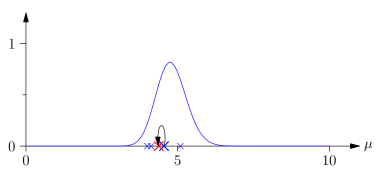
Advanced Machine Learning

MCMC

$$\mathcal{D} = \{4, 4, 6, 4, 2, 2, 5, 9, 5, 4, 3, 2, 5, 4, 4, 11, 6, 2, 3, 11\}$$



Monte Carlo methods, MCMC, Variational Methods

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1. Sampling

3. MCMC

2. Random Number Generation

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T = 10000000, acceptence rate = 0.897

Bayesian Inference Gets Hard

- We saw that in some cases if we had a simple likelihood (normal, binomial, Poisson, multinomial) you can choose a conjugate prior (gamma-normal/Wishart, beta, gamma, Dirchlet) so that the posterior has the same form as the prior
- Very often we are working with more complex models where no conjugate prior exists
- The posterior is not described by a known distribution
- We have to work a lot harder—particularly with multivariate distributions

Bayesian Inference

- Recall our problem is that we are given some data \mathcal{D}
- Our posterior is given by

$$\mathbb{P}(\boldsymbol{\theta}|\mathcal{D}) = \frac{\mathbb{P}(\mathcal{D}|\boldsymbol{\theta})\mathbb{P}(\boldsymbol{\theta})}{\mathbb{P}(\mathcal{D})} \qquad \text{or} \qquad f(\boldsymbol{\theta}|\mathcal{D}) = \frac{f(\mathcal{D}|\boldsymbol{\theta})f(\boldsymbol{\theta})}{f(\mathcal{D})}$$

- ullet Where heta are the parameters we are trying to infer
- But our likelihood (and/or prior) might be quite complicated
- Typically we don't have a closed form representation for our posterior distibution

Outline

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Histograms, Samples and Means

- We could represent our posterior as a histogram, although for multivariate distributions (i.e. when we are modelling more than one variable) a histogram can be unwieldy!
- A sample from the posterior distribution is often sufficient e.g. in our topic models (LDA) a typical set of topics is what we are after
- However, when samples vary a lot, often the most useful quantities are expectation, e.g.

$$\begin{split} \mathbb{E}[\boldsymbol{\Theta}] & & \mathbb{E}\left[\boldsymbol{\Theta}_i^2\right] - \mathbb{E}[\boldsymbol{\Theta}_i]^2 \\ \mathbb{E}[\boldsymbol{\Theta}_i \boldsymbol{\Theta}_j] - \mathbb{E}[\boldsymbol{\Theta}_i] \mathbb{E}[\boldsymbol{\Theta}_j] & & \mathbb{E}\left[\boldsymbol{\Theta} \boldsymbol{\Theta}^\mathsf{T}\right] - \mathbb{E}[\boldsymbol{\Theta}] \mathbb{E}[\boldsymbol{\Theta}]^\mathsf{T} \end{split}$$

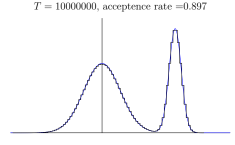
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Outline

- 1. Sampling
- 2. Random Number Generation
- 3. MCMC

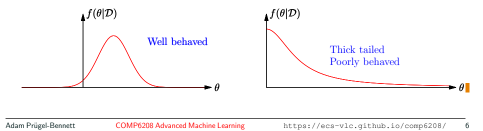


Sample Estimation

• If we can draw independent **deviates** (aka **variates**), Θ_i , from our posterior distribution then we can obtain an estimate of our expectation

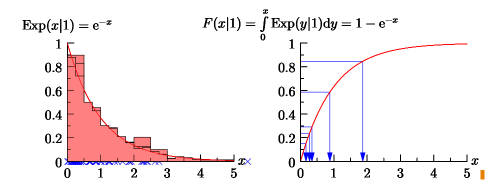
$$\mathbb{E}[g(\mathbf{\Theta})] \approx \frac{1}{n} \sum_{i=1}^{n} g(\mathbf{\Theta}_i) \mathbf{I}$$

• Provided our posterior distribution is well behaved the relative error in our estimate will drop off as $1/\sqrt{n}$



Drawing Random Samples

- Drawing (pseudo) random variables from a distribution is known as Monte Carlo
- For some very simple distributions we can use the transformation methods to transform a uniform distribution.



