Maximum Likelihood Estimation

MLE

- tool for parameter estimation
- good approach for cases when OLS (ordinary least squares) assumptions are violated
- e.g. for non-linear models with non-normal data
- in MLE, we estimate the parameters of a model that maximize the likelihood of your data

- assume an observed data vector
 y = (y I, y2, ..., ym)
- goal of MLE is to identify the population (the model) that is **most likely** to have generated the data

- Here we assume population (model) is associated with a corresponding probability distribution
- Each probability distribution is characterized by a unique value of the model's parameter(s)

- As model parameters change, different probability distributions are generated
- Model = the family of probability distributions indexed by the model's parameter(s)

- f(y|w) is the probability density function (PDF) specifying the probability of observing data y, given model parameter(s) w
- note: w may be a parameter vectorw = (w1, w2, ..., wk)
- e.g. for a normal PDF: w = (mu, sigma)

• If observations yi are statistically independent, then by probability theory, the PDF for the data as a whole, y = (y1, ..., ym) given the parameter vector w, can be expressed as the multiplication of PDFs for individual observations:

$$f(y = (y_1, y_2, \dots, y_n)|w) = f_1(y_1|w)f_2(y_2|w)\dots f_n(y_n|w)$$

- e.g. let's say our data vector Y is made up of 3 observations y1=80, y2=110, y3=130
- and we want to compute the PDF for a normal distribution

$$p(y_i|\mu,\sigma) = \frac{1}{\sigma\sqrt{2\pi}}e^{-\frac{(y_i-\mu)^2}{2\sigma^2}}$$

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$$p(y = (y_1, y_2, y_3)|\mu, \sigma) = p(y_1|\mu, \sigma)p(y_2|\mu, \sigma)p(y_3|\mu, \sigma)$$

assume our mu=100 and sigma=15

$$p(80|\mu = 100, \sigma = 15) = \frac{1}{\sigma\sqrt{2\pi}}e^{-\frac{(80-\mu)^2}{2\sigma^2}} = 0.010934$$

$$p(110|\mu = 100, \sigma = 15) = \frac{1}{\sigma\sqrt{2\pi}}e^{-\frac{(80-\mu)^2}{2\sigma^2}} = 0.021297$$

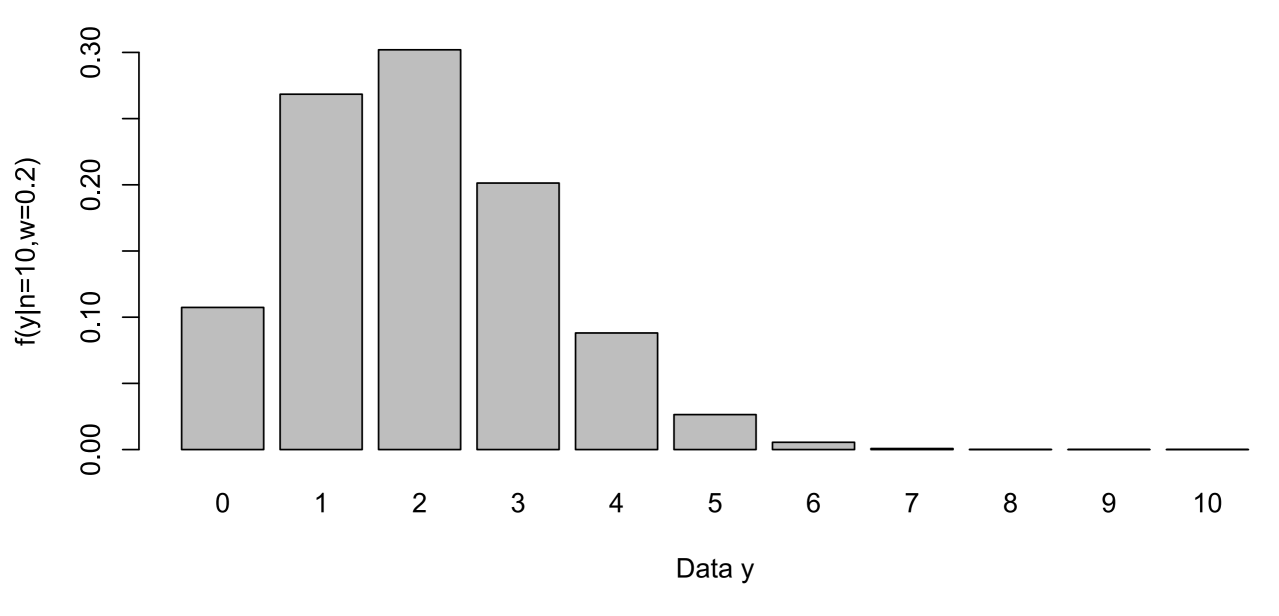
$$p(130|\mu = 100, \sigma = 15) = \frac{1}{\sigma\sqrt{2\pi}}e^{-\frac{(80-\mu)^2}{2\sigma^2}} = 0.003599$$

$$p(y = (y_1, y_2, y_3)|\mu, \sigma) = (.010934)(.021297)(.003599) = .000000838$$

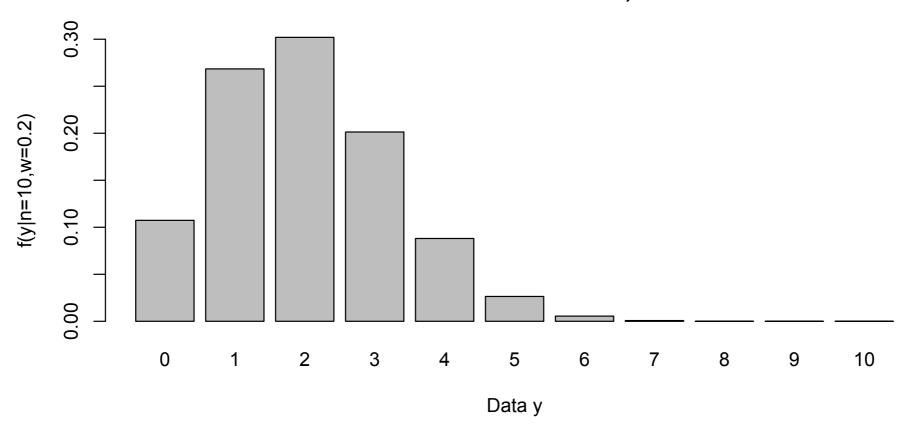
PDF: an example

- y is # of successes in a sequence of 10
 Bernoulli trials* (e.g. tossing a coin 10 x)
- assume probability of a success on any one trial is 0.2 (a biased coin)
- parameter vector w is n=10, w=0.2
- PDF is: $f(y|n = 10, w = 0.2) = \frac{10!}{y!(10-y)!}(0.2)^y(0.8)^{10-y}$ (y = 0, 1, ..., 10)
- this is binomial distribution with n=10, w=0.2

