Bayesian Methods

A data generating process is a model characterized by a parameter vector, and the data is generated from the model at a **fixed parameter** vector, θ_0 .

Bayesians view data as given, and update beliefs about a parameter using the information about the parameter contained in the data.

There's nothing obviously contradictory in these views. Nevertheless, it's not hard to find discussions where there are disagreements.

Here, I'm trying to address a model with a fixed non-random parameter about which we would like to learn. As long as the object of interest (the dgp and it's parameter) is agreed upon, then we can contemplate using any convenient tool.

Frequentist

- θ is fixed, but unknown
- Uncertainty comes from the sampling uncertainty. That is, from the fact that we can get different samples.
- All probabilistic statements are statements about sampling uncertainty.

Bayesian

- θ is random
- · Data is treated as fixed after observed
- All probabilistic statements are about uncertainty about heta

As long as one takes the view that there is a fixed unknown parameter value θ_0 which generates all samples, then frequentist and Bayesian methods are trying to inform us about the same object, and the choice between tools may become one of convenience.

Definitions

The Bayesian approach summarizes beliefs about parameters using a density function:

- There is a true unknown parameter vector, θ_0 , and the density, $\pi(\theta)$, which is known as the prior, reflects current beliefs about the parameter, before observing the sample. It is assumed that the econometrician can provide this density.
- We also have sample information, $y = \{y_1, y_2, \dots, y_n\}$. We're already familiar with the likelihood function, $f(y|\theta)$, which is the density of the sample given a parameter value.

Given these two pieces, we can write the joint density of the sample and the beliefs:

$$f(v, \theta) = f(v|\theta)\pi(\theta)$$

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We can get the marginal likelihood by integrating out the parameter, integrating over its support Θ :

$$f(y) = \int_{\Theta} f(y, \theta) d\theta$$

The last step is to get the **posterior** of the parameter. This is simply the density of the parameter conditional on the sample, and we get it in the normal way we get a conditional density, using **Bayes' theorem** (https://www.youtube.com/watch?v=HZGCoVF3YvM):

$$f(\theta|y) = \frac{f(y,\theta)}{f(y)} = \frac{f(y|\theta)\pi(\theta)}{f(y)}$$

- The movement from the prior to the posterior reflects the **learning** that occurs about the parameter when one receives the sample information.
- The sources of information used to make the posterior are the prior and the likelihood function.
- Once we have the posterior, one can provide a complete probabilistic description about our updated beliefs about the parameter, using quantiles or moments of the posterior.

So far, this is pretty straightforward. The complications are mostly computational. To illustrate, the posterior mean is

$$E(\theta|y) = \int_{\Theta} \theta f(\theta|y) d\theta = \frac{\int_{\Theta} \theta f(y|\theta) \pi(\theta) d\theta}{\int_{\Theta} f(y,\theta) d\theta}$$

- · One can see that a means of integrating will be needed.
- Only in very special cases will the integrals have analytic solutions.
- Otherwise, computational methods will be needed. Advances in computational methods are what have lead to the increased use of Bayesian methods.

Extensions

<u>Chernozhukov and Hong (2003) "An MCMC Approach to Classical Estimation"</u>
<u>(http://www.sciencedirect.com/science/article/pii/S0304407603001003)</u> is a very interesting article that shows how Bayesian methods may be used with criterion functions that are associated with classical estimation techniques.

For example, when the criterion function $L_n(\theta)$ in their paper is set to the log-likelihoodfunction and the pseudo-prior $\pi(\theta)$ is a real Bayesian prior, the class of estimators discussed by CH reduces to the ordinary Bayesian posterior mean. this estimator is consistent and asymptotically normally distributed. In particular, the Bayesian posterior mean has the same asymptotic distribution as does the ordinary maximum likelihood estimator.

- the intuition is clear: as the amount of information coming from the sample increases, the
 likelihood function brings an increasing amount of information, relative to the prior. Eventually,
 the prior is no longer important for determining the shape of the posterior.
- when the sample is large, the shape of the posterior depends on the likelihood function. The likelihood function collapses around θ_0 when the sample is generated at θ_0 . The same is true of the posterior, it narrows around θ_0 . This causes the posterior mean to converge to the true parameter value. In fact, all quantiles of the posterior converge to θ_0 . Chernozhukov and Hong discuss estimators defined using quantiles.
- For an econometrician coming from the frequentist perspective, this is attractive. The
 Bayesian estimator has the same asymptotic behavior as the MLE. There may be
 computational advantages to using the Bayesian approach, because there is no need for

optimization. If the objective function that defines the classical estimator is irregular (multiple local optima, nondifferentiabilities, noncontinuities...), then optimization may be very difficult. However, Bayesian methods that use integration may be more tractable. This is the main

motivation of CH's paper. Additional advantages include the benefits if an informative prior is available. When this is the case, the Bayesian estimator can have better small sample performance than the maximum likelihood estimator.

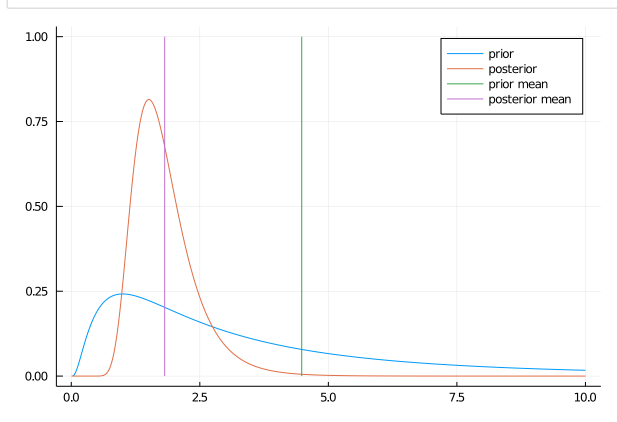
Example (Exponential Distribution)

Suppose data is generated by i.i.d. sampling from an exponential distribution with mean θ . An exponential random variable takes values on the positive real numbers. Waiting times are often modeled using the exponential distribution.

- The density of a typical sample element is $f(y|\theta) = \frac{1}{\theta}e^{-y/\theta}$. The likelihood is simply the product of the sample contributions.
- Suppose the prior for θ is $\theta \sim \text{lognormal}(1,1)$. This means that the logarithm of θ is standard normal. We use a lognormal prior because it enforces the requirement that the parameter of the exponential density be positive.

```
In [2]:
          1 # the prior is Lognormal(1,1)
             function prior(theta)
          2
          3
                 d = LogNormal(1.0, 1.0)
                 p = pdf.(Ref(d), theta)
          4
          5
                 pmean = exp(1.5) # mean of lognormal is exp(mu+sig/2)
          6
                 return p, pmean
          7
             end
          8
          9
             # the likelihood function
            function likelihood(y, theta)
         10
                 dens = zeros(size(theta))
         11
                 for i = 1:size(theta,1)
         12
         13
                     d = Exponential(theta[i])
                     dens[i] = prod(pdf.(Ref(d), y))
         14
         15
                 end
                 return dens
         16
         17
            end
         18
         19
         20 | # joint is prior X likelihood
         21 function joint(y, theta)
                 1 = likelihood(y, theta)
         22
         23
                 p, junk = prior(theta)
         24
                 dens = 1.*p
         25 end
         26
             # compute marginal likelihood of Y by integrating out theta (crude, only ill
         27
            function marginal(y)
         28
         29
                 dens = 0.0
         30
                 theta = 0.0
         31
                 delta = 0.01
         32
                 # evaluate joint over grid
         33
                 for r = 1:1000
         34
                     theta += delta
         35
                     dens += joint(y, theta)
         36
                 end
         37
                 # marginalize by integrating the joint (sum up height X width)
                 dens = dens*delta
         38
         39
            end
         40
         41
            # the posterior, by Bayes' Law
            function posterior(y, theta)
         42
         43
                 m = marginal(y)
                 j = joint(y, theta)
         44
         45
                 dens = j ./ m
         46
                 thetas = range(0.01, stop=10, length=1000)
         47
                 pmean = sum(dens.*thetas.*0.01)
         48
                 return dens, pmean
         49 end
         50
            ;
```