Rejection Method

- The transformation method only works when we can easily compute the inverse *cumulative distribution function* (CDF)
- A more general technique is the **rejection method** where we generate deviates from $g_Y(y)$ such that $cg_Y(x) \ge f_X(x)$
- To draw deviates from $f_X(x)$ we draw a deviate $Y \sim g_Y$ and then accept the deviate with probability $f_X(Y)/(cg_Y(Y))$
- The expected rejection rate is c-1
- Need to choose a good distribution $q_Y(y)$

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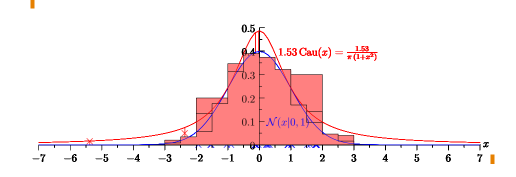
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Problems with Rejection

- The rejection method is very general and often the method of choice (although for normal deviates there is a clever transformation method which is faster)
- However, for complicated probability distributions it can be difficult to find a good proposal distribution $q_Y(y)$
- This is particular true for multivariate distributions
- \bullet If the proposal distribution is poor c might be very high and the number of rejections is stupidly high

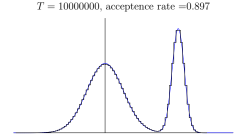
Drawing Normal Deviates



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Outline

- 1. Sampling
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Detailed Balance

- ullet Suppose we have a set of states ${\cal S}$ and want to draw sample from a probability distribution $\boldsymbol{\pi} = (\pi_i | i \in \mathcal{S})$
- ullet We invent a dynamical system with a transition probability M_{ij} from state i to state i such that

$$M_{ij}\pi_j = M_{ji}\pi_i$$

- This is known as detailed balance
- Summing both sides over *j*

$$\sum_{j} M_{ij} \pi_j = \sum_{j} M_{ji} \pi_i = \pi_i$$
 $\mathbf{M} \boldsymbol{\pi} = \boldsymbol{\pi}$

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• It I apply M many times then

maximum eigenvalue

• Now $\|\mathbf{M}\mathbf{v}\|_1 \leq \|\mathbf{M}\|_1 \|\mathbf{v}\|_1 = \|\mathbf{v}\|_1$ so the maximum eigenvalue is

• A very easy way to achieve detailed balance is starting from state j choose a "neighbouring" state, i with equal probability

Metropolis Algorithm

- We accept the move if either
 - $\star \pi_i > \pi_i$ or
 - \star we make the move with a probability π_i/π_i
- If $\pi_i > \pi_i$ then $M_{ij} = 1$ and $M_{ji} = \pi_j/\pi_i$. Thus

$$M_{ij}\pi_j = \pi_j$$
 $M_{ji}\pi_i = \frac{\pi_j}{\pi_i}\pi_i$ $= \pi_j$

• Note that we require the state i to have the same number of neighbours as state j so that detailed balance is satisfied

1 with eigenvector π (M is known as a **stochastic matrix**) Adam Prügel-Bennett COMP6208 Advanced Machine Learning

ullet And $\lim oldsymbol{x}(t) = oldsymbol{v}^*$ where $oldsymbol{v}^*$ is the eigenvector with the

Continuous Variables

ullet If we are working with continuous variables heta then the equation for detailed balance for the transition probability $W({m heta} o {m heta}')$ is

$$W(\boldsymbol{\theta} \to \boldsymbol{\theta}')\pi(\boldsymbol{\theta}) = W(\boldsymbol{\theta}' \to \boldsymbol{\theta})\pi(\boldsymbol{\theta}')$$

- where $\pi(\theta)$ is the probability distribution we wish to sample from
- The update rule is to choose a nearby value θ' , compute $r = \pi(\theta')/\pi(\theta)$ and accept the update with probability $\min(1,r)$
- We require that the probability of choosing θ from θ' is the same as the reverse

• Suppose we start from a state $x(0) = \sum_i c_i v^{(i)}$ where the $v^{(i)}$'s

are an eigenvectors of the transition matrix M with eigenvalues λ_i

 $oldsymbol{x}(t) = oldsymbol{\mathsf{M}}^t oldsymbol{x}(0)$ | $oldsymbol{\mathsf{M}}^t \sum_i c_i oldsymbol{v}^{(i)}$ | $oldsymbol{\mathsf{L}} = \sum_i \lambda_i^t c_i oldsymbol{v}^{(i)}$ |

What Makes MCMC Nice

What Makes MCMC Nasty

• It can take a long time until our states occur with the probability

- Because we are free to choose where we move (and choose close by neighbours) $\pi(\theta') \approx \pi(\theta)$ so that moves are not too infrequent
- Also very importantly the updates depend only on the ratio $\pi(\boldsymbol{\theta}')/\pi(\boldsymbol{\theta})$
- We only need to know our probabilities up to a multiplicative scaling factor
- For sampling from the posterior we only need to know the likelihood and prior $\mathbb{P}(\mathcal{D}|\boldsymbol{\theta})\mathