                  3.44
                          1.41
                                 0.88
                                        2.47
                                               3.33
                                                      4.33
                                                             6.48
                  -0.09
                          0.04 -0.17
                                       -0.11
                                             -0.08 -0.06 -0.02
## age
## femcM
                  -1.59
                          0.74 -3.13
                                       -2.08
                                              -1.57
                                                    -1.08
                                                            -0.18
                  0.44
                                                      0.51
## mean_PPD
                          0.09
                                 0.27
                                        0.38
                                               0.44
                                                             0.62
## log-posterior -32.54
                         1.35 -35.91 -33.11 -32.20 -31.60 -31.06
##
## Diagnostics:
```

```
mcse Rhat n_eff
##
                 0.03 1.00 3123
## (Intercept)
## age
                 0.00 1.00 2841
## femcM
                 0.01 1.00 3328
## mean_PPD
                 0.00 1.00 4033
## log-posterior 0.03 1.00 1503
## For each parameter, mcse is Monte Carlo standard error, n_eff is a crude measure of effective sample
# probability that a male more likely to die than a female of same age= P[femc < 0]
donner.post <- as.matrix(donner.stan)</pre>
mean(donner.post[,3] < 0)</pre>
## [1] 0.98725
Incomplete blocks, with random block effects
Benefit of Bayes - do not assume block variance is known exactly
ib <- read.csv('IBtest.csv')</pre>
ib.stan <- stan lmer(</pre>
   y ~ trt.f + (1 | block.f),
    data=ib,
    chains = 4,
    cores = 4
    )
## Warning: There were 4 divergent transitions after warmup. Increasing adapt_delta above 0.95 may help
## http://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## Warning: Examine the pairs() plot to diagnose sampling problems
ib.stan <- stan_lmer(</pre>
    y ~ trt.f + (1 | block.f),
    data=ib,
    chains = 4,
    cores = 4,
    adapt_delta = 0.98
    )
prior_summary(ib.stan)
## Priors for model 'ib.stan'
## -----
## Intercept (after predictors centered)
##
  ~ normal(location = 0, scale = 10)
##
        **adjusted scale = 9.07
##
## Coefficients
## ~ normal(location = 0, scale = 2.5)
##
        **adjusted scale = 2.27
##
## Auxiliary (sigma)
   ~ exponential(rate = 1)
##
        **adjusted scale = 0.91 (adjusted rate = 1/adjusted scale)
```

##

```
## Covariance
## ~ decov(reg. = 1, conc. = 1, shape = 1, scale = 1)
## See help('prior_summary.stanreg') for more details
summary(ib.stan, digits=2,
  pars=c('(Intercept)', 'trt.fb', 'sigma'),
 regex_pars='Sigma*')
##
## Model Info:
##
## function:
                 stan lmer
## family:
                 gaussian [identity]
## formula:
                 y ~ trt.f + (1 | block.f)
## algorithm:
                 sampling
## priors:
                  see help('prior_summary')
## sample:
                  4000 (posterior sample size)
## observations: 80
## groups:
                 block.f (60)
##
## Estimates:
##
                                            mean
                                                  sd
                                                         2.5%
                                                                25%
                                                                      50%
## (Intercept)
                                           0.06
                                                  0.14 -0.22 -0.04 0.06
## trt.fb
                                           0.19
                                                  0.18 -0.17
                                                               0.07 0.19
                                                  0.11 0.56
                                                               0.67 0.75
## sigma
                                           0.76
## Sigma[block.f:(Intercept),(Intercept)]
                                           0.27
                                                  0.19 0.00
                                                               0.12 0.26
                                            75%
                                                  97.5%
## (Intercept)
                                           0.16 0.34
## trt.fb
                                           0.32 0.55
## sigma
                                           0.84 0.99
## Sigma[block.f:(Intercept),(Intercept)] 0.40 0.66
## Diagnostics:
##
                                          mcse Rhat n_eff
## (Intercept)
                                          0.00 1.00 3533
                                          0.00 1.00 5222
## trt.fb
                                          0.01 1.01 382
## sigma
## Sigma[block.f:(Intercept),(Intercept)] 0.01 1.01 343
## For each parameter, mcse is Monte Carlo standard error, n_eff is a crude measure of effective sample
```

Overdispersed count data

Benefit of Bayes - do not assume known amount of overdispersion

