Joachim Vandekerckhove

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 - Ideally, we could just show a graph of it and leave the interpretation to the reader
 - But often the posterior will have many dimensions
 - And also that seems lazy
- We need a way to describe the posterior distribution
 - Mean? SD? Skew? Kurtosis? Mass at or around a certain value? $p(.8 \le P_R \le .9 | \#R, \#W)$?

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Here, with the discrete domain of P_R :

$$p(.8 \le P_R \le .9 | \#R, \#W) =$$

$$p(P_R = .80 | \#R, \#W)$$

$$+p(P_R = .85 | \#R, \#W)$$

$$+p(P_R = .90 | \#R, \#W)$$

 \bullet A more continuous case on $x=\frac{P_R+1}{2}$

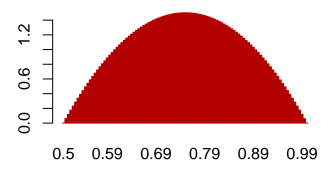
- \bullet A more continuous case on $x=\frac{P_R+1}{2}$
 - $p(x) = Kx^{\alpha-1}(1-x)^{\beta-1} = B(x|\alpha,\beta)$

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 - Likelihood is the same: $p(\#R, \#W|P_R) = CP_R^{\#R}(1-P_R)^{\#W}$
 - So the posterior must be: $p(P_R|\#R,\#W) \propto \left(\frac{P_R+1}{2}\right)^{\alpha-1} \left(1-\frac{P_R+1}{2}\right)^{\beta-1}$

Functional programming

Sometimes it is useful in R to turn a function into a variable to change it quickly

You can make a function "on the fly" inside a function or script file like this:

```
funcname <- function(n, x) { rep(x, n) }</pre>
```

So that prior and likelihood can be written like:

```
prior <- function(p) { dbeta(2 * (p-.5), 2, 2) }
likelihood <- function(p) { dbinom(5, 6, p) }</pre>
```

Functional programming

Prior and likelihood:

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prior <- function(p) { dbeta(2 * (p-.5), 2, 2) }
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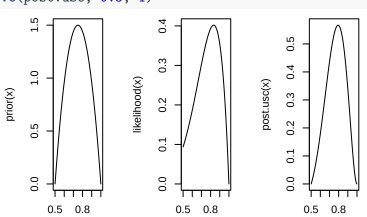
Given those, building the posterior is trivial:

```
post.usc <- function(p) { prior(p) * likelihood(p) }</pre>
```

Exercise: implement this, plot the three functions

Functional programming

```
par(mfrow=c(1,3))
curve(prior, 0.5, 1)
curve(likelihood, 0.5, 1)
curve(post.usc, 0.5, 1)
```



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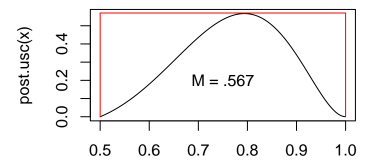
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- How do we determine the mean of an arbitrary, somewhat complicated function?
- As it turns out, drawing random samples from a distribution is an efficient way to do that
 - Methods for doing this are called Monte Carlo methods
 - Math win: Monte Carlo methods don't need those hard-to-compute K and C scaling constants

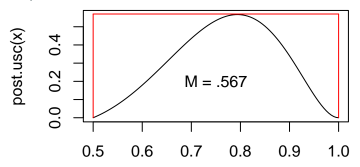
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1) Draw a sample from some basic distribution S(x|...)



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- 1) Draw a sample from some basic distribution S(x|...)
- 2) Reject the sample with probability $q=\frac{fx}{M\times S(x)}$, where M is chosen so that this is always ≤ 1 (but ideally sometimes close to 1)



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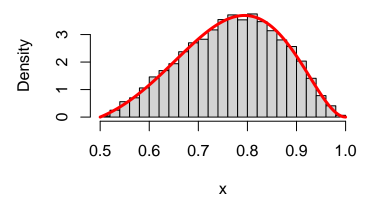
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- 3. Repeat many times to get a few thousand samples
- 4. Make a histogram and compare the shapes of the distribution

```
N < -10000
x <- vector(,N)
c <- 1
M < -.567
while(c <= N) {
  x[c] \leftarrow runif(1, 0.5, 1)
  u \leftarrow runif(1, 0, M)
  if (u < post.usc(x[c])) c <- c + 1;
hist(x, breaks=25,freq=FALSE)
K <- integrate(post.usc, 0.5, 1)$value</pre>
lines(domain, post.usc(domain)/K, lwd=3, col='red')
```





