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- Mild conditions guarantee the convergence of the algorithm but not the convergence rate!
- We have looked at two special proposal densities:
 - ▶ The independence proposal Q(y|x) = Q(y)
 - ▶ The RW proposal Q(y|x) = Q(x|y)
- Importance of the tuning parameter

Review: Special cases Metropolis-Hastings

 Metropolis algorithm: The proposal density is symmetric around the current value, that means

$$Q(x_{i-1}|y) = Q(y|x_{i-1}).$$

Hence,
$$\alpha = \min\left(1, \frac{\pi(y)}{\pi(x_{i-1})} \times \frac{Q(x_{i-1}|y)}{Q(y|x_{i-1})}\right) = \min\left(1, \frac{\pi(y)}{\pi(x_{i-1})}\right)$$

Acceptance rates should be between 20% and 50%.

 Independence sampler: The proposal distribution does not depend on the current value x_{i-1}

$$Q(x|x_{i-1})=Q(x).$$

Q(x) is an approximation to $\pi(x) \Rightarrow$ acceptance rate should be high.

MCMC and iterative conditioning

The MH-algorithm can be applied iteratively on the different elements of a stochastic vector x conditioning on all other elements.

To be concrete, one uses

- a proposal kernel $Q(y^j|x_{i-1}^j, \pmb{x}_{i-1}^{-j}), j=1,\ldots,p$.
- with acceptance probability

$$\alpha = \min\left(1, \frac{\pi(y^j|\mathbf{x}_{i-1}^{-j})}{\pi(x_{i-1}^j|\mathbf{x}_{i-1}^{-j})} \times \frac{Q(x_{i-1}^j|y^j, \mathbf{x}_{i-1}^{-j})}{Q(y^j|x_{i-1}^j, \mathbf{x}_{i-1}^{-j})}\right)$$

This algorithm converges to the stationary distribution with density $\pi(x)$, as long as all components are arbitrary often updated.

Conditional densities

The full conditional densities $\pi(x^j|\mathbf{x}^{-j})$, $j=1,\ldots,p$, and the joint density $\pi(\mathbf{x})$ are related as follows:

$$\pi(\mathbf{x}^{j}|\mathbf{x}^{-j}) = \frac{\pi(\mathbf{x})}{\pi(\mathbf{x}^{-j})} \propto \pi(\mathbf{x})$$

Thus, the (non-normalised) conditional densities of $x^j | x^{-j}$ can be directly derived from $\pi(x)$ by omitting all multiplicative factors, that do not depend on x^j .

Gibbs sampling

Idea: Sequentially sampling from univariate conditional distributions (which are often available in closed form).

- 1. Select starting values x_0 and set i = 0.
- 2. Repeatedly:

$$\begin{array}{ll} \text{Sample} & x_{i+1}^1| \cdot \sim \pi(x^1|x_i^2, \dots, x_i^p) \\ \\ \text{Sample} & x_{i+1}^2| \cdot \sim \pi(x^2|x_{i+1}^1, x_i^3, \dots, x_i^p) \\ \\ \vdots \\ \\ \text{Sample} & x_{i+1}^{p-1}| \cdot \sim \pi(x^{p-1}|x_{i+1}^1, x_{i+1}^2, \dots, x_{i+1}^{p-2}, x_i^p) \\ \\ \text{Sample} & x_{i+1}^p| \cdot \sim \pi(x^p|x_{i+1}^1, \dots, x_{i+1}^{p-1}) \end{array}$$

where $|\cdot|$ denotes conditioning on the most recent updates of all other elements of x.

3. Increment i and go to step 2. TMA4300 - Part 2

Gibbs sampling

The Gibbs sampler is a special case of the MH-algorithm.

The acceptance probability is always $\boldsymbol{1}$

Plan for today

- Gibbs sampling
- Blocking strategies
- Convergence diagnostic

Example: Simple linear regression

Let

$$Y_i = a + bx_i + e_i, \ e_i \sim \mathcal{N}(0, 1/\tau), \ i = 1, ..., n$$

and

$$a \sim \mathcal{N}(0, 1/ au_a)$$

$$b \sim \mathcal{N}(0, 1/\tau_b)$$

$$\tau \sim \mathsf{Gamma}(\alpha, \beta)$$

we are interested in the posterior:

$$\pi(a, b, \tau|y)$$

(See notes, show R-code demo_linear_regression.R)

Remarks on Gibbs sampling

- High dimensional updates of x can be boiled down to scalar updates.
- Visiting schedule: Various approaches exist (and can be justified) to ordering the variables in the sampling loop. One approach is random sweeps: variables are chosen at random to resample.
- Gibbs sampling assumes that it is easy to sample from the full-conditional distribution. This is sometimes not so easy.
 Alternatively, a Metropolis-Hastings proposal can be used for the *j*-th component, i.e. Metropolis-within-Gibbs ⇒ Hybrid Gibbs sampler.

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Remarks on Gibbs sampling

- Blocking or grouping is possible, that means not all elements
 of x are treated individually. Might be useful when elements of
 x are correlated.
- Care must be taken when improper prior are used, which may lead to an improper posterior distribution. Impropriety implies that there does not exist a joint density to which the full-conditional distributions correspond.

Hobert, J. P. and Casella, G. (1996), JASA, 91: 1461-1473.

Example: Conjugate gamma-Poisson hierarchical model

Example from George et al. (1993) regarding the analysis of 10 power plants.

- yi number of failures of pump i
- t_i length of operation time of pump i (in kilo hours)

Model:

$$y_i \mid \lambda_i \sim Po(\lambda_i t_i)$$

Conjugate prior for λ_i :

$$\lambda_i \mid \alpha, \beta \sim \mathsf{G}(\alpha, \beta)$$

Hyper-prior on α and β :

$$\alpha \sim \text{Exp}(1.0)$$

$$\beta \sim G(0.1, 10.0)$$

Conjugate gamma-Poisson hierarchical model (II)

The posterior of the 12 parameters $(\alpha, \beta, \lambda_1, \dots, \lambda_{10})$ given y_1, \dots, y_{10} is proportional to

$$\pi(\alpha, \beta, \lambda_1, \dots, \lambda_{10} \mid y_1, \dots, y_{10}) \propto \pi(\alpha)\pi(\beta) \prod_{i=1}^{10} [\pi(\lambda_i \mid \alpha, \beta)\pi(y_i \mid \lambda_i)]$$

$$\propto e^{-\alpha}\beta^{0.1-1}e^{-10\beta} \left\{ \prod_{i=1}^{10} \exp(-\lambda_i t_i)\lambda_i^{y_i} \right\} \left\{ \prod_{i=1}^{10} \exp(-\beta\lambda_i)\lambda_i^{\alpha-1} \right\} \left[\frac{\beta^{\alpha}}{\Gamma(\alpha)} \right]^{10}.$$

This posterior is not of closed form.

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This posterior is not of closed form.

Work out the full conditional distributions yourself!

Update scheme for gamma-Poisson hierarchical model

For each iteration i

- For k = 1, ..., 10
 - ▶ Simulate new value $\lambda_k \sim \mathsf{Gamma}(y_i + \alpha, t_i + \beta)$ Gibbs step
- Simulate new value $eta \sim {\sf Gamma} (10 \alpha + 0.1, \sum \lambda_k + 1)$ Gibbs step
- Propose new value $\alpha_{new} \sim \mathcal{N}(\alpha_{i-1}, \tau)$ MH step
- Compute acceptance probability

$$a = \min \left\{ 1, \frac{\pi(\alpha_{new}|\dots)}{\pi(\alpha_{old}|\dots)} \right\}$$

- if *u* < *a*
 - ightharpoonup set $\alpha_i = \alpha_{new}$
- else
 - ightharpoonup set $\alpha_i = \alpha_{old}$

Blocking Strategies

Blocking (ie simulating some variables together) might improve the algorithm especially when variables are correlated.

Example: Korsbetningen

In the year of our Lord 1361, on the third day after S:t Jacob, the Goth fell outside the gates of Visby at the hands of the Danish. They are buried here. Pray for them.

- Archeoloical excavation found 493 femurs, 256 right and 237 left
- At least 256 person were buried here....but how many more??

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A simple model

Let x_1 and x_2 be the number of left and right femurs found.

Assume x_1 and x_2 to be two independent observations from a $Bin(N, \phi)$ distribution.

With

- N total number of people buried
- ullet ϕ probability of finding a femur, left or right

The unkown parameter vector is $\theta = (N, \phi)$. Assume a Beta(a, b) prior for ϕ , and a Unif(256, 2500) prior for N.

