Bootstrapping & Resampling Methods

general problem

- scientific Qs are about populations
- we can't measure entire populations
- experiments generate samples
- samples -> estimate population parameters
- "parametric" approaches come with assumptions

general problem

- what if assumptions are violated?
- data are not normally distributed
- variances unequal
- sample size unequal
- nonlinear model
- etc etc

bootstrapping

- I. a way to estimate the precision of sample-based population estimates (without having access to the entire population)
 - doesn't rely on parametric assumptions (e.g. normality)
- 2. a way to do hypothesis testing
 - non-parametric, by simulating the null
- 3. a way to do power calculations
 - not restricted by assumptions

- we saw earlier:
 - best estimate of a population mean is the sample mean (assuming $\hat{\mu} = \bar{X} = \frac{\sum X_i}{N}$ normality)
 - estimate of sd of sampling distribution of means is standard error of mean:

$$s_{\bar{x}} = \frac{s_x}{\sqrt{N}}$$

 \bullet can use this to generate 95% Cls of $~\bar{\chi} \pm t_{\alpha}(s_{\bar{x}})$ population mean

- bootstrapping can estimate sampling distribution of means
- no need to assume any particular theoretical distribution
- use resampling with replacement to simulate repeatedly sampling from the population
- uses sample as proxy for population

assume you have a sample X1...Xn and a statistic of interest (e.g. the mean) repeat M times (where M is large, e.g. 10,000)

generate a new sample of size n by resampling, with replacement, from X1..Xn compute the statistic based on the new sample set that statistic aside (e.g. save it in a list)

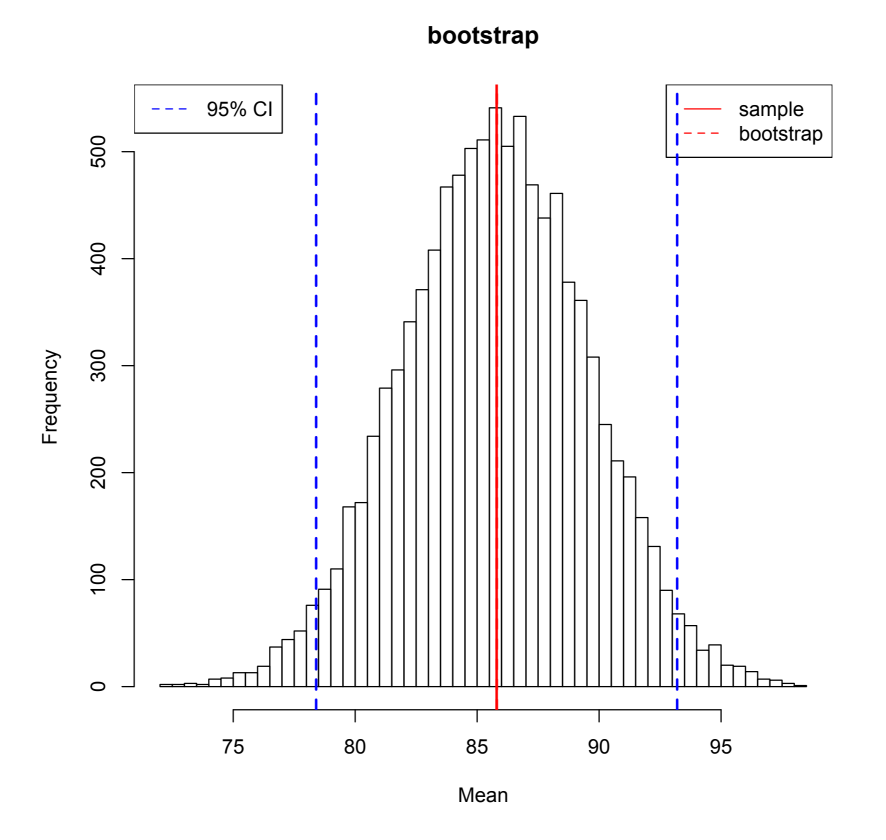
now you have a list of M versions of the statistic, one for each resampling

that list represents an empirical bootstrap distribution of the statistic of interest

now you can compute relevant quantities of that distribution (e.g. 95% Cls)