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 - ... or the proportion of samples that are > .85: 0.2304
 - ... or indeed any quality we fancy

Basic unit of a Bayesian model is a distribution function

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- Basic unit of a Bayesian model is a distribution function
- The posterior is defined through a generative model representation
 - which is basically a sequence of distributional assumptions
 - In Bayesian statistics, a "model" is just a special kind of distribution function over parameters and data (and so with possibly very many dimensions)

Let's define a really trivial model \mathcal{M}_t in which we estimate the parameters μ and τ (= $1/\sigma^2$) of a normal distribution, applied to some data points d_j :

$$\mathcal{M}_t: \begin{cases} \forall j \in (1, \dots, J) : d_j \sim N(\mu, \tau) \\ \mu \sim N(0, 0.1) \\ \tau \sim \Gamma(4, 0.01) \end{cases}$$

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$$p(d_1, \dots, d_J \mid \mathcal{M}_t) = \left(\prod_{j=1}^J N(d_j \mid \mu, \tau) \right) \times N(\mu \mid 0, 0.1) \times \Gamma(\tau \mid 4, 0.01)$$

JAGS code is (almost) perfect

```
\mathcal{M}_s: \begin{cases} \forall j \in (1, \dots, J) : d_j \sim N(\mu, \tau) \\ \mu \sim N(0, 0.1) \\ \tau \sim \Gamma(4, 0.01) \end{cases}
```

The program needs to know the specifics of the model:

```
model {
    for (j in 1:J) {
        d[j] ~ dnorm(mu, tau)
    }
    mu ~ dnorm(0,0.1)
    tau ~ dgamma(4,0.01)
}
```

A psychological model: Signal detection theory

$$\mathcal{M}_{sdt}: \begin{cases} \delta \sim N(1,1) & \beta \sim N(0,1) \\ \phi_h = \Phi(\delta/2 - \beta) & \phi_f = \Phi(-\delta/2 - \beta) \\ h \sim B(\phi_h, n_s) & f \sim B(\phi_f, n_n) \end{cases}$$

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```

```
model {
    d ~ dnorm(1, 1)
    b ~ dnorm(0, 1)

phih <- phi( d / 2 - b)
    phif <- phi(-d / 2 - b)

h ~ dbin(phih, sigtrials)
    f ~ dbin(phif, noistrials)
}</pre>
```

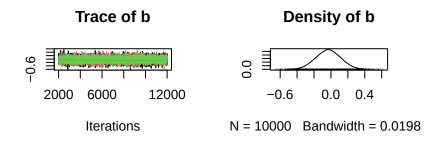
Signal detection theory \sim implementation

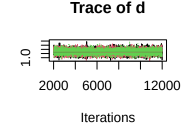
```
library(rjags)
data \leftarrow list(h = 32, sigtrials = 39,
               f = 12 , noistrials = 60 )
modelString = "
    model {
        d \sim dnorm(1, 1)
        b \sim dnorm(0, 1)
        phih \leftarrow phi(d/2-b)
        phif \leftarrow phi(-d / 2 - b)
        h ~ dbin(phih, sigtrials)
         f ~ dbin(phif, noistrials)
11
```

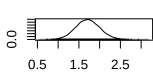
Signal detection theory \sim implementation

```
writeLines( modelString , con = "sdt.txt" )
jagsModel = jags.model( file = "sdt.txt" ,
                      data =
                                    data ,
                                    3,
                      n.chains =
                      n.adapt = 1000)
set.seed(0)
update( jagsModel , n.iter = 1000 ) # burn-in
samples = coda.samples( jagsModel ,
                      variable.names = c("d", "b"),
                      n.iter
                                           10000)
```

Signal detection theory ~ results







Density of d

Signal detection theory ~ summary statistics

```
## Mean SD Naive SE Time-series SE
## b -0.03205525 0.1468885 0.0008480611 0.001147812
## d 1.71389374 0.2878064 0.0016616508 0.002213016

summary(samples)$quantiles
## 2.5% 25% 50% 75% 97.5%
```

b -0.3202336 -0.1306705 -0.03203559 0.06660237 0.256125 ## d 1.1586158 1.5191690 1.71022464 1.90596253 2.292419

Signal detection theory ~ convergence

```
effectiveSize(samples)
          b
##
## 16407.95 16954.86
gelman.diag(samples)
## Potential scale reduction factors:
##
##
     Point est. Upper C.I.
## b
## d
##
## Multivariate psrf
##
## 1
```

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JAGS (and Stan) have a straightforward modeling language that lets users specify a model

They then draw samples from the posterior distributions, so we can calculate any summaries we like