Out[4]:



```
In [3]:
          1 n = 50 # sample size
          2 | theta = 3 # true theta
          3 y = rand(Exponential(theta), n) # sample from exponential(theta)
          4 # make plots
          5 thetas = range(0.01,stop=10,length=1000)
          6 p, priormean = prior(thetas)
          7
             post, postmean = posterior(y, thetas)
          8 plot(thetas, [p post], label = ["prior" "posterior"])
             plot!([priormean, priormean], [0.0, 1.0], label = "prior mean")
         plot!([postmean, postmean], [0.0, 1.0], label = "posterior mean")
Out[3]:
          1.00
                                                                         prior
                                                                         posterior
                                                                         prior mean
                                                                         posterior mean
          0.75
          0.50
          0.25
```

Note how the posterior is more concentrated around the true parameter value when sample has size 50. Also note how the posterior mean is closer to the prior mean when the sample is small. When the sample is small, the likelihood function has less weight, and more of the information comes from the prior. When the sample is larger, the likelihood function will have more weight, and its effect will dominate the prior's.

Markov Chain Monte Carlo (MCMC)

To compute the posterior mean, we need to evaluate

$$E(\theta|y) = \int_{\Theta} \theta f(\theta|y) d\theta$$
$$= \frac{\int_{\Theta} \theta f(y|\theta) \pi(\theta) d\theta}{\int_{\Theta} f(y,\theta) d\theta}.$$

- Note that both of the integrals are multiple integrals, with the dimension given by that of the parameter, θ .
- Under some special circumstances, the integrals may have analytic solutions: e.g., Gaussian likelihood with a Gaussian prior leads to a Gaussian posterior.
- When the dimension of the parameter is low, quadrature methods may be used. What we have done with marginal(y) and posterior(y, theta) is an unsophisticated example of this. More

- sophisticated methods use an intelligently chosen grid to reduce the number of function evaluations. Still, these methods only work for dimensions up to 3 or so.
- Otherwise, some form of simulation-based "Monte Carlo" integration must be used. The basic idea is that $E(\theta|y)$ can be approximated by $(1/S)\sum_{s=1}^S \theta^s$, where θ^s is a random draw from the posterior distribution $f(\theta|y)$.

How to make draws from the posterior when in general we can't compute the posterior?

- the law of large numbers tells us that this average will converge to the desired expectation as
 S gets large
- convergence will be more rapid if the random draws are independent of one another, but insisting on independence may have computational drawbacks.

Monte Carlo methods include importance sampling, Markov chain Monte Carlo (MCMC) and sequential Monte Carlo (SMC, also known as particle filtering). The great expansion of these methods over the years has caused Bayesian econometrics to become much more widely used than it was in the not so distant (for some of us) past. There is much literature - here we will only look at a basic example that captures the main ideas.

Metropolis-Hastings

Let's consider the basic **Metropolis-Hastings MCMC** algorithm. We will generate a long realization of a Markov chain process for θ , as follows:

The prior density is $\pi(\theta)$, as above.

Let $g(\theta^*; \theta^s)$ be a proposal density, which describes the density of a trial value θ^* conditional on starting at θ^s . It must be possible to sample from the proposal. This gives a new trial parameter value θ^* , given the most recently accepted parameter value θ^s . A proposal will be accepted if

$$\frac{\dot{f}(\theta^*|y)}{f(\theta^s|y)}\frac{g(\theta^s;\theta^*)}{g(\theta^*;\theta^s)} > \alpha$$

where α is a U(0,1) random variate.

There are two parts to the numerator and denominator: the posterior, and the proposal density.

- The numerator, when the trial value of the proposal has a higher posterior, acceptance is favored.
- The density associated with returning to θ^s when starting at θ^* , which has to do with the reversibility of the Markov chain. If this is too low, acceptance is not favored. We don't want to jump to a new region if we will never get back, as we need to sample from the entire support of the posterior.

The two together mean that we will jump to a new area only if we are able to eventually jump back with a reasonably high probability. The probability of jumping is higher when the new area has a higher posterior density, but lower if it's hard to get back.

The idea is to sample from all regions of the posterior, those with high and low density, sampling more heavily from regions of high density. We want to go occasionally to regions of low density, but it is important not to get stuck there.