Updating schemes

Single site update

- Simulate new $\phi \sim \mathsf{Beta}(\cdot, \cdot)$ (Gibbs step)
- Propose $N_{new} \sim$ Unif $(N_{old} d, N_{old} + d)$
- Compute

$$\alpha = \min \left\{ 1, \frac{\pi(\textit{N}_{\textit{new}}|\dots)}{\pi(\textit{N}_{\textit{old}}|\dots)} \right\}$$

 Accept or reject the new value for N

Block update

- Propose a new value N_{new} for N from
 Unif(N_{old} d, N_{old} + d)
- Propose a new value $\phi_{\textit{new}}$ for ϕ from

$$Beta(\alpha+x_1+x+2,\beta+2N_{new}-x_1-x_2)$$

- Compute α
- Accept or reject N_{new} and ϕ_{new} simultaneously (Show R-code Vikings.R)

Implementation and convergence diagnostics



Source: http://i.telegraph.co.uk/multimedia/archive/02365/coding_alamy_2365972b.jpg

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Convergence

- If well constructed, the Markov chain is guaranteed to have the posterior as limiting distribution.
- However, this does not tell you how long you have to run the MCMC algorithm til convergence.
 - ► The initial position may have a big influence.
 - ▶ The proposal distribution may lead to low acceptance rates.
 - The chain may get caught in a local maximum of the likelihood surface.
- We say the Markov chain mixes well if it can
 - reach the posterior quickly, and
 - moves quickly around the posterior modes.

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Convergence diagnostics

Valid inferences from sequences of MCMC outputs are based on the assumption that the outputs are from the desired target distribution.

- There is no overall minimum number of samples to ensure approximation.
- Consequently methods for testing convergence, known as convergence diagnostics, have to be applied.
- However it has to emphasised that these diagnostics do not guarantee convergence.

Trace plots

An initial possibility for deciding if a MCMC output does not converge to the desired posterior distributions is to look at the sample trace for each variable.

 If our chain is taking a long time to move around the parameter space, then it will take longer to converge.

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Trace plots

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- If the samples form a homogeneous band (no wave movements or other rare fluctuations), convergence might be indicated.

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- If our chain is taking a long time to move around the parameter space, then it will take longer to converge.
- If the samples form a homogeneous band (no wave movements or other rare fluctuations), convergence might be indicated.
- Vastly different values at the beginning of the trace indicate burn-in iterations, which should be discarded.

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Burn-in

In practice, one waits until the Markov chain is converged. Let K denote the burn-in period. Then the realisations

 $\mathbf{x}_{K+1}, \mathbf{x}_{K+2}, \dots, \mathbf{x}_{K+N}$ are used to estimate characteristics of the target distribution.

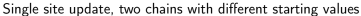
The empirical determination of K is difficult. Often it is determined based on the trace plot of the Markov chain.

Output analysis

Standard starting point to evaluate convergence:

- Look at the trace plot for each variable
- consider different scalar function of x
- may run different Markov chain with different (extreme) starting values

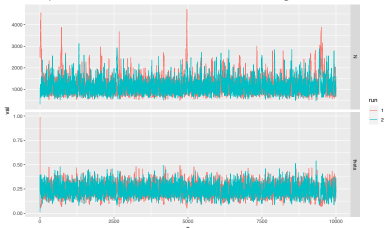
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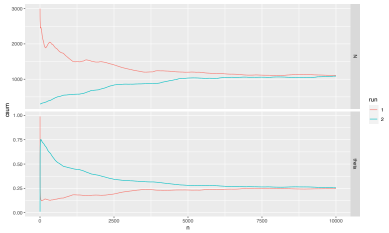
Block update, two chains with different starting values



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Single site update, two chains with different starting values.

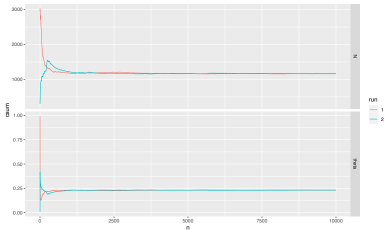
Estimate of the mean



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Block update, two chains with different starting values.

Estiamte of the mean



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Convergence Diagnostic

With a fixed cpu-time shoud we:

- use all time in one long Markov chain, or
- run several shorter Markov chains?

Convergence Diagnostic

With a fixed cpu-time shoud we:

- use all time in one long Markov chain, or
- run several shorter Markov chains?
- One long chain:
 - only one bunr-in period to discard
 - more likely that you really have converged
- Several shorter runs:
 - easier to evaluate convergence
 - easier to estimate the variance of the estimator (the chains are independent)

In practice one often use a combination of the two strategies

Variance of the MCMC estimator

Recall: We want to estimate $\mu = \int g(x)\pi(x) dx$ with

$$\hat{\mu} = \frac{1}{n} \sum g(x_i)$$
 where $x_i \sim \pi(x)$.

