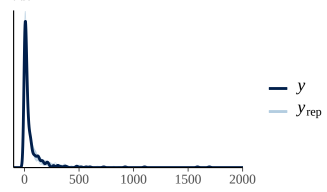
Model 1

brm(visits ~ autonomous_selfing_level + (1|net_id) + (1|gr(phylo, cov = A))

This model uses both quantitative and qualitative data of selfing levels which is grouped in 4 categories: "high", "medium", "low" and "none". The response variable is the integer of visits. Note that the quantitative measurement that we use is autonomous fruit production and is grouped as follows: 0=none, >0-25=low, >25-75=medium; >75-100=high.

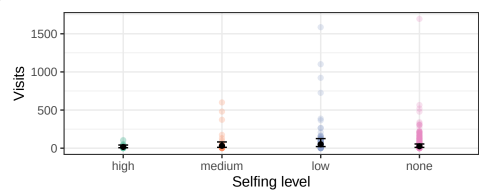
Check now the fit of the model:



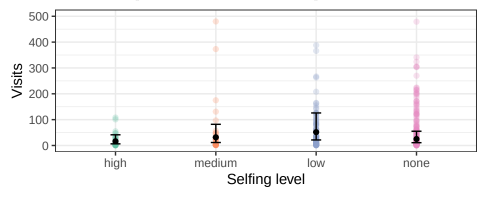
The fit seems right, check now estimates, se and credible intervals:

autonomous_selfing_level	estimate	se	lower	upper
high	16.22890	7.237954	6.388558	41.40229
medium	31.81053	15.517126	11.542339	81.92444
low	51.62325	21.704897	21.417291	125.88574
none	25.07522	9.871306	10.825766	55.17484

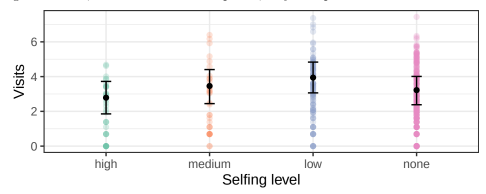
Plot output:



Set ylim max to 500 in order to improve visualization Some data points not shown.



Try now to logit estimates, credible intervals and points, maybe improves visualization?



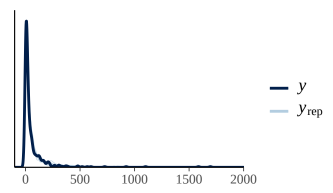
So far we haven't found big differences between groups. Also note that we have a greater number of low and none self species. This could be discussed. Keep in mind.

Now I'm going to exclude the dioecious and monoecioues species that were considered as non-selfers.

Model 2 Excluding species with flowers with separated sexes

brm(visits ~ autonomous_selfing_level + (1|net_id) + (1|gr(phylo, cov = A))

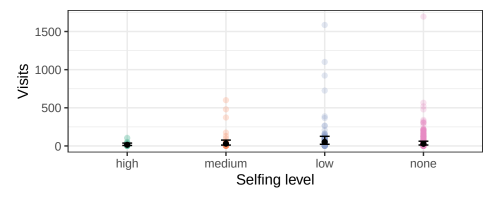
Check now the fit of the model:



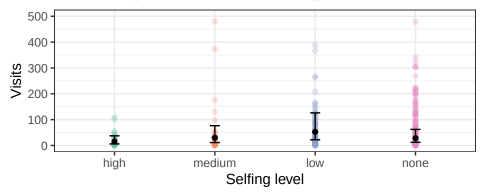
Again, the fit seems right, check now estimates, se and credible intervals:

autonomous_selfing_level	estimate	se	lower	upper	
high	14.99806	6.823373	6.126669	37.44073 76.56629	
medium	30.11032	14.079376	11.333166		
low	52.28513	21.083391	22.036199	126.20680	
none	28.43006	10.913705	12.308011	62.22853	

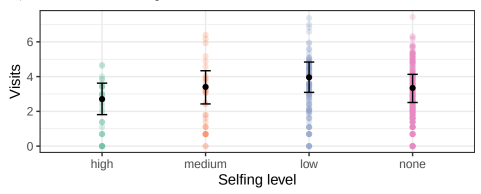
Plot output:



Set ylim max to 500 in order to improve visualization Some data points not shown.



Logit estimates, credible intervals and points



Including or excluding dioecious and monoecious species does not make a very big differences in the models.

