# STA 360/602L: MODULE 3.1

#### MONTE CARLO APPROXIMATION AND SAMPLING

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- Monte Carlo integration is very key for Bayesian computation and using simulations in general.
- While we will focus on using Monte Carlo integration for Bayesian inference, the development is general and applies to any pdf/pmf  $p(\theta)$ .
- For our purposes, we will want to evaluate expectations of the form

$$
H=\int h(\theta)\cdot p(\theta)d\theta,
$$

for many different functions  $h(.)$  (usually scalar for us).



- **Procedure:** 
	- 1. Generate a random sample  $\theta_1, \ldots, \theta_m \stackrel{ind}{\sim} p(\theta)$ .
	- 2. Estimate  $H$  using

$$
\bar{h}=\frac{1}{m}\sum_{i=1}^m h(\theta_i).
$$

In this course,  $p(\theta)$  would often be the posterior distribution  $\pi(\theta | y).$ 



- We have  $\mathbb{E}[h(\theta_i)]=H.$
- Assuming  $\mathbb{E}[h^2(\theta_i)] < \infty$ , so that the variance of each  $h(\theta_i)$  is finite, we have
	- 1. LLN:  $\bar{h} \overset{a.s.}{\rightarrow} H$ .
	- 2. CLT:  $\bar{h}-H$  is is asymptotically normal, with asymptotic variance

$$
\frac{1}{m}\int (h(\theta)-H)^2 p(\theta)d\theta,
$$

which can be approximated by

$$
v_m=\frac{1}{m^2}\sum_{i=1}^m(h(\theta_i)-\bar{h})^2.
$$

 $\sqrt{v_m}$  is often called the Monte Carlo standard error.



- That is, generally, taking large Monte Carlo sample sizes  $m$  (in the thousands or tens of thousands) can yield very precise, and cheaply computed, numerical approximations to mathematically difficult integrals.
- What this means for us: we can approximate just about any aspect of the posterior distribution with a large enough Monte Carlo sample.



For samples  $\theta_1, \ldots, \theta_m$  drawn iid from  $\pi(\theta | y)$ , as  $m \to \infty$ , we have

$$
\begin{aligned}\n\overline{\theta} &= \frac{1}{m} \sum_{i=1}^{m} \theta_i \to \mathbb{E}[\theta|y] \\
\overline{\theta} &= \frac{1}{m-1} \sum_{i=1}^{m} (\theta_i - \overline{\theta})^2 \to \mathbb{V}[\theta|y] \\
\overline{\theta} &= \frac{1}{m} \sum_{i=1}^{m} 1[\theta_i \le c] = \frac{\# \theta_i \le c}{m} \to \Pr[\theta \le c|y] \\
\overline{\theta} &= \frac{\alpha}{m} \quad \text{if } \theta < c \end{aligned}
$$

 $\rightarrow 100 \times (1 - \alpha)$  quantile-based credible interval.  $\left[\frac{\alpha}{2}$ th percentile of  $(\theta_1,\ldots,\theta_m)$ ,  $(1-\frac{\alpha}{2})$ th percentile of  $(\theta_1,\ldots,\theta_m)$ ] 2 2



#### BACK TO BIRTH RATES

- Suppose we randomly sample two "new" women, one with degree and one without.
- To what extent do we expect the one without the degree to have more To what extent do we expect the one without the degree to hard kids than the other, e.g.  $\tilde{y}_1 > \tilde{y}_2 | y_{11}, \ldots, y_{1n_1}, y_{21}, \ldots, y_{2n_2}$ ?
- **Using R,**

```
set.seed(01222020)
a \leftarrow 2; b \leftarrow 1; #prior
n1 <- 111; sumy1 <- 217; n2 <- 44; sumy2 <- 66 #data
y1<sub>pred</sub> \leftarrow rnbinom(100000,size=(a+sumy1),mu=(a+sumy1)/(b+n1))
y2 pred <- rnbinom(10000,size=(a+sumy2),mu=(a+sumy2)/(b+n2))
mean(v1 pred > v2 pred)
```
## [1] 0.48218