In standard MC we have

$$x_1, x_2, ..., x_n \sim \pi(x)$$
, i.i.d.

This gives

$$\mathsf{E}(\hat{\mu}) = \mu \text{ and } \mathsf{Var}(\hat{\mu}) = \frac{\mathsf{Var}(g(X))}{n}$$

We can estimate the variance $Var(\hat{\mu})$ as

$$\widehat{\operatorname{Var}(\widehat{\mu})} = \frac{\widehat{\operatorname{Var}(g(X))}}{n}$$

$$\widehat{\operatorname{Var}(g(X))} = \frac{1}{n-1} \sum_{i=1}^{n} (g(x_i) - \widehat{\mu})^2$$

MCMC gives dependence samples, what is the variance then??

Autocorrelation

To examine dependencies of successive MCMC samples, the autocorrelation function can be used. Let x_1, \ldots, x_N , where N denotes the number of samples, denote our MCMC chain.

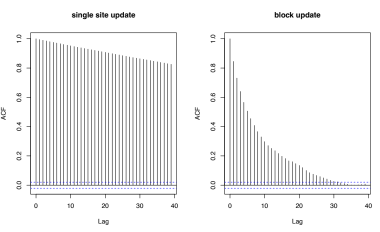
The lag k autocorrelation $\rho(k)$ is the correlation between every draw and its k-th lag. For N reasonably large

$$\rho(k) \approx \frac{\sum_{i=1}^{N-k} (x_i - \bar{x})(x_{i+k} - \bar{x})}{\sum_{i=1}^{N} (x_i - \bar{x})^2},$$

where $\bar{x} = \frac{1}{N} \sum_{i=1}^{N} x_i$ is the overall mean.

- With increasing lag k we expect lower autocorrelations.
- If autocorrelation is still relatively high for higher values of k, this indicates high degree of correlation between our draws and slow mixing.

Autocorrelation function for N (after discarding the burn-in period)



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Effective sample size

A useful measure to compare the performance of different MCMC samplers is the effective sample size (ESS) Kass et al. (1998) American Statistician 52, 93–100..

 The ESS is the estimated number of independent samples needed to obtain a parameter estimate with the same precision as the MCMC estimate based on N dependent samples.

$$\mathsf{ESS} = \frac{N}{\tau}, \quad \tau = 1 + 2 \cdot \sum_{k=1}^{\infty} \rho(k),$$

where τ is the autocorrelation time and $\rho(k)$ the autocorrelation at lag k.

Estimate of ESS

$$\mathsf{ESS} = \frac{N}{\tau}, \quad \tau = 1 + 2 \cdot \sum_{k=1}^{\infty} \rho(k),$$

Estimate τ as

$$\tau = 1 + 2 \cdot \sum_{k=1}^{m} \hat{\rho}(k)$$

where $\hat{\rho}(k)$ is the sample autocorrelation function at lag k, and m is choosen to fullfill some criteria.

Different criteria exists.

Example: Korsbetningen - Effective sample size (ESS)

```
> library(coda)
> nsamples
[1] 8000
> ## single site
> effectiveSize(as.mcmc(res1))
       N
            theta
20.28367 16.30797
> ## block update
> effectiveSize(as.mcmc(res2))
       N
            theta
506,6220 845,4659
```

The precision of the MCMC estimate of the posterior mean of *N* based on 8000 samples from a single site update is a good as taking 16 independent samples!

Geweke diagnostics

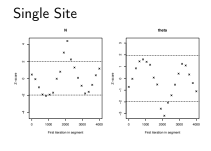
The MCMC chain is divided into two windows

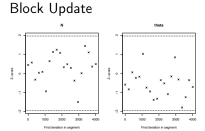
- the first x%, and
- the last y% of the iterates

(coda default: x = 10, y = 50). For both windows the mean is calculated.

If the chain is stationary both values should be equal and Geweke's test statistic (z-score) follows an asymptotical standard normal distribution.

Example: Korsbetningen - Geweke plot





Further reading

There are several convergence diagnostics:

- some are based on a single Markov chain run
- some are based on several Markov chain runs

There are no guarantees!

For further reading see for example

Gilks, W. R., Richardson, S. and Spiegelhalter, D.J. (1996)
 Markov Chain Monte Carlo in Practice, Chapman & Hall,
 London,

Different approaches are implemented in the

• R-package coda.

(Plummer et al., 2006)