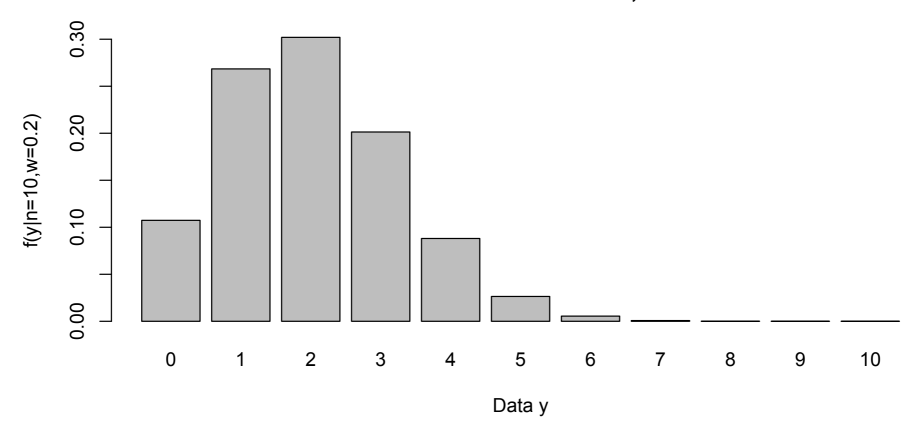
PDF for binomial with n=10, w=0.2



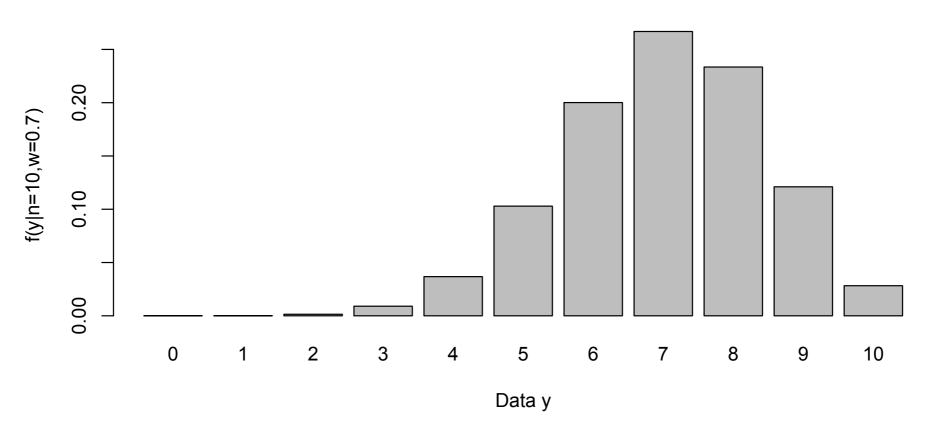
PDF for binomial with n=10, w=0.2

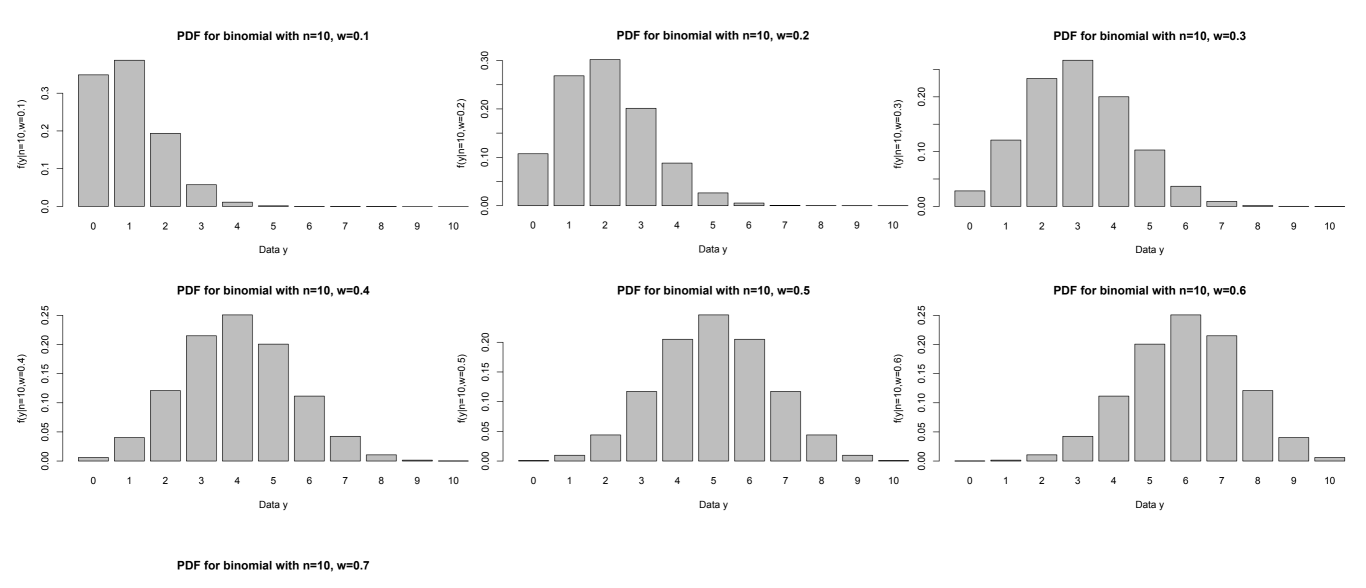


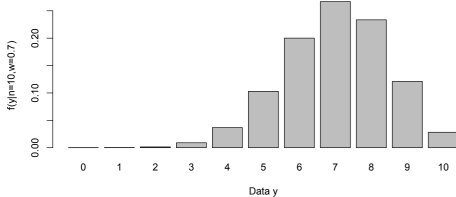
PDF for binomial with n=10, w=0.2



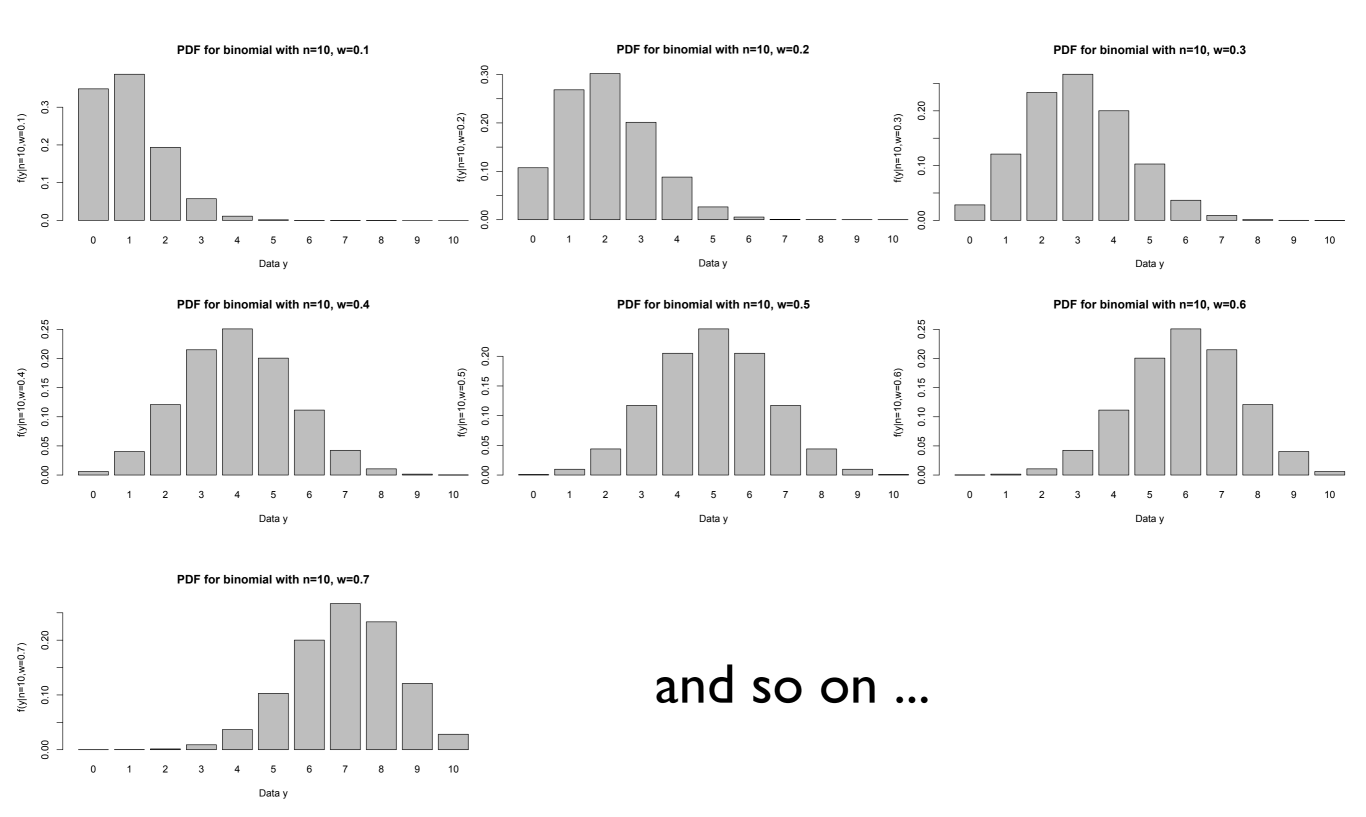
PDF for binomial with n=10, w=0.7







and so on ...



 The collection of all such PDFs generated by varying the parameter across its range defines a **model**

- Given a set of parameter values, the corresponding PDF will show that some data are more probable than other data
- In fact we have already observed the data

- We are faced with the inverse problem
- Given the observed data, and a model of the process by which the data was generated,

find the **one PDF**, among all the probability densities that the model prescribes, that is **most likely to have produced the data**

 we define the likelihood function by reversing the roles of the data vector y and the parameter vector w in f(y|w):

$$L(w|y) = f(y|w)$$

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- L(w|y) represents the likelihood of the parameter w given the observed data y
- For our one-dimensional binomial example the likelihood function for y=7 and n=10 is

