# Summarizing the Posterior

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June 17, 2021

## Summarizing the Posterior

- Once our Bayesian model produces a posterior distribution, it is necessary to summarize and interpret the posterior distribution.
- A posterior distribution is (usually) a high dimensional object that is hard to visualize and work with [Murphy, 2021].
- In this class we will learn how to draw estimates (e.g., point estimates, intervals) to summarize and interpret a posterior distribution.
- Exactly how it is summarized depends upon our purpose.
- Common guestions include:
  - How much posterior probability lies below some parameter value?
  - How much posterior probability lies between two parameter values?
  - Which parameter value marks the lower 5% of the posterior probability?
  - Which range of parameter values contains 90% of the posterior probability?
  - Which parameter value has highest posterior probability?

### Sampling to summarize

- These questions can be usefully divided into questions about:
  - intervals of defined boundaries
  - intervals of defined probability mass
  - point estimates
- In the theoretical world (when the posterior has a closed mathematical expressions), answering these questions implies calculating complicated integrals to cancel out (or average) different variables.
- In the practical world, however, the same results can be approximated using samples from the posterior.
- In this class we will approach the above questions using samples from the posterior.
- Another reason to learn to work with posterior samples is that methods like MCMC produce nothing but samples from the posterior.
- This class is based on Chapter 3 of [McElreath, 2020].

### Sampling from a grid-approximate posterior

- Before beginning to work with samples, we need to generate them.
- Here's a reminder for how to compute the posterior for the globe tossing model, using grid approximation:

```
p_grid <- seq( from=0 , to=1 , length.out=1000 )
prior <- rep( 1 , 1000 )
likelihood <- dbinom( 6 , size=9 , prob=p_grid )
posterior <- likelihood * prior
posterior <- posterior / sum(posterior)</pre>
```

- Now we wish to draw 10,000 samples from this posterior.
- Imagine the posterior is a bucket full of parameter values, numbers such as 0.1, 0.7, 0.5, 1, etc.
- Within the bucket, each value exists in proportion to its posterior probability, such that values near the peak are much more common than those in the tails.

## Sampling from a grid-approximate posterior

- We're going to scoop out 10,000 values from the bucket.
- Provided the bucket is well mixed, the resulting samples will have the same proportions as the exact posterior density.
- Therefore the individual values of *p* will appear in our samples in proportion to the posterior plausibility of each value.
- Here's how you can do this in R, with one line of code:

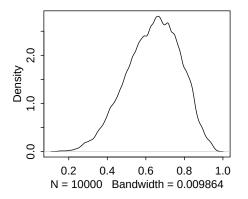
```
samples <- sample( p_grid , prob=posterior , size=1e4 ,
replace=TRUE )</pre>
```

 We are randomly pulling values from the grid of parameter values where the probability of each value is given by the posterior.

## Sampling from a grid-approximate posterior

We can visualize a density plot of our posterior sample as follows:

```
library(rethinking)
dens(samples)
```



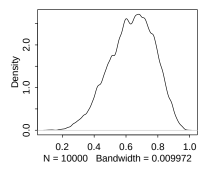
 We can see that the estimated density is very similar to to ideal posterior we computed via grid approximation in previous class.

### Sampling from the theoretical posterior

 We could get the same results by sampling from the theoretical posterior using the beta distribution:

```
teo.samples<-rbeta(1e4,7,4)
dens(teo.samples)</pre>
```

• We can see that the estimated density is very similar to the theoretical posterior obtained from the beta distribution:



 However, we should keep in mind that for complex models we will not have access to the posterior closed form, so it is better to get used to working with samples.

#### Intervals of defined boundaries

- Suppose I ask you for the posterior probability that the proportion of water is less than 0.5.
- We could calculate this from the theoretical posterior:

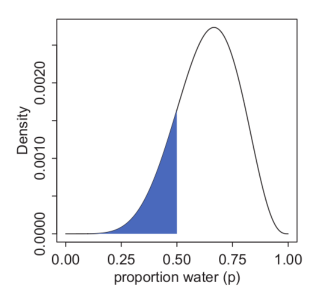
```
> pbeta(0.5,7,4)
[1] 0.171875
```

 Or alternatively we could calculate it from the grid-approximate posterior by adding up all of the probabilities where the corresponding parameter value is less than 0.5.

```
> sum( posterior[ p_grid < 0.5 ] )
[1] 0.1718746</pre>
```

So about 17% of the posterior probability is below 0.5.

#### Intervals of defined boundaries



#### Intervals of defined boundaries

- Now, let's perform the same calculation, using samples from the posterior.
- Recall than in more complex models neither a grid-approximation nor a closed-form posterior will be available.
- All we have to do is add up all samples less than 0.5 and divide the resulting count by the total number of samples.

```
> sum( samples < 0.5 ) / 1e4
[1] 0.1752</pre>
```

#### Conclusions

Blabla

#### References I



McElreath, R. (2020).

Statistical rethinking: A Bayesian course with examples in R and Stan. CRC press.



Murphy, K. P. (2021).

Probabilistic Machine Learning: An introduction. MIT Press.