Note that the ratio of posteriors is equal to the ratio of likelihoods times the ratio of priors:

$$\frac{f(\theta^*|y)}{f(\theta^s|y)} = \frac{f(y|\theta^*)}{f(y|\theta^s)} \frac{\pi(\theta^*)}{\pi(\theta^s)}$$

because the marginal likelihood f(y) is the same in both cases. We don't need to compute that integral! We don't need to know the posterior, either. The acceptance criterion can be written as: accept if

$$\frac{f(y|\theta^*)}{f(y|\theta^s)} \frac{\pi(\theta^*)}{\pi(\theta^s)} \frac{g(\theta^s; \theta^*)}{g(\theta^*; \theta^s)} > \alpha$$

otherwise, reject

From this, we see that the information needed to determine if a proposal is accepted or rejected is the prior, the proposal density, and the likelihood function $f(y|\theta)$.

- in principle, the prior is non-negotiable. In practice, people often chose priors with convenience in mind
- the likelihood function is what it is
- · the place where artistry comes to bear is the choice of the proposal density
- when the proposal density is symmetric, so that $g(\theta^s; \theta^*) = g(\theta^*; \theta^s)$, the acceptance criterion simplifies to

$$\frac{f(y|\theta^*)}{f(y|\theta^s)} \frac{\pi(\theta^*)}{\pi(\theta^s)} > \alpha$$

A random walk proposal, where the trial value is the current value plus a shock that doesn't depend on the current value, satisfies symmetry.

The steps are:

- 1. the algorithm is initialized at some θ^1
- 2. for s = 2, ..., S,
 - A. draw θ^* from $g(\theta^*; \theta^s)$
 - B. according to the acceptance/rejection criterion, if the result is acceptance, set $\theta^{s+1} = \theta^*$, otherwise set $\theta^{s+1} = \theta^s$
 - C. iterate

Once the chain is considered to have stabilized, say at iteration r, the values of θ^s for s > r are taken to be draws from the posterior. The posterior mean is computed as the simple average of the value. Quantiles, etc., can be computed in the appropriate fashion.

The art of applying these methods consists of providing a good proposal density so that the acceptance rate is reasonably high, but not too high. There is a vast literature on this, and the vastness of the literature should serve as a warning that getting this to work in practice is not necessarily a simple matter. If it were, there would be fewer papers on the topic.

- too high acceptance rate: this is usually due to a proposal density that gives proposals very
 close to the current value, e.g, a random walk with very low variance. This means that the
 posterior is being explored inefficiently, we travel around through the support at a very low
 rate, which means the chain will have to run for a (very, very...) long time to do a thorough
 exploration.
- too low acceptance rate: this means that the steps are too large, and we attempt to move to
 low posterior density regions too frequently. The chain will become highly autocorrelated, as it
 stays in the same place due to rejections, so long periods convey little additional information
 relative to a subset of the values in the interval

Example

The simple exponential example with log-normal prior can be implemented using MH MCMC

```
In [6]:
                function mcmc(θ, reps::Int64, burnin::Int64, Prior::Function, lnL::Function,
            1
             2
                     lnL\theta = lnL(\theta)
                     chain = zeros(reps, size(\theta,1)+1)
            3
                     for rep = 1:burnin+reps
            4
            5
                          \theta^{t} = Proposal(\theta) # new trial value
                          pt = Prior(\theta^t)
            6
            7
                          accept = false
            8
                          if pt > 0.0
            9
                               lnL\theta^{t} = lnL(\theta^{t})
                               accept = rand() < exp(lnL\theta^t - lnL\theta) * pt/Prior(\theta)
           10
                               if accept
           11
           12
                                    \theta = \theta^t
                                    lnL\theta = lnL\theta^t
           13
           14
                               end
           15
                          end
           16
                          if rep > burnin
           17
                               chain[rep-burnin,:] = [\theta; accept]
           18
                          end
           19
                     end
           20
                     return chain
           21
               end
```

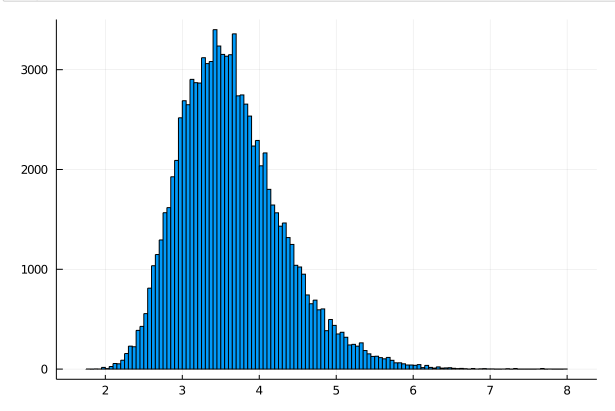
Out[6]: mcmc (generic function with 1 method)

```
In [10]:  # sample is from exponential, prior is lognormal, proposal is random walk to
2 y = rand(Exponential(3.0),30)
3 # set prior, likelihood and proposal
4 Prior = θ -> pdf(LogNormal(1.0,1.0), θ)
5 lnL = θ -> sum(logpdf.(Ref(Exponential(θ)),y))
6 tuning = 0.75
7 Proposal = θ -> rand(LogNormal(log(θ),tuning))
8 # get the chain, plot posterior, and descriptive stats
9 chain = mcmc(1.0, 100000, 10000, Prior, lnL, Proposal); # start value, chain
10
```

```
In [11]:
              chain
Out[11]: 100000×2 Array{Float64,2}:
           2.98767
                    0.0
           5.24983
                    1.0
           5.24983
                    0.0
           5.24983
                    0.0
           5.24983
                    0.0
           5.22463
                    1.0
           5.22463
                    0.0
           5.22463
                    0.0
           5.22463
                    0.0
           5.22463
                    0.0
           5.22463
                    0.0
           5.22463
                    0.0
           5.22463
                    0.0
           3.41251
                    1.0
           3.40802
                    1.0
           3.40802
                    0.0
           3.40802
                    0.0
In [12]:
           1 mean(chain,dims=1)
Out[12]: 1x2 Array{Float64,2}:
           3.63477 0.27977
In [13]:
              plot(chain[end-500:end,1])
Out[13]:
           7
                                                                                   у1
           6
           5
           4
```

In [14]: 1 histogram(chain[:,1],legend=false)

Out[14]:



Now let's graph:

```
In [87]:
           1
              function npdensity(z)
                  #pyplot()
           2
           3
                  n = size(z,2)
           4
                  p = zeros(n)
           5
                  for i = 1:n
           6
                      x = z[:,i]
           7
                      y = kde(x)
           8
                      q05 = quantile(x, 0.05)
           9
                      Plots.plot(range(minimum(x), stop=q05, length=100),z->pdf(y,z), colo
          10
                      q95 = quantile(x, 0.95)
          11
                      Plots.plot!(range(q05, stop=q95, length=100),z->pdf(y,z), color=:gre
          12
                      Plots.plot!(range(q95, stop=maximum(x), length=100),z->pdf(y,z), col
                      m = mean(x)
          13
                      Plots.plot!([m,m],[0,pdf(y,m)],color=:blue, label="mean")
          14
          15
                      m = median(x)
          16
                      if i == 1
          17
                           p = Plots.plot!([m,m],[0,pdf(y,m)],color=:black, label="median")
          18
                      else
          19
                           p = [p, Plots.plot!([m,m],[0,pdf(y,m)],color=:black, label="medi")
          20
                      end
          21
                  end
          22
                  return p
          23
              end
```

Out[87]: npdensity (generic function with 1 method)

Gibbs Samling

The algorithm:

```
0. Assign starting values for \theta with size k: \theta^0
```

```
1. Set j = j + 1
```

2. Sample
$$(\theta_1^j | \theta_2^{j-1}, \dots, \theta_k^{j-1})$$

2. Sample
$$(\theta_2^j | \theta_1^{j-1}, \dots, \theta_k^{j-1})$$

:

k+1. Sample
$$(\theta_k^j | \theta_2^{j-1}, \dots, \theta_{k-1}^{j-1})$$

k+2. Return to step 1.

Gibbs sampling is a special case of Metropolis-Hastings algorithm where the acceptance rate is equal to 1.

In other words, Gibbs sampling involves ordering the parameters and sampling from the conditional distribution for each parameter given the current value of all the other parameters and repeatedly cycling through this updating process.

For Gibbs sampling, the full conditional density for a parameter needs only to be known up to a normalizing constant. This implies that we can use the joint density with the other parameters set at their current values. This fact makes Gibbs sampling relatively simple for most problems in which the joint density reduces to known forms for each parameter once all other parameters are treated as fixed.