 $mean(y1$ <sub>pred</sub> ==  $y2$ <sub>pred</sub>)

## [1] 0.21842



#### BACK TO BIRTH RATES

- That is,  $\Pr({{\tilde y}_1} > {{\tilde y}_2}|y_{11}, \ldots ,y_{1n_1}, y_{21}, \ldots ,y_{2n_2}) \approx 0.48$  and  $Pr(\tilde{y}_1 = \tilde{y}_2 | y_{11}, \ldots, y_{1n_1}, y_{21}, \ldots, y_{2n_2}) \approx 0.22.$
- Notice that strong evidence of difference between two populations does not really imply the difference in predictions is large.



- This general idea of using samples to "approximate" averages (expectations) is also useful when trying to approximate posterior predictive distributions.
- Quite often, we are able to sample from  $p(y_i|\theta)$  and  $\pi(\theta|\{y_i\})$  but not from  $p(y_{n+1}|y_{1:n})$  directly.
- We can do so indirectly using the following Monte Carlo procedure:

$$
\begin{aligned} \text{sample } \theta^{(1)} &\sim \pi(\theta|\{y_i\}), \ \ \text{then sample } y_{n+1}^{(1)} \sim f(y_{n+1}|\theta^{(1)}) \\ \text{sample } \theta^{(2)} &\sim \pi(\theta|\{y_i\}), \ \ \text{then sample } y_{n+1}^{(2)} \sim f(y_{n+1}|\theta^{(2)}) \\ &\vdots \\ \text{sample } \theta^{(m)} &\sim \pi(\theta|\{y_i\}), \ \ \text{then sample } y_{n+1}^{(m)} \sim f(y_{n+1}|\theta^{(m)}). \end{aligned}
$$

- The sequence  $\{(\theta,y_{n+1})^{(1)},\ldots,(\theta,y_{n+1})^{(m)}\}$  constitutes : independent samples from the joint posterior of  $(\theta, Y_{n+1})$ .  $,\ldots, (\theta, y_{n+1})^{(m)}\}$  constitutes  $m$
- In fact,  $\{y_{n+1}^{(1)},\ldots,y_{n+1}^{(m)}\}$  are independent draws from the posterior predictive distribution we care about.  $, \ldots, y$ <sub>1</sub>  $\binom{m}{n+1}$



#### BACK TO BIRTH RATES

Let's re-compute  $\Pr({{\tilde y}_1} > {{\tilde y}_2}|{y_{11}}, \ldots ,{y_{1n_1}},y_{21}, \ldots ,{y_{2n_2}})$  and Pr $(\tilde{y}_1 = \tilde{y}_2 | y_{11}, \ldots, y_{1n_1}, y_{21}, \ldots, y_{2n_2})$  using this method.

**Using R,** 

```
set.seed(01222020)
a \leftarrow 2; b \leftarrow 1; #prior
n1 <- 111; sumy1 <- 217; n2 <- 44; sumy2 <- 66 #data
theta1 pred \leq rgamma(10000,219,112); theta2 pred \leq rgamma(10000,68,45)
y1_pred <- rpois(10000,theta1_pred); y2_pred <- rpois(10000,theta2_pred)
mean(v1 pred > v2 pred)
```
## [1] 0.4765

mean( $v1$  pred ==  $v2$  pred)

## [1] 0.2167

Again,  $\Pr({{\tilde y}_1} > {{\tilde y}_2}|y_{11}, \ldots ,y_{1n_1}, y_{21}, \ldots ,y_{2n_2}) \approx 0.48$  and  $\Pr({\tilde y}_1={\tilde y}_2|y_{11},\ldots,y_{1n_1},y_{21},\ldots,y_{2n_2}) \approx 0.22.$ 



#### WHAT' S NEXT?

MOVE ON TO THE READINGS FOR THE NEXT MODULE!



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