- e.g. we have a sample of size 20:
- 66 79 93 86 69 79 101 97 91 95
 72 106 105 75 70 85 92 74 88 93
- estimate of population mean (using sample mean) is 85.8
- how precise is that estimate?

```
X = c(66, 79, 93, 86, 69, 79, 101, 97, 91, 95, 72, 106, 105, 75, 70, 85, 92, 74, 88, 93)
# compute a statistic of interest
(Xm = mean(X))
# use resampling to generate an empirical bootstrap distribution of that statistic
# how many simulated experiments?
boot_m = 1000000
# create a list to store our bootstrap values
Xm\_boot = array(NA, boot\_m)
# do it
for (i in 1:10000) {
   Xb = sample(X, length(X), replace=TRUE) # generate new sample
   Xm\_boot[i] = mean(Xb)
                                   # compute statistic of interest
# display results
hist(Xm_boot, xlab="Mean", main="bootstrap")
abline(v=Xm, col="red")
abline(v=mean(Xm_boot), col="red", lty=2)
legend(x="topright", lty=c(1,2), col=c("red","red"), legend=c("sample","bootstrap"))
# compute 95% CI
(CI95 = quantile(Xm_boot, probs=c(.025,.975)))
abline(v=CI95[1], lty=2, col="blue")
abline(v=CI95[2], lty=2, col="blue")
legend(x="topleft", lty=2, col="blue", legend="95% CI")
```



- here we used a bootstrap to estimate the sampling distribution of the mean
- we can do the same procedure to estimate the sampling distribution of any statistic we want
- e.g. variance, or median, or skew, ...
- or anything we make up
- bootstrapping will estimate sampling distribution

- example: comparing two populations
- drug vs control
- null hypothesis: drug has no effect
 - drug & control sampled from same population
- alternate hypothesis: drug has an effect
 - drug & control not sampled from same population

- choose a test statistic (e.g. the difference between means... but could be anything; t, F, sd, whatever your scientific question calls for)
- do many many times (e.g. 10,000):

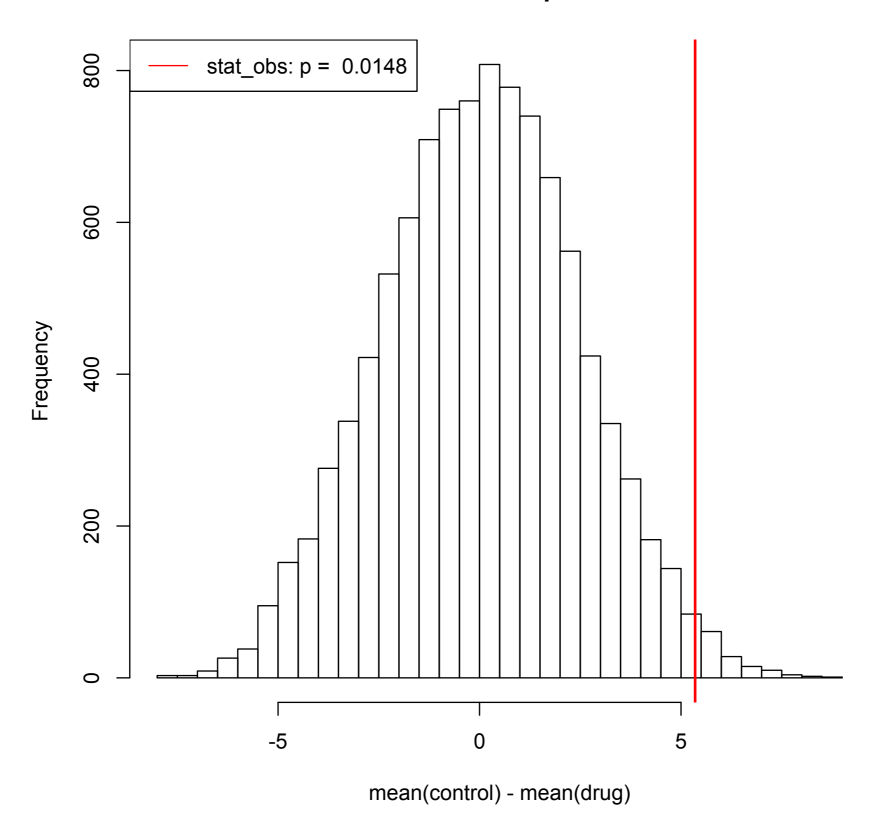
 simulate the null hypothesis (that drug & control are sampled from same population)