$$L(w|n = 10, y = 7) = f(y = 7|n = 10, w)$$

$$= \frac{10!}{7!3!} w^7 (1 - w)^3 \quad (0 \le w \le 1)$$

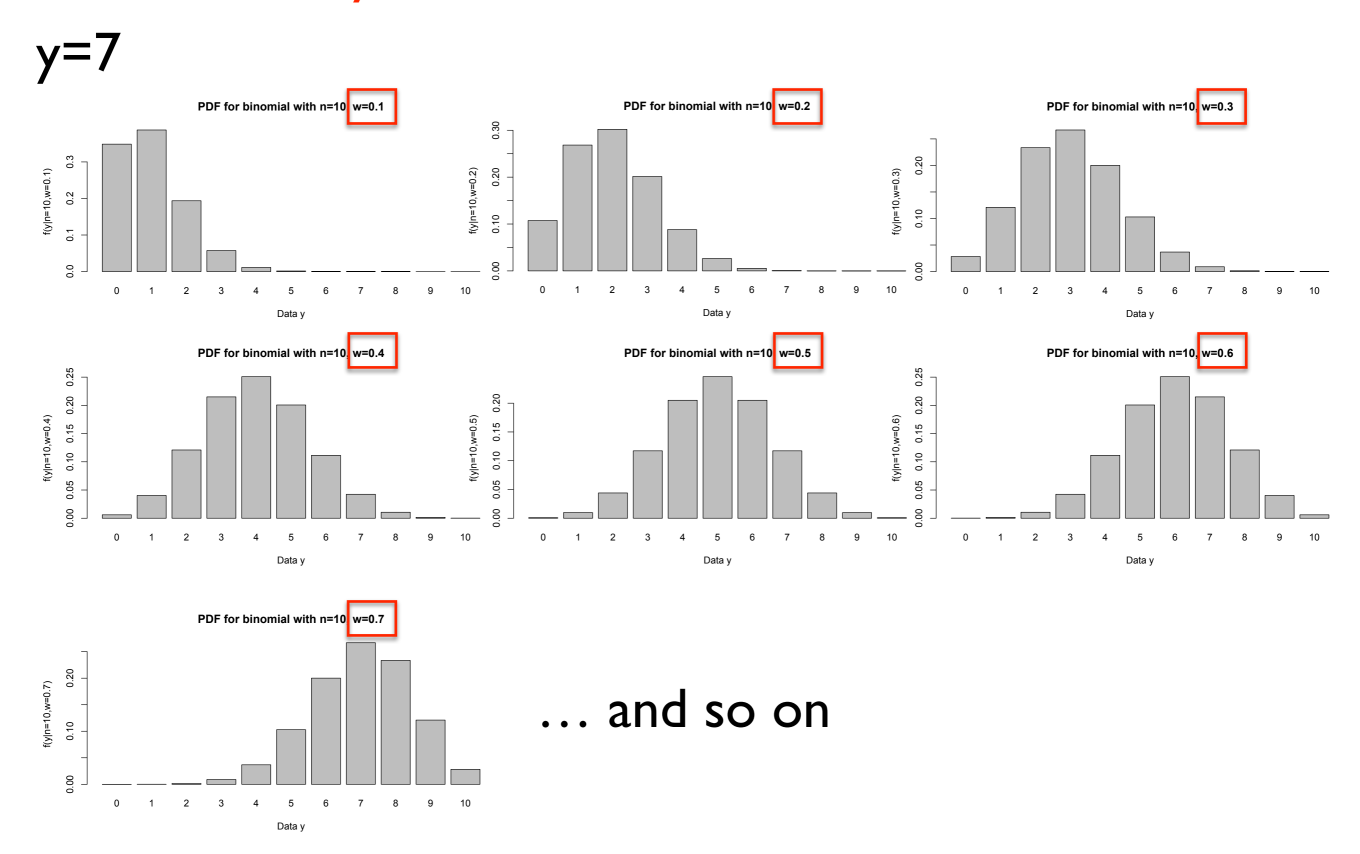
$$L(w|y) = f(y|w)$$

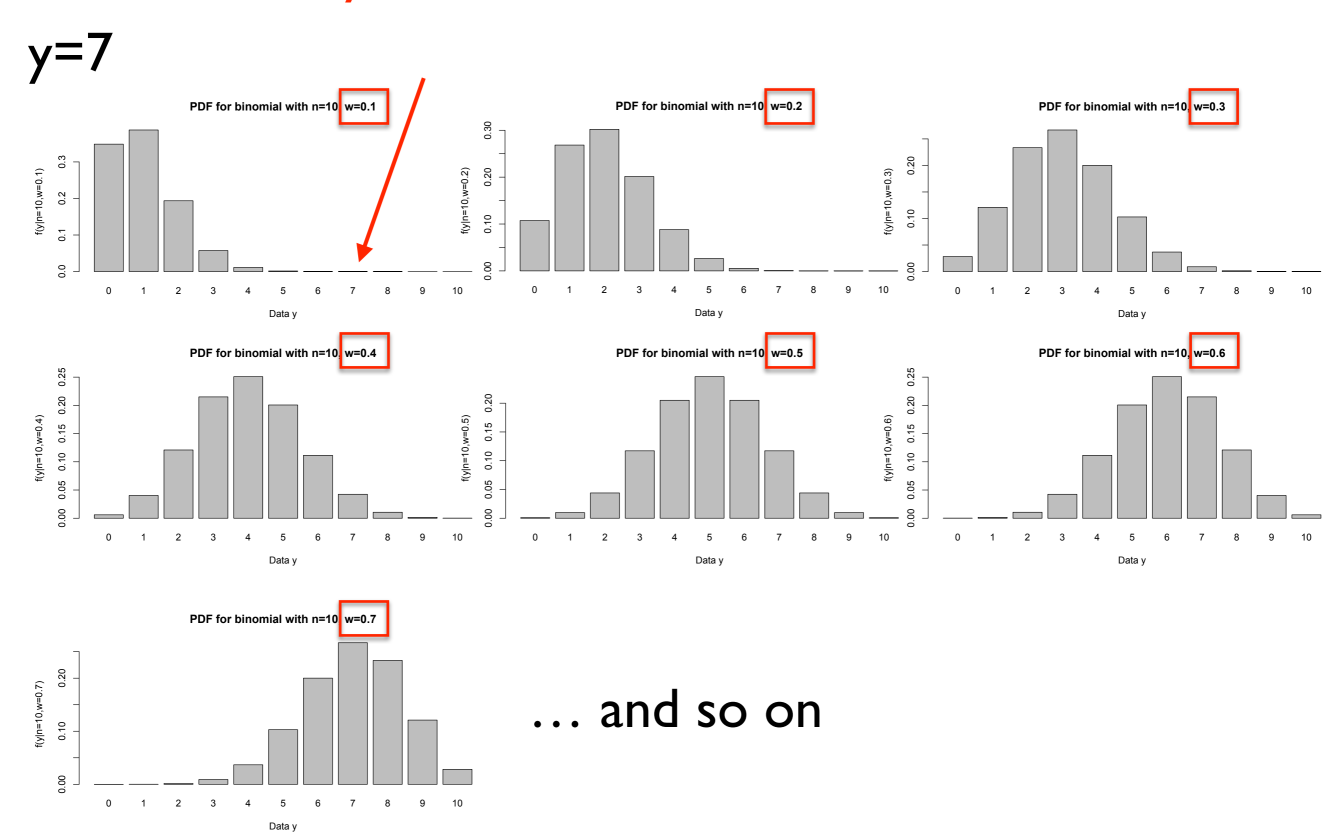
- L(w|y) represents the likelihood of the parameter w given the observed data y
- For our one-dimensional binomial example the likelihood function for y=7 and n=10 is

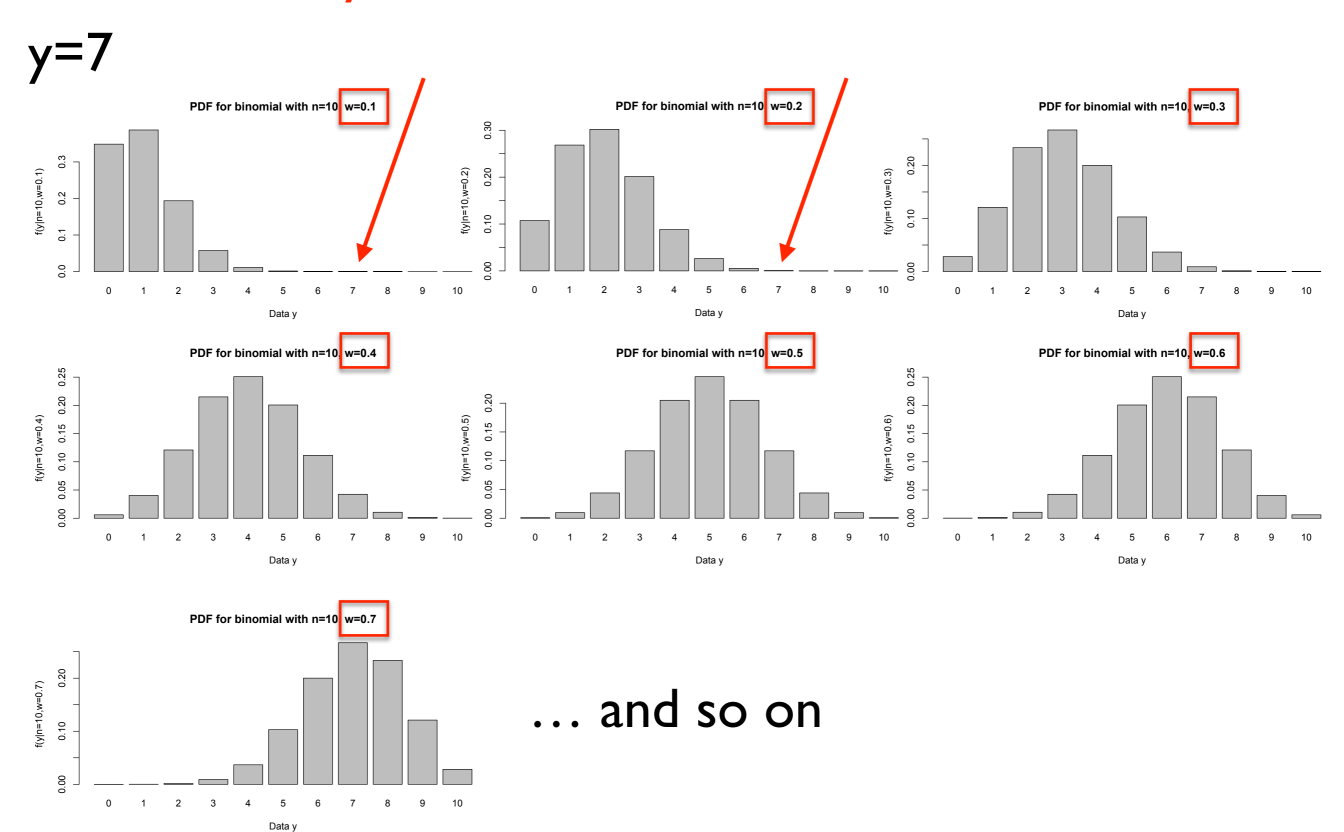
$$L(w|n = 10, y = 7) = f(y = 7|n = 10, w)$$