Let's construct a **Gibbs sampler** for the bivariate distribution:

$$f(x, y) = kx^{2}e^{-xy^{2}-y^{2}+2y-4x}$$

where x > 0, $y \in \mathcal{R}$ with unknown normalizing constant k > 0 with unknown normalizing constant k > 0.

You can think of (x, y) as previously noted θ .

The statistical motivation is not important for this post, but this is the kind of distribution which arises naturally in Bayesian inference for the mean and variance of a normal random sample. Gibbs sampling is a simple MCMC scheme which samples in turn from the full-conditional distributions.

In this case, then, are:

$$x|y \sim Gamma(3, y^2 + 4)$$
$$y|x \sim N\left(\frac{1}{1+x}, \frac{1}{\sqrt{2(1+x)}}\right)$$

Given the full conditionals, alternately sample from them to construct a Gibbs sampler for the target distribution. We will run a Gibbs sampler with a thin of 1000 and obtain a final sample of 50000.

```
In [17]:
              function Gibbs(reps::Int, burnin::Int)
                   chain = fill(NaN, (reps,2))
           3
                       = 0.
           4
                       = 0.
           5
                   for i = 1:reps
           6
                       for j = 1:burnin
           7
                           x = rand(Gamma(3,(y*y + 4)))
                           y = rand(Normal(1/(x+1), 1/sqrt(2*(x + 1))))
           8
           9
          10
                       chain[i,:] = [x,y]
          11
                   end
          12
                   return chain
          13
              end
```

Out[17]: Gibbs (generic function with 1 method)

```
In [18]:
               chain = Gibbs(100000,1000)
            8.05318
                      -0.139629
            7.86046
                       0.537612
            7.25506
                       0.506774
            5.22541
                      -0.102717
            7.58817
                      -0.237087
            9.44253
                       0.185475
                      -0.0363721
            3.63053
            6.59441
                      -0.0988208
           10.7533
                      -0.0555952
           30.5729
                      -0.00567513
            5.0229
                       0.137309
            2.70492
                       1.06705
           23.0421
                      -0.286004
            7.18185
                      -0.128605
            7.10599
                       0.048754
            1.33589
                       0.223614
           13.0347
                       0.266302
           13.802
                      -0.147551
In [19]:
               plot(chain,layout=2,seriestype=[:histogram],legend=false,title=["x" "y"])
Out[19]:
                                 Χ
                                                                           У
                                                      4000
            3000
                                                      3000
            2000
                                                      2000
            1000
                                                      1000
              0
                                                        0
                        20
                               40
                                                                         0
                                     60
                                            80
In [21]:
              mean(chain,dims=1)
Out[21]: 1x2 Array{Float64,2}:
```

Convergence

12.2124 0.101961

Assume that we have a Markov chain X_t generator with a help of Metropolis-Hastings algorithm (Gibb's sampling is a special case of it). The starting point is X_0 . We are interested in convergence of the chain to its invariant distribution π . In other words, we expect: $P[X_t \in A|X_0]$ to converge to π as $t \to \infty$.

If all values are visited with strictly positive probability and no partition that has strictly 1 probability, then:

$$|P[X_t \in A | X_0 = x] - \pi(A)|$$
 ast $\to \infty$

To detect convergence in practice one may do the following:

- check whether some characteristics of the chain distribution (such as mean, variance, median, quartiles) are stabilize with time for a single long run;
- compare a ratios of a current estimate of a target density and a target density itself (known up
 to a constant) for a single long run of a chain;
- compare some characteristics of the chain distribution across several independent runs of a chain.

General practical recommendations:

- There is no unique way to implement MCMC. In majority of cases you have many choices, try
 several of them. It relates to choosing between different variants of M-H and Gibbs sampling
 as well as to picking up sampling density q. For better results q should be appropriately
 centered and have tails dominating π.
- I've seen a recommendation to tune your M-H algorithm to roughly 30% acceptance rate. I still
 don't know where the number comes from... The general wisdom is: if your acceptance rate is
 too low, then you are producing very dependent draws; if your acceptance rate is too high, it
 may be the sign that you are visiting only the high probability area and not moving too far from
 it.
- You should be very careful with choosing "non-informative" priors, since many of them tend to be improper and may produce improper posteriors (ooo-u-ps!)