 how many times did you get a test statistic as large or larger as the original one? < 5%? then reject H0

- choose a test statistic (e.g. the difference between means... but could be anything; t, F, sd, whatever your scientific question calls for)
- do many many times (e.g. 10,000):
 - throw both groups into a bucket
 - randomly reconstitute the two groups, disregarding their original group membership
 - recompute the statistic of interest
- how many times did you get a test statistic as large or larger as the original one? < 5%? then reject H0

```
# our control group
q_{control} \leftarrow c(87,90,82,77,71,81,77,79,84,86,78,84,86,69,81,75,70,76,75,93)
# our drug group
q_druq < c(74,67,81,61,64,75,81,81,81,67,72,78,83,85,56,78,77,80,79,74)
# our statistic of interest here is the difference between means
(stat_obs <- mean(q_control) - mean(q_drug))</pre>
# how many simulated experiments?
n\_boot = 10000
# create a list to store our bootstrap values
stat_boot = array(NA, n_boot)
# now do a bootstrap to simulate the null hypothesis,
# namely that both groups were sampled from the same population
n_c = length(q_control)
n_d = length(g_drug)
g_bucket = c(q_control, q_drug)
for (i in 1:n_boot) {
    # reconstitute both groups, ignoring original labels
    permuted_order <- sample(1:(n_c+n_d), n_c+n_d, replace=FALSE)</pre>
    permuted_bucket <- q_bucket[permuted_order]</pre>
    boot_control <- permuted_bucket[1:n_c]</pre>
    boot_drug <- permuted_bucket[(n_c+1):(n_c+n_d)]</pre>
    stat_boot[i] <- mean(boot_control) - mean(boot_drug)</pre>
}
# visualize the empirical bootstrap distribution of our statistic of interest
hist(stat_boot, xlab="mean(control) - mean(drug)", main="bootstrap")
abline(v=stat_obs, col="red")
# how many times in the bootstrap did we observe a stat_boot as big or bigger than our stat_obs?
(p_boot <- length(which(stat_boot >= stat_obs)) / n_boot)
legend(x="topleft", lty=1, col="red", legend=paste("stat_obs: p = ", p_boot))
```

bootstrap



- here we tested the difference between means
- but we can apply this method to any statistic of interest that we can calculate
- no need to assume theoretical distribution
- compute probability under H0 empirically by simulating the null hypothesis

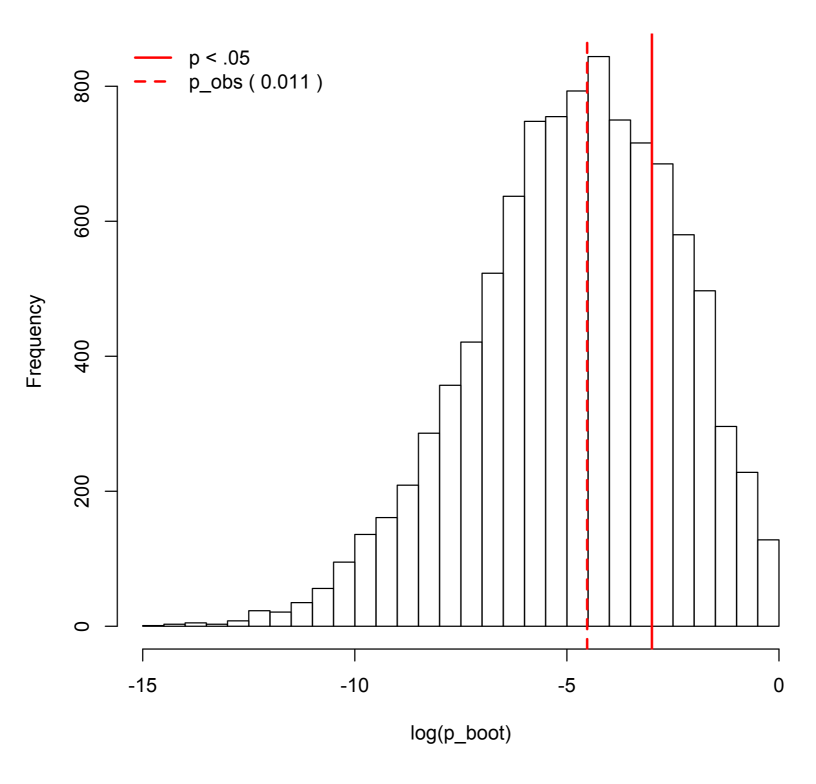
- we can use random resampling to simulate experiments not only under the null hypothesis but under any alternate hypothesis of our choosing
- we can use simulations to answer questions about statistical power

- what's the probability of detecting a given effect with a given number of subjects?
- how many subjects are required to detect a given effect 80% of the time? (or any other % of your choosing)
- again a bootstrapping/resampling approach doesn't require assumptions about a theoretical distribution

- example: 2 groups, drug and control
- control
 87 90 82 87 71 81 77 79 84 86
 78 84 86 69 81 75 70 76 75 93
- drug
 74 73 81 65 64 75 76 81 81 67
 72 78 83 75 66 78 77 80 79 74
- Mann-Whitney U test:
 t = 2.0613 what is our statistical power?
 p = 0.04626

```
# our two groups
g_control <- c(87,90,82,87,71,81,77,79,84,86,78,84,86,69,81,75,70,76,75,93)</pre>
q_druq \leftarrow c(74,73,81,65,64,72,76,81,81,67,72,78,83,75,66,78,77,80,79,74)
# do a Mann-Whitney U test (nonparametric version of a t-test)
out <- wilcox.test(q_control, q_drug)</pre>
w_obs <- out$statistic</pre>
p_obs <- out$p.value</pre>
n boot <- 10000
w_{boot} = array(NA, n_{boot})
p_boot = array(NA, n_boot)
for (i in 1:n_boot) {
    b_control <- sample(q_control,length(q_control),replace=TRUE)</pre>
    b_drug <- sample(q_drug,length(q_drug),replace=TRUE)</pre>
    out <- wilcox.test(b_control, b_drug)</pre>
    w_boot[i] <- out$statistic
    p_boot[i] <- out$p.value</pre>
}
(power <- length(which(p_boot <= .05)) / n_boot)</pre>
hist(log(p_boot), 100, main=paste("p_boot, power=", power), xlab="p_boot")
abline(v=log(0.05), col="red", lty=1, lwd=2)
abline(v=log(p_obs), col="red", lty=2, lwd=2)
legend(x="topleft", col="red", lty=c(1,2), lwd=2, legend=c("p < .05", paste("p_obs (",round(p_obs,3),")")), box.lty=0)
```