```
pod <- read.csv('PODtest.csv')
pod.glmm <- glmer(y ~ xc + (1|obs), data=pod, family=poisson)
summary(pod.glmm)

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: poisson ( log )
## Formula: y ~ xc + (1 | obs)
## Data: pod</pre>
```

```
##
##
       ATC
                BIC logLik deviance df.resid
              349.8 -169.8
##
      345.6
                                339.6
##
## Scaled residuals:
                  1Q
                     Median
                                    3Q
       Min
                                            Max
## -1.13140 -0.04849 -0.00471 0.02156 0.12427
##
## Random effects:
## Groups Name
                       Variance Std.Dev.
           (Intercept) 2.241
                               1.497
## Number of obs: 30, groups: obs, 30
## Fixed effects:
              Estimate Std. Error z value Pr(>|z|)
                        0.2800 13.610 < 2e-16 ***
## (Intercept)
                3.8104
## xc
                 0.8154
                            0.1818 4.486 7.26e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
      (Intr)
## xc -0.040
confint(pod.glmm)
## Computing profile confidence intervals ...
                   2.5 %
                         97.5 %
## .sig01
              1.1458084 2.045047
## (Intercept) 3.2219027 4.367875
              0.4516029 1.196259
pod.stan <- stan_glmer(</pre>
    y \sim xc + (1 \mid obs),
    data=pod,
   family=poisson,
    chains = 4,
    cores = 4
summary(pod.stan, digits=2,
  pars=c('(Intercept)', 'xc'),
  regex_pars='Sigma*')
## Model Info:
##
## function:
                  stan glmer
## family:
                  poisson [log]
## formula:
                 y \sim xc + (1 \mid obs)
## algorithm:
                 sampling
                  see help('prior_summary')
## priors:
## sample:
                  4000 (posterior sample size)
## observations: 30
## groups:
                  obs (30)
##
```

```
## Estimates:
##
                                                      2.5%
                                                              25%
                                                                    50%
                                                                          75%
                                                 sd
                                          mean
## (Intercept)
                                               0.30 3.20
                                                                  3.80
                                                                       4.00
                                        3.80
                                                            3.61
                                               0.19 0.41
                                        0.80
                                                            0.67
                                                                  0.80 0.92
## xc
## Sigma[obs:(Intercept),(Intercept)] 2.55
                                               0.85 1.34
                                                            1.98
                                                                  2.41
                                                                        2.98
##
                                          97.5%
## (Intercept)
                                        4.38
## xc
                                        1.16
## Sigma[obs:(Intercept),(Intercept)] 4.62
##
## Diagnostics:
                                       mcse Rhat n_eff
##
## (Intercept)
                                        0.01 1.00 728
                                        0.01 1.00 812
## xc
## Sigma[obs:(Intercept),(Intercept)] 0.03 1.00 917
##
```

For each parameter, mcse is Monte Carlo standard error, n_eff is a crude measure of effective sample

lmer() and glmer() condition on the estimated variance components.

I.e., inference on fixed effect parameters considers those variance components to be known precisely. Bayes accounts for that uncertainty

Some models (e.g. RCBD) variance has no effect on estimates, just uncertainty

Other models (e.g. OD, IB), different variances changes the estimates - here's where Bayes matters

Final words:

the prior is an important part of the model: be critical of both from the WinBUGS reference manual: BEWARE: MCMC sampling can be dangerous

Resources:

- - Articles: Muth, Oravecz and Gabry (2018) User-friendly Bayesian regression modeling: a tutorial with rstandarm and shinystan. The Quantitative Methods for Psychology 14(2):99-119 with code at https://osf.io/ebz2f/
 - STAN project has wonderful vignettes about using rstanarm. Start with http://mc-stan.org/rstanarm/articles/rstanarm.html
 - Then look at the model-specific vignettes (vignettes tab) mixed models are in the Group Specific Terms vignette
- Bayes in Ecology: now lots of great books
 - Barker and Link: Bayesian Inference (my favorite)
 - McCarthy: Bayesian Methods for Ecology
 - King et al.: Bayesian Analysis for Population Ecology
 - Korner-Niervergelt et al.: Bayesian Analysis in Ecology using linear models with R, BUGS and
 - Parent and Rivot: Introduction to Hierarchical Bayesian Modeling for Ecological Data