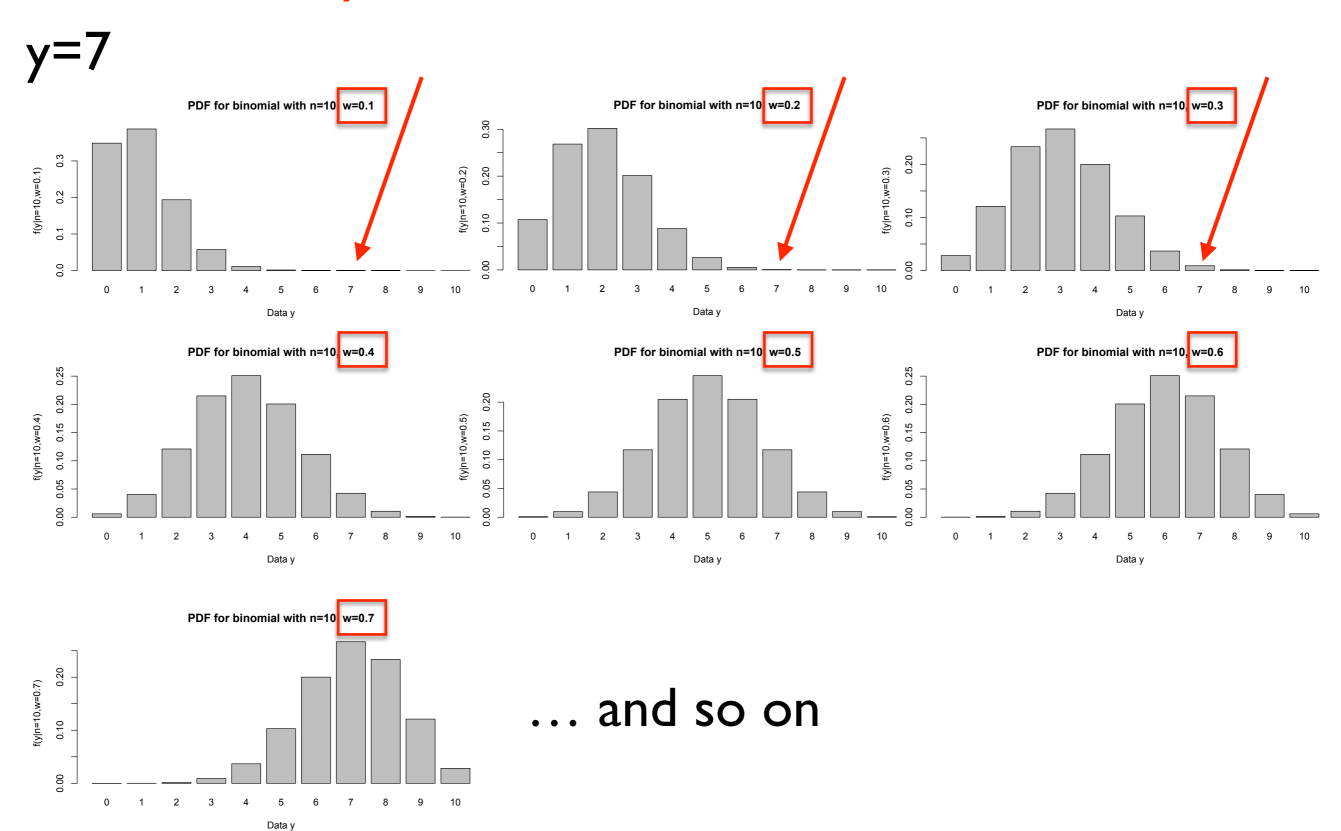
$$= \frac{10!}{7!3!} w^7 (1 - w)^3 \quad (0 \le w \le 1)$$

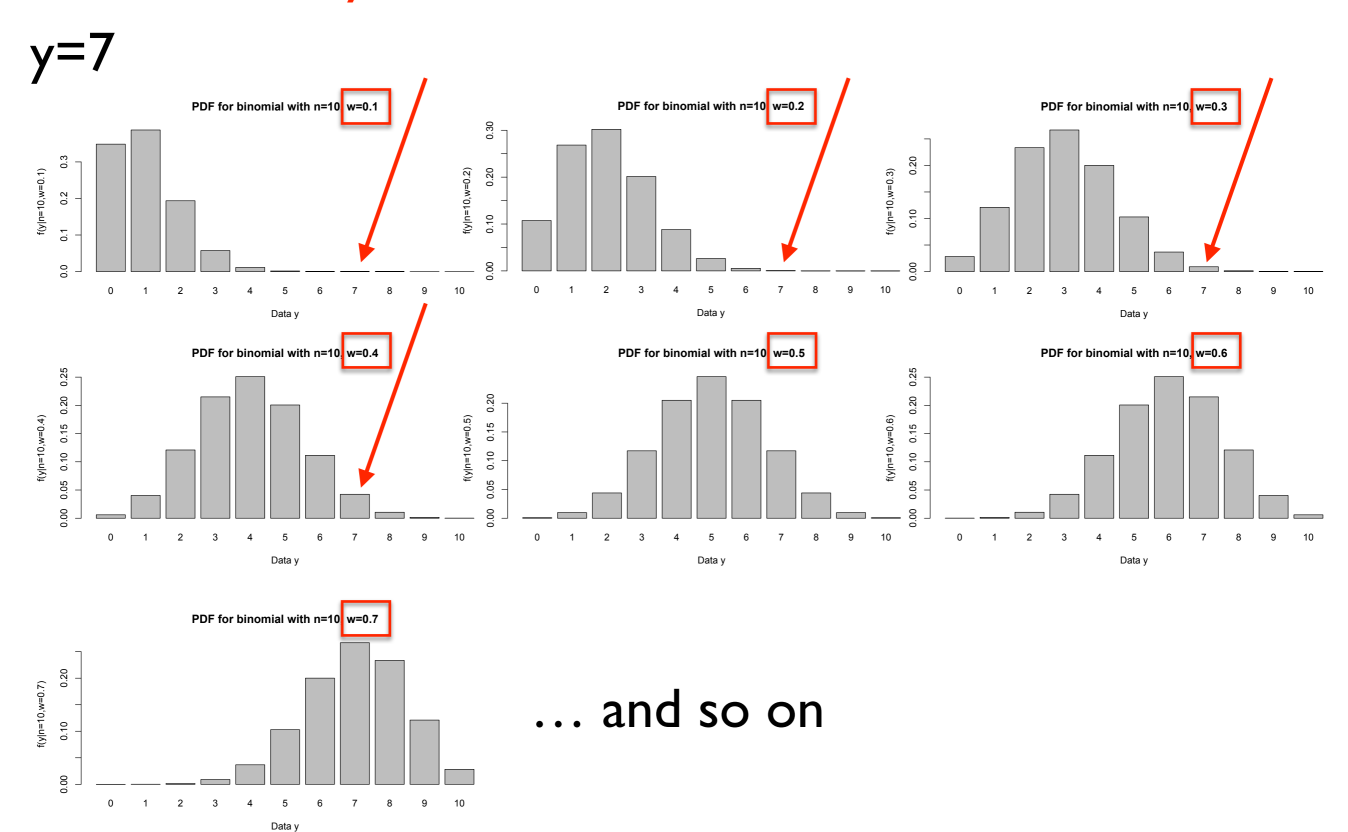
but what value of w?

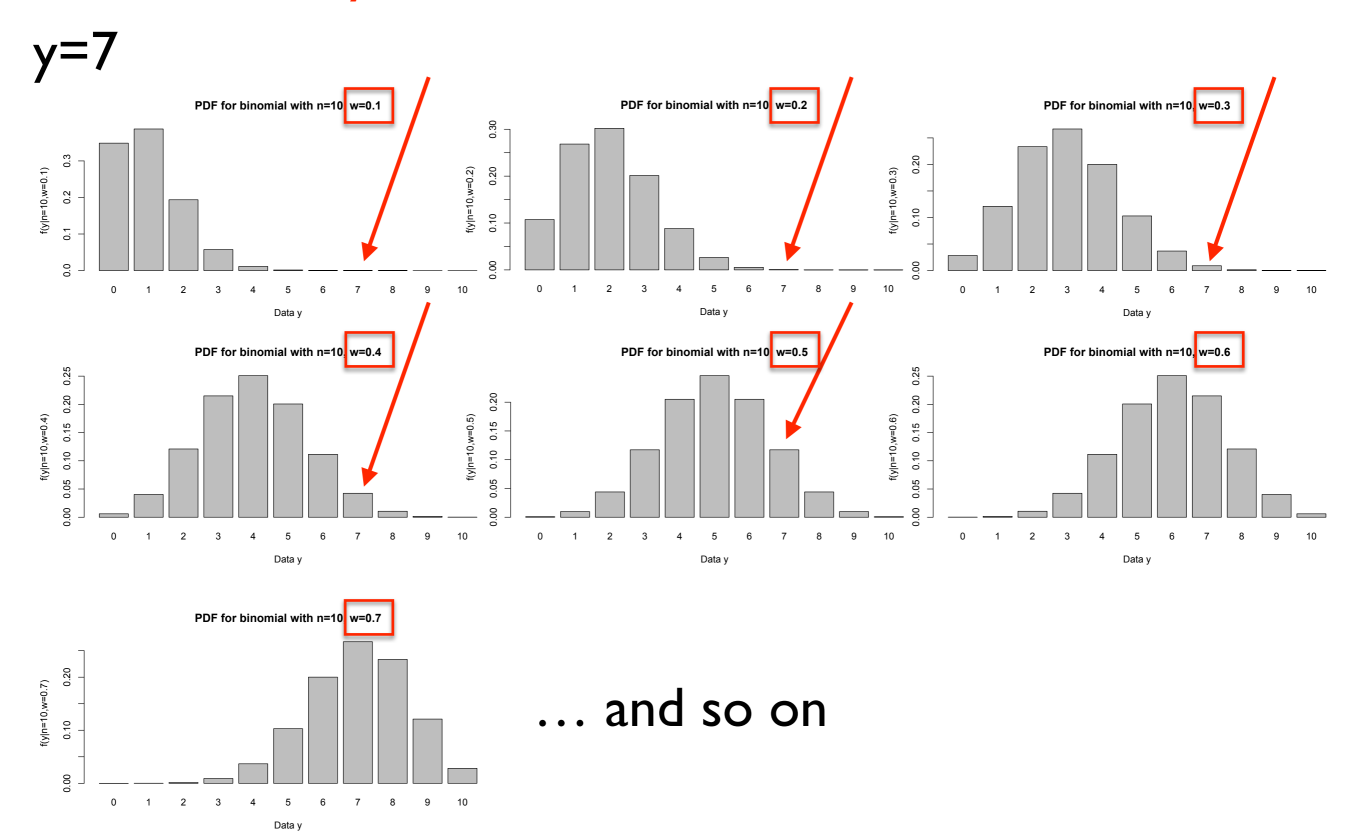


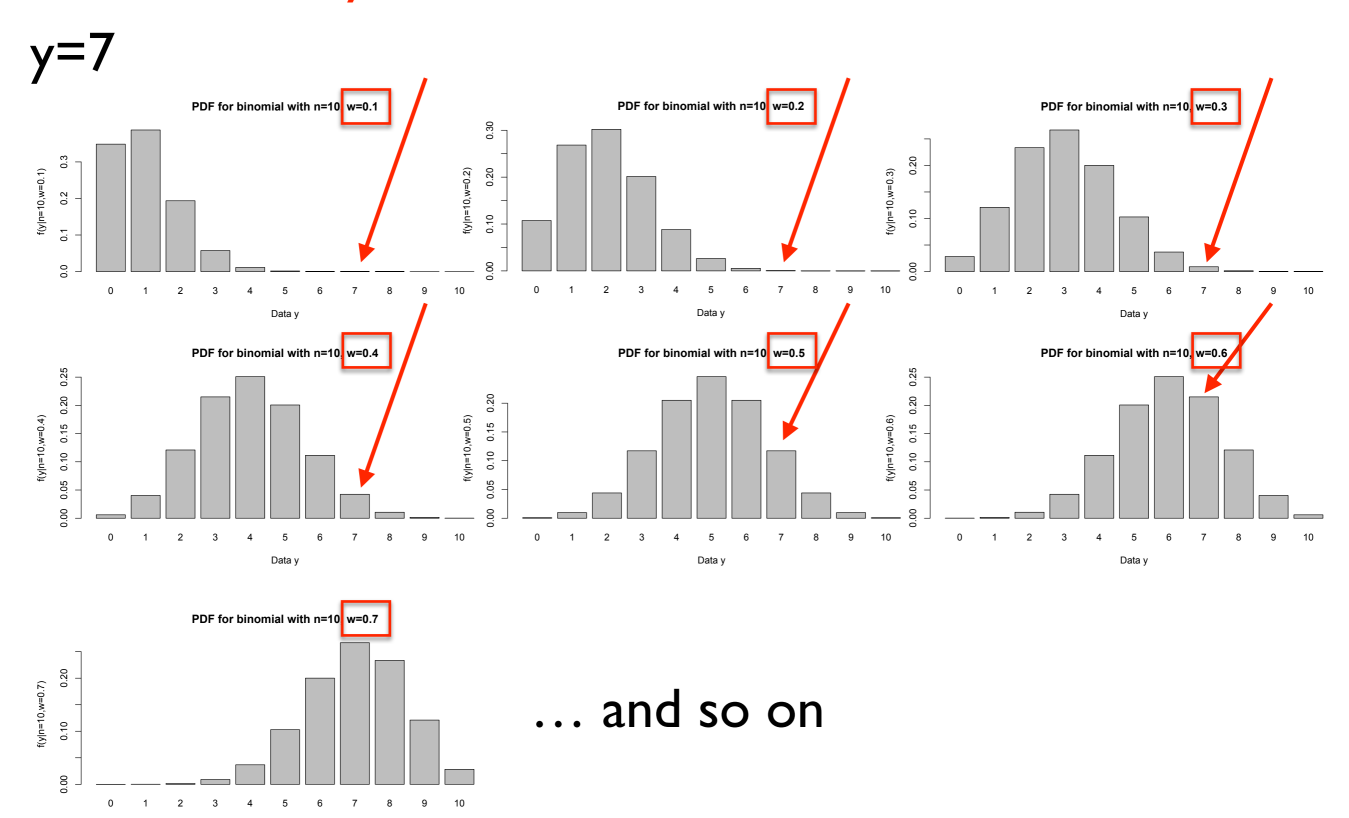




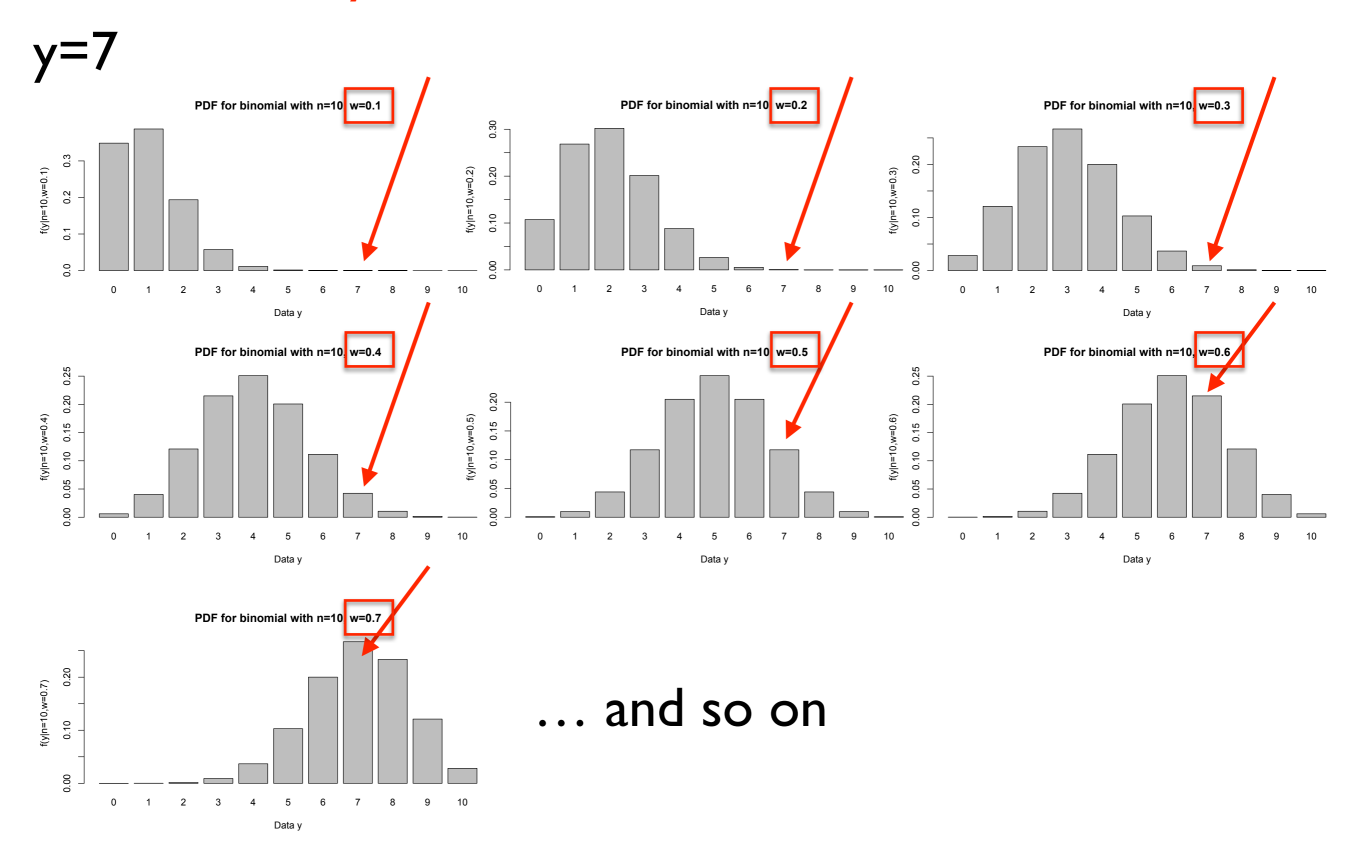


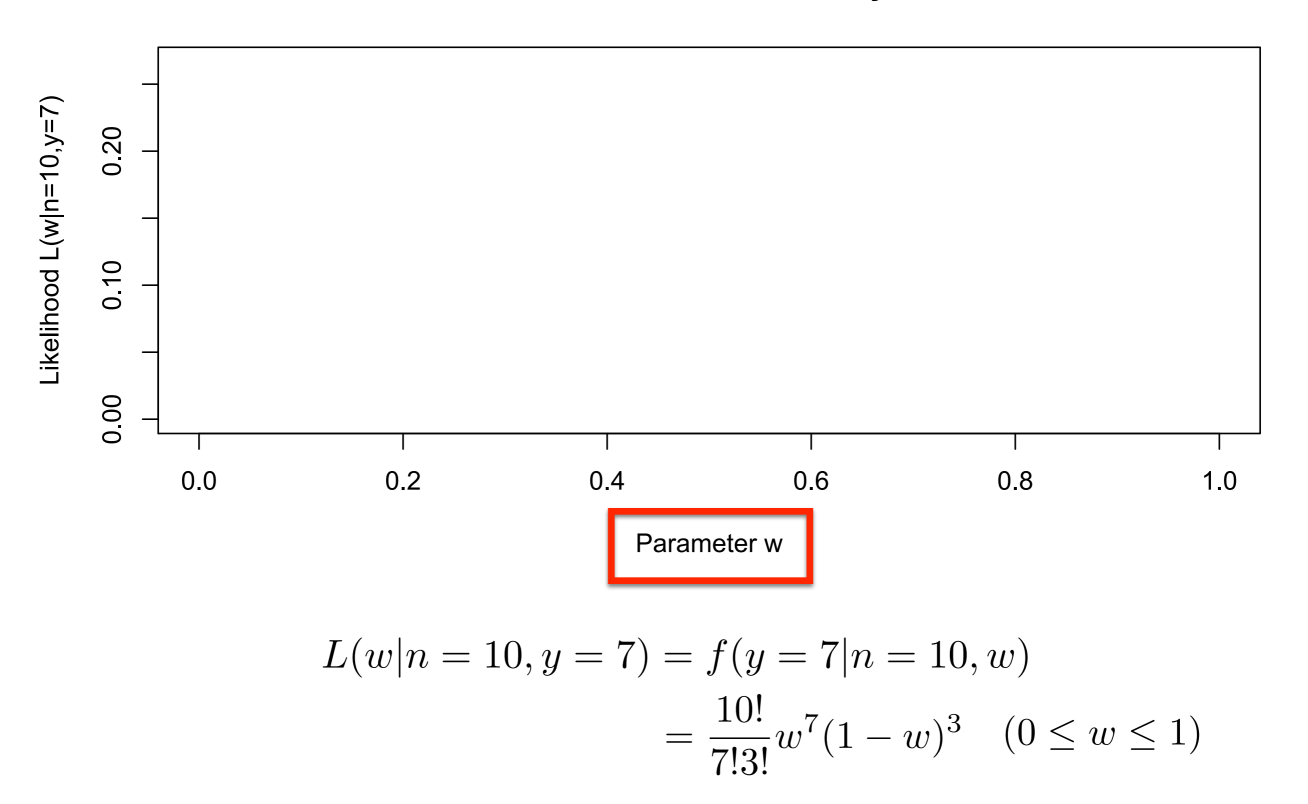


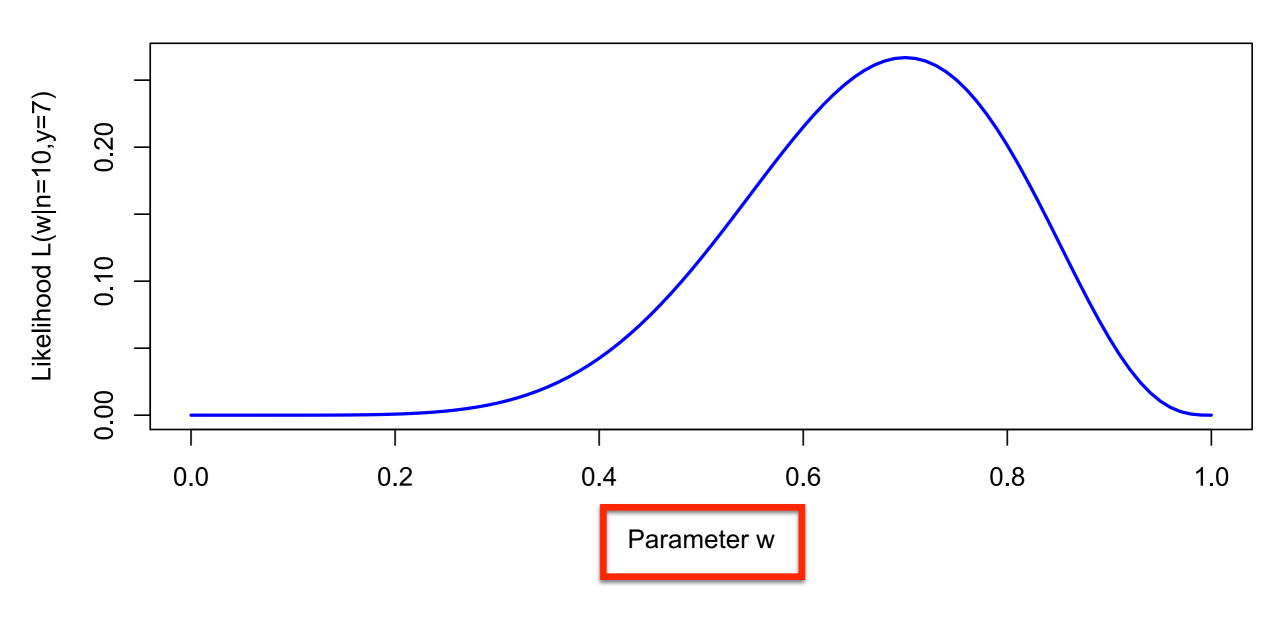




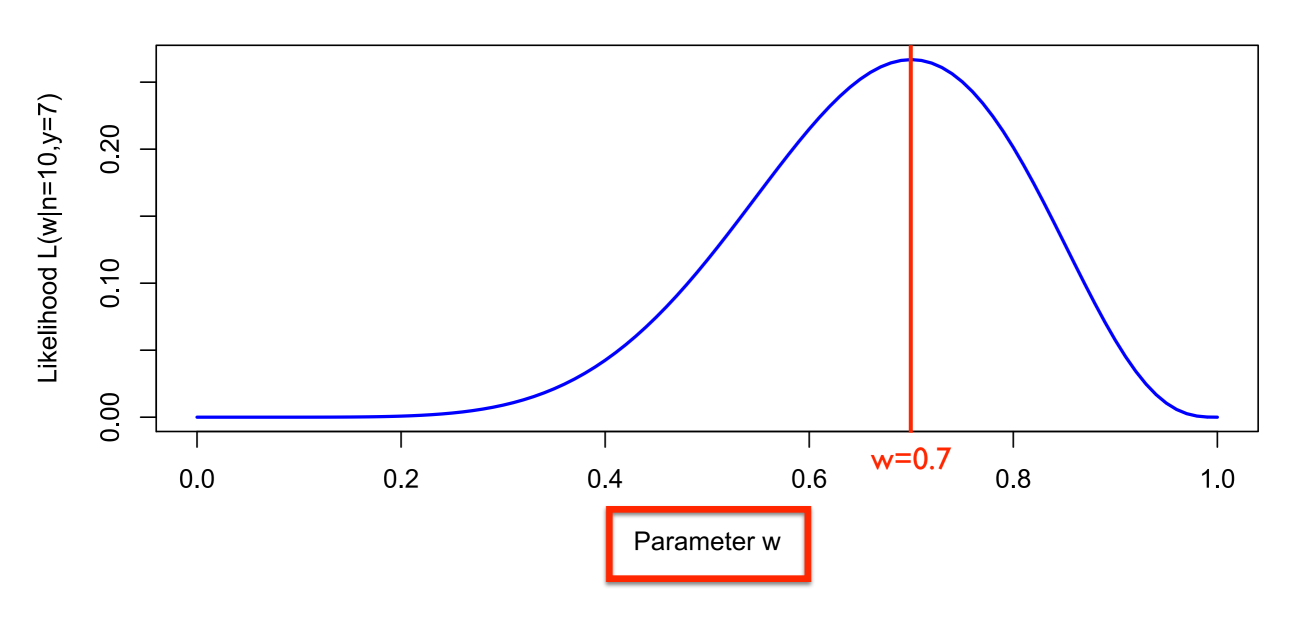
Data y





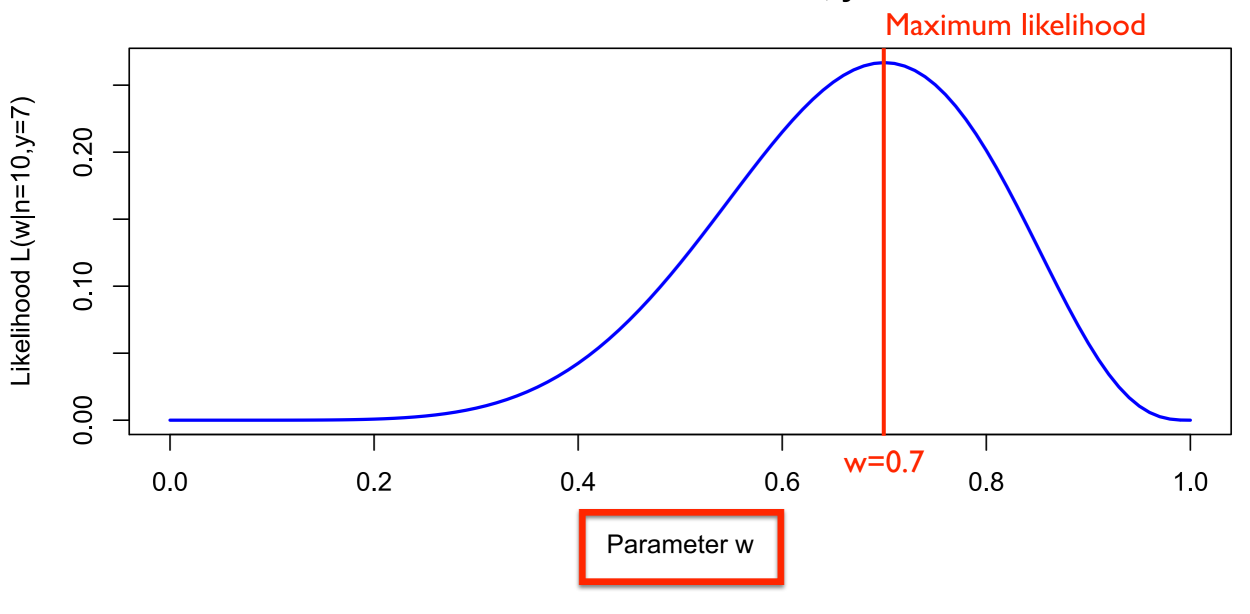


$$L(w|n = 10, y = 7) = f(y = 7|n = 10, w)$$
$$= \frac{10!}{7!3!} w^7 (1 - w)^3 \quad (0 \le w \le 1)$$



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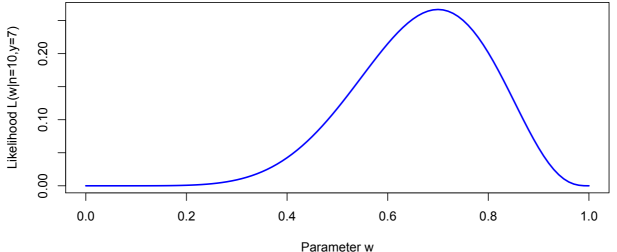


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$$= \frac{10!}{7!3!} w^7 (1 - w)^3 \quad (0 \le w \le 1)$$

Maximum Likelihood Estimation

- find the probability distribution (the model) that makes the observed data most likely
- seek the value of the parameter vector w that maximizes the likelihood function L(w|y)
- the resulting parameter vector w is known as the MLE estimate

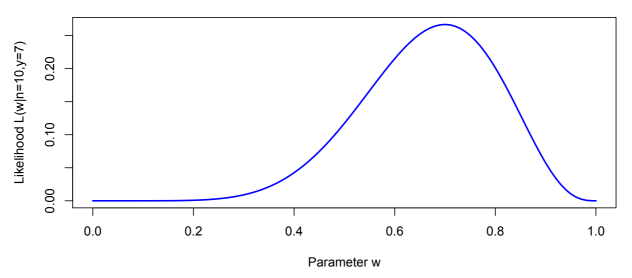
Maximum Likelihood Estimation



- three ways of finding the MLE
- I. analytically: use calculus to solve for the parameter value(s) w that result in a peak
- zero derivative and a negative second derivative

$$\frac{\partial L}{\partial w} = 0 \qquad \frac{\partial^2 L}{\partial^2 w} < 0$$

Maximum Likelihood Estimation Likelihood of w for n=10, y=7



- three ways of finding the MLE
- 2. grid search: exhaustive search through parameter space
- (inefficient, could take long time for high dimensional parameter vector)

Maximum Likelihood Estimation

Likelihood of w for n=10, y=7

(kulu=10,y=7)
(V=0,y=10,y=10)

- three ways of finding the MLE
- 3. **numerically**: use non-linear optimization (e.g. gradient descent) to iteratively find the peak

we saw before that the PDF for observed data, y = (y I, ..., ym) given a parameter vector w, can be expressed as the product (multiply) of PDFs for individual observations

$$L(w|y = (y_1, y_2, \dots, y_n)) = L_1(w|y_1)L_2(w|y_2)\dots L_n(w|y_n)$$

$$f(y = (y_1, y_2, \dots, y_n)|w) = f_1(y_1|w)f_2(y_2|w)\dots f_n(y_n|w)$$

 $p(y = (y_1, y_2, y_3)|\mu, \sigma) = (.010934)(.021297)(.003599) = .000000838$

- multiplying together a lot of values that lie between 0 and 1, (as many as there are data points) will result in a very small number
- in fact the more data, the smaller the resulting product will be
- computers are not good at representing very small numbers

- solution: take the logarithm
- this reformulates the series of products, as a series of sums
- the more data, the higher the resulting sum

```
\ln \left[ L_1(w|y_1) L_2(w|y_2) \dots L_n(w|y_n) \right] = \ln \left[ L_1(w|y_1) \right] + \ln \left[ L_2(w|y_2) \right] + \dots + \ln \left[ L_n(w|y_n) \right]
```

- another problem: most optimization algorithms are formulated in terms of **minimizing** an objective function, not maximizing
- solution: rather than maximizing the log-likelihood, we will minimize the negative log-likelihood

```
find w that minimizes : -\ln[L(w|y)]
find w that minimizes : -\ln[L_1(w|y_1)] - \ln[L_2(w|y_2)] - \cdots - \ln[L_n(w|y_n)]
```

An Example

- Let's say I claim I can correctly identify espresso brewed with Illy beans (as opposed to Lavazza beans)
- My lab designs an experiment to test me