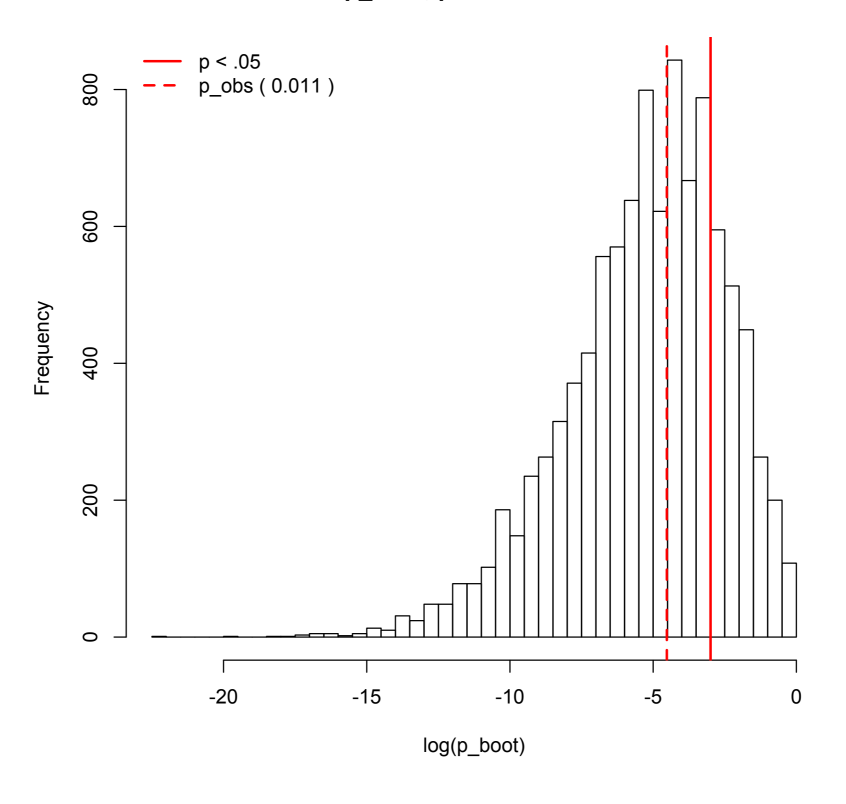
p_boot, **power= 0.7589**



- here we used bootstrap to simulate re-doing an experiment many times
- we used a Mann-Whitney U test as our statistical test
- but one could use anything (e.g. a t-test)
- If you are OK with assuming a theoretical distribution (e.g. a t distribution) then you can perform a parametric bootstrap

```
n_control <- length(q_control)</pre>
m_control <- mean(g_control)</pre>
sd_control <- sd(g_control)</pre>
n_drug <- length(g_drug)</pre>
m_drug <- mean(g_drug)</pre>
sd_drug <- sd(g_drug)</pre>
for (i in 1:n_boot){
    b_control <- rnorm(n_control, mean=m_control, sd=sd_control)</pre>
    b_drug <- rnorm(n_drug, mean=m_drug, sd=sd_drug)</pre>
    out <- wilcox.test(b_control, b_drug)</pre>
    w_boot[i] <- out$statistic
    p_boot[i] <- out$p.value</pre>
(power <- length(which(p_boot <= .05)) / n_boot)</pre>
hist(log(p_boot), 50, main=paste("p_boot, power=", power), xlab="log(p_boot)")
abline(v=log(0.05), col="red", lty=1, lwd=2)
abline(v=log(p_obs), col="red", lty=2, lwd=2)
legend(x="topleft", col="red", lty=c(1,2), lwd=2, legend=c("p < .05", paste("p_obs (",round(p_obs,3),")")), box.lty=0)
```

p_boot, **power= 0.7872**



- in a parametric bootstrap instead of simulating the experiment by resampling from your sample,
- instead you sample from the best estimate of the population distribution
- e.g. for the previous example, if we're ok to assume a normal distribution, then
- control: Normal(mean=80.55, sd=6.70)
 drug: Normal(mean=74.8, sd=5.74)

non-parametric statistical tests

- unpaired t-test: Mann-Whitney U test
- paired t-test: Wilcoxon test
- one factor ANOVA: Kruskal-Wallis test
- correlation: Spearman rank-order correlation
- etc etc