Lecture 9: Brief Reminder

We are learning about the MCMC algorithm:

• What it is and why does it work

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- Elements of the algorithm:
 - ▶ Target distribution $\pi(x)$ Problem determined
 - Proposal distribution Q(y|x) Choosen by us
 - Acceptance probability $\alpha(y|x)$ Computed s.t. the detailed balance holds
- Mild conditions guarantee the convergence of the algorithm but not the convergence rate!

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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.

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Conditional densities

The full conditional densities $\pi(\mathbf{x}^j|\mathbf{x}^{-j})$, $j=1,\ldots,p$, 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 $x^j | \mathbf{x}^{-j}$ can be directly derived from $\pi(\mathbf{x})$ by omitting all multiplicative factors, that do not depend on x^j .

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 $Q(y^{j}|x_{i-1}^{j}, x_{i-1}^{-j}), j = 1, ..., p.$
- 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.

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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:

Sample
$$x_{i+1}^1| \cdot \sim \pi(x^1|x_i^2, \dots, x_i^p)$$

Sample $x_{i+1}^2| \cdot \sim \pi(x^2|x_{i+1}^1, x_i^3, \dots, x_i^p)$
:
Sample $x_{i+1}^{p-1}| \cdot \sim \pi(x^{p-1}|x_{i+1}^1, x_{i+1}^2, \dots, x_{i+1}^{p-2}, x_i^p)$
Sample $x_{i+1}^p| \cdot \sim \pi(x^p|x_{i+1}^1, \dots, x_{i+1}^{p-1})$

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

3. Increment *i* and go to step 2.
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Gibbs sampling

Plan for today

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

The acceptance probability is always 1

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Example: Simple linear regression

Let

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

and

$$a \sim \mathcal{N}(0, 1/ au_a)$$
 $b \sim \mathcal{N}(0, 1/ au_b)$ $au \sim \mathsf{Gamma}(lpha, eta)$

we are interested in the posterior:

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

(See notes, show R-code demo_linear_regression.R) TMA4300 - Part 2

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Gibbs sampling

- Blocking strategies
- Convergence diagnostic

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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 \Rightarrow 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.

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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.

Example: Conjugate gamma-Poisson hierarchical model

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

- y_i 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 :

$$\alpha \sim \mathsf{Exp}(1.0)$$
 $\beta \sim \mathsf{G}(0.1, 10.0)$

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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 $\lambda_k \sim \text{Gamma}(y_i + \alpha, t_i + \beta)$ Gibbs step

- Simulate new value $\beta \sim \mathsf{Gamma}(10\alpha + 0.1, \sum \lambda_k + 1)$ 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 $\alpha_i = \alpha_{old}$

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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
- \bullet 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.

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
- At least 256 person were buried here....but how many more??

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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

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

 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

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

- Compute α
- Accept or reject N_{new} and

 $\phi_{\it new}$ 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 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.

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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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.

Trace plots

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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.

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Burn-in

target distribution.

In practice, one waits until the Markov chain is converged. Let K denote the burn-in period. Then the realisations $x_{K+1}, x_{K+2}, \ldots, x_{K+N}$ are used to estimate characteristics of the

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

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.
- 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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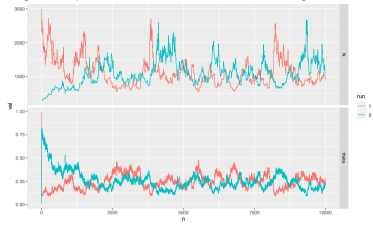
Output analysis

Standard starting point to evaluate convergence:

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

Example: Korsbetningen

Single site update, two chains with different starting values

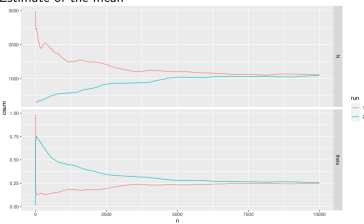


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Example: Korsbetningen

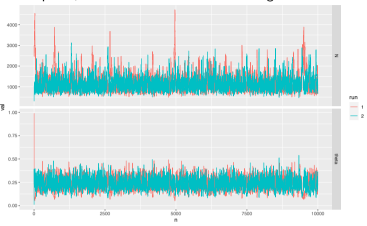
Single site update, two chains with different starting values.

Estimate of the mean



Example: Korsbetningen

Block update, two chains with different starting values

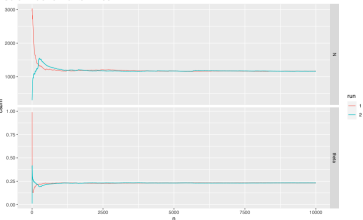


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Example: Korsbetningen

Block update, two chains with different starting values.

Estiamte of the mean



Convergence Diagnostic

With a fixed cpu-time shoud we:

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

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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

$$x_1, x_2, \dots, x_n \sim \pi(x)$$
, 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

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

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

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

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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?
- 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

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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

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

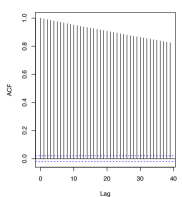
where $\bar{x} = \frac{1}{N} \sum_{i=1}^{N} x_i$ is the overall mean.

- 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.

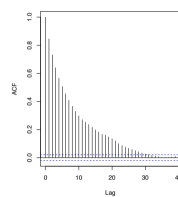
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Example: Korsbetningen

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



single site update



block update

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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.

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.

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Example: Korsbetningen - Effective sample size (ESS)

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.

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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,

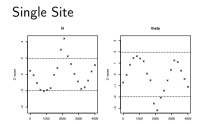
London,

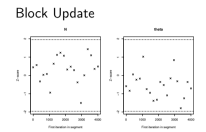
Different approaches are implemented in the

• R-package coda. (Plummer et al., 2006)

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Example: Korsbetningen - Geweke plot





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