- They give me 20 cups of coffee in random order and I have to say "Illy" or "Lavazza"
- Observed data: I get 16 correct, 4 incorrect

An Example

- Observed data: I get 16 correct, 4 incorrect
- This experiment can be modelled as 20
 Bernoulli trials (outcome of each trial is random and can be either of two possible outcomes,
 "success" and "failure")
- we know PDF is binomial, which has 2
 parameters: n (# trials) and w (prob of a success
 on a given trial)

An Example

- we know PDF is binomial, which has 2 parameters: n
 (# trials) and w (prob of a success on a given trial)
- what model explains the observed data?
- equivalent to asking, what is the value of the parameter w?
- high w (e.g. near 1.0) means I have a good ability to discriminate
- w near 0.5 means I am flipping a coin

Likelihood function

 binomial distribution: gives probability of observing y successes in n trials, given probability w of success on any single trial

$$prob(y|n, w) = \frac{n!}{y!(n-y)!}w^y(1-w)^{n-y}$$

Likelihood function

- in our experiment, n=20, y=16 and w is unknown
- our likelihood function needs to provide likelihood of a particular value of parameter w, given n=20 and y=16

$$L(w|n = 20, y = 16) = \frac{20!}{16!4!}w^{16}(1-w)^4$$

Likelihood function

now let's take the logarithm:

$$L(w|n = 20, y = 16) = \frac{20!}{16!4!}w^{16}(1-w)^4$$

$$\ln\left[L(w|n=20,y=16)\right] = \ln\left[\frac{20!}{16!4!}\right] + 16\ln\left[w\right] + 4\ln\left[(1-w)\right]$$

Find MLE w

$$\ln\left[L(w|n=20, y=16)\right] = \ln\left[\frac{20!}{16!4!}\right] + 16\ln\left[w\right] + 4\ln\left[(1-w)\right]$$