BAYES R2 Model 1 including all species $\frac{1}{2}$

```
## R2 0.296992 0.07886342 0.1680863 0.4678135
```

##		autonomous	s_selfing_l	evel	visits	net_id	phylo	cond	effect1	estimate
##	1			high	65.28024	NA	NA	1	high	14.99806
##	2		me	dium	65.28024	NA	NA	1	medium	30.11032
##	3			low	65.28024	NA	NA	1	low	52.28513
##	4			none	65.28024	NA	NA	1	none	28.43006
##		se	lower	upp	per					
##	1	6.823373	6.126669	37.4	14073					
##	2	14.079376	11.333166	76.5	56629					

```
## 3 21.083391 22.036198 126.20680
## 4 10.913705 12.308011 62.22853
```

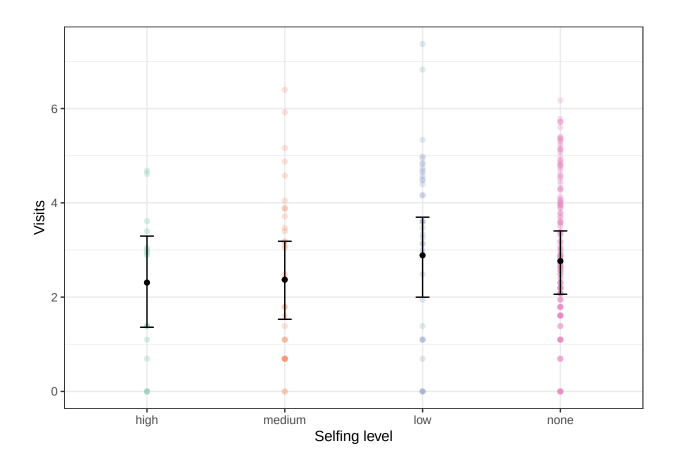
BAYES R2 Model 1 without monoecious and dioecious species

```
##
      Estimate Est.Error
                                02.5
                                         Q97.5
## R2 0.246085 0.07870143 0.1224886 0.4225047
##
     autonomous_selfing_level
                               visits net_id phylo cond__ effect1__ estimate__
## 1
                         high 65.28024
                                            NA
                                                   NA
                                                           1
                                                                  high
                                                                         14.99806
## 2
                       medium 65.28024
                                            NA
                                                  NA
                                                           1
                                                                medium
                                                                         30.11032
## 3
                           low 65.28024
                                                           1
                                                                         52.28513
                                            NA
                                                   NA
                                                                   low
                         none 65.28024
## 4
                                            NA
                                                   NA
                                                           1
                                                                  none
                                                                         28.43006
##
                 lower__
          se__
                            upper__
     6.823373 6.126669
                          37.44073
## 2 14.079376 11.333166 76.56629
## 3 21.083391 22.036198 126.20680
## 4 10.913705 12.308011 62.22853
```

Because the qualitative data can add some noise to the analysis now I'm going to exclude it and just consider species that have quantitative information of the auntonomous selfing level.

MODEL 3 JUST QUANTITATIVE DATA BUT GROUPED IN 4 CATEGORIES NONE, LOW, MEDIUM, HIGH.

Two distributions fit the data: **Negative binomial** and **skew gaussian** with logit transformation of visits and because for the negative binomial I have to to log transform the data in order to improve visualization. I'm going to keep from now on with a **skew gaussian** with the logit of visits (the output is quite similar).



```
##
     Estimate Est.Error
                          Q2.5
                                 Q97.5
## R2 0.3663574 0.07309651 0.2143842 0.499171
    autonomous_selfing_level log(visits) visits net_id phylo cond__ effect1__
##
## 1
                           4.106142 60.71206
                                             NA
                                                  NA
                    high
                                                         1
                                                              high
                          4.106142 60.71206
## 2
                  medium
                                             NA
                                                  NA
                                                             medium
## 3
                     low
                           4.106142 60.71206
                                             NA
                                                  NA
                                                               low
                                                         1
## 4
                    none
                           4.106142 60.71206
                                             NA
                                                  NA
                                                              none
##
    estimate__
                 se__ lower__ upper__
## 1
     2.308604 0.4979512 1.361141 3.294633
    2.369555 0.3962446 1.530127 3.184184
## 2
```