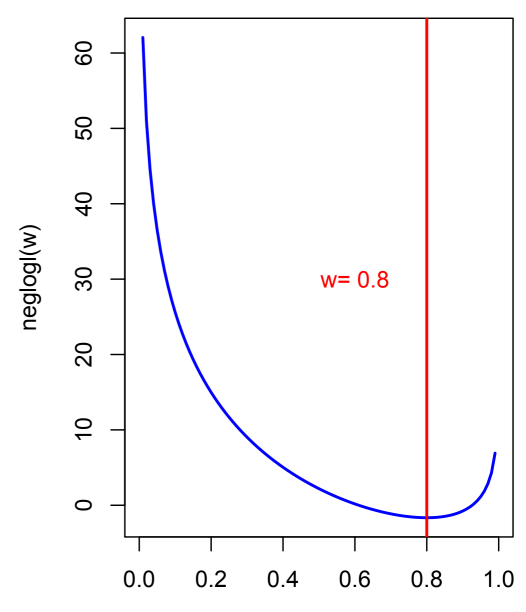
- we have our log-likelihood function
- now we need to find w that minimizes the negative log-likelihood

Find MLE for w: brute force

$$\ln\left[L(w|n=20,y=16)\right] = \ln\left[\frac{20!}{16!4!}\right] + 16\ln\left[w\right] + 4\ln\left[(1-w)\right]$$

```
> neglogl <- function(w) {
    loglik <- log(116280) + 16*log(w) + 4*log(1-w)
    return(-1*loglik)
}
> w <- seq(0,1,.01)
> plot(w, neglogl(w), type="l", col="blue", lwd=2)
> imin <- which(neglogl(w)==min(neglogl(w)))
> abline(v=w[imin], col="red", lwd=2)
> text(.6, 30, paste("w=",w[imin]),col="red")
```

the MLE for w given the data y=16 (and n=20) is w=0.80



W

Find MLE for w: optimize

$$\ln\left[L(w|n=20, y=16)\right] = \ln\left[\frac{20!}{16!4!}\right] + 16\ln\left[w\right] + 4\ln\left[(1-w)\right]$$

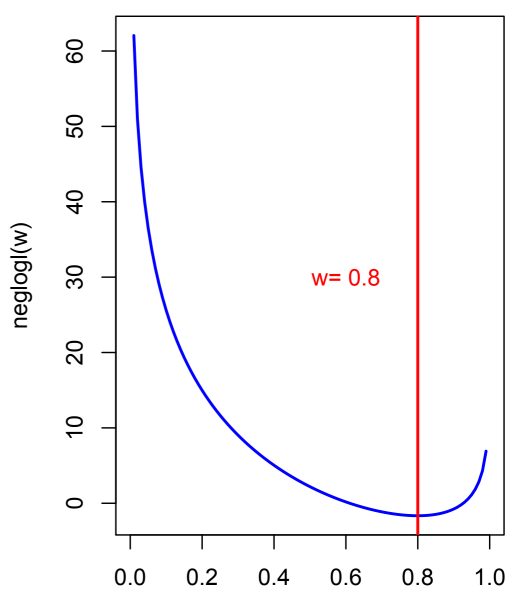
```
> neglogl <- function(w) {
        loglik <- log(116280) + 16*log(w) + 4*log(1-w)
        return(-1*loglik)
}
> nlm(f=neglogl, p=0.5)
$minimum
[1] -1.655708

$estimate
[1] 0.7999995

$gradient
[1] -8.881784e-10

$code
[1] 1

$iterations
[1] 7
```

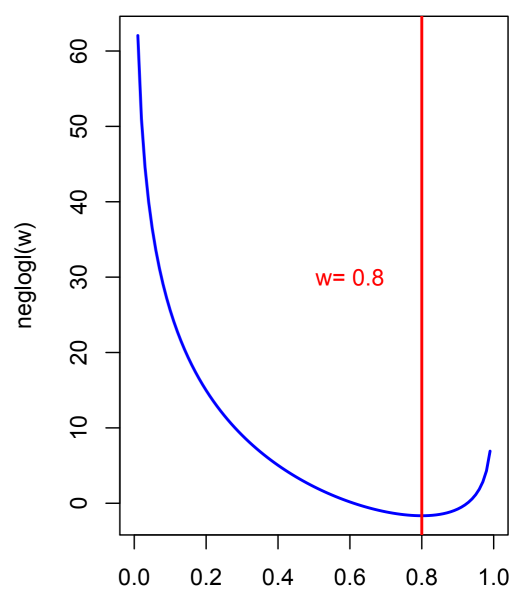


W

Find MLE for w: optimize

$$\ln\left[L(w|n=20, y=16)\right] = \ln\left[\frac{20!}{16!4!}\right] + 16\ln\left[w\right] + 4\ln\left[(1-w)\right]$$

[1] 7



W

MLE for binomial

- in fact it is known for binomial that MLE for w is equal to y/n
- 16/20
- \bullet = 0.80

MLE for binomial

- if we approximate the binomial distribution with a normal distribution (OK for large #s of observations)
- confidence interval is $\hat{w} \pm z_{1-\frac{\alpha}{2}} \sqrt{\frac{\hat{w}(1-\hat{w})}{n}}$
- so 95% confidence interval for Illy is

$$0.8 \pm 1.96\sqrt{\frac{0.8(1-0.8)}{20}} = 0.8 \pm 0.175$$

 \bullet = 0.625 - 0.975

MLE in general

- MLE for many distributions are known (look it up)
- MLE for more complex models can sometimes be determined analytically
- Often however not possible/feasible
- iterative optimization is a common method in these cases

Optimization: Local Minima

 repeat optimization starting from different initial guesses

Optimization: Local Minima

 use stochastic optimization algorithms like simulated annealing

The Bottom Line

- If you can write an equation for the Likelihood function
- i.e. probability of obtaining your observed data, given a model with parameter(s) w
- then you can find the MLE for w
- i.e. you can find the model that is most likely to generate your data

Analytic Solutions: Bernoulli Distribution

find
$$w$$
 for $\frac{\partial (L(w|n,y))}{\partial w} = 0$

gives
$$w = \frac{\sum y_i}{n}$$

http://mathworld.wolfram.com/MaximumLikelihood.html

Normal Distribution

$$f(x_1, \dots, x_n | \mu, \sigma) = \prod \frac{1}{\sigma \sqrt{2\pi}} e^{-(x_i - \mu)^2 / (2\sigma^2)}$$

$$= \frac{(2\pi)^{-n/2}}{\sigma^n} \exp\left[-\frac{\sum (x_i - \mu)^2}{2\sigma^2}\right]$$
so $\ln f = -\frac{1}{2} n \ln(2\pi) - n \ln \sigma - \frac{\sum (x_i - \mu)^2}{2\sigma^2}$
and $\frac{\partial (\ln f)}{\partial \mu} = \frac{\sum (x_i - \mu)}{\sigma^2} = 0$
giving $\hat{\mu} = \frac{\sum x_i}{n}$

http://mathworld.wolfram.com/MaximumLikelihood.html

Normal Distribution

Similarly,
$$\frac{\partial(\ln f)}{\partial\sigma} = -\frac{n}{\sigma} + \frac{\sum(x_i - \mu)^2}{\sigma^3} = 0$$

gives
$$\hat{\sigma} = \sqrt{\frac{\sum (x_i - \hat{\mu})^2}{n}}$$

http://mathworld.wolfram.com/MaximumLikelihood.html

Hypothesis Testing

- We can use the Likelihood Ratio Test to compare two models
- e.g. Illy vs Lavazza example:
- 16 correct out of 20 trials
- our MLE for p was 0.80
- let's test this against a null hypothesis that p=0.50

- test statistic D is a ratio:
- D = -2 In ((likelihood for null model) / (likelihood for alternative model))
- D = -2 In (likelihood null) + 2 In (likelihood alt)

- the probability distribution of test statistic
 D is approximately a chi-squared
 distribution with df = df2-df1
- df2 and df1 are number of free parameters of models 1 (null) and 2 (alternative)

- Illy vs Lavazza:
- null model is L(p=0.5|data)
- alternative model is p for max(L(p|data))
 (p=0.8)
- df for null = 0 (no parameters are free to vary)
- df for alt = I (p is free to vary)

$$L(p|y,n) = \frac{n!}{y!(n-y)!}p^y(1-p)^{n-y}$$

- D = -2 In (likelihood null) + 2 In (likelihood alt)
- our data: 16 correct and 4 incorrect
- -2 In (L(p=0.5 | y=16, n=20)) = 16.29966
- MLE of p is p=0.8, so
- 2 In (L(p=0.8 | y=16, n=20)) = -4.82984
- D = 16.29966 4.82984 = 11.46982

- \bullet D = 11.46982
- now compute a p-value using chi-square distribution with df = I-0 = I

```
pval <- pchisq(q=11.46982, df=1, lower.tail=FALSE)
0.0007073553</pre>
```

- p-value = 0.00071
- we can reject the null with a Type-I error rate of 0.0007 I (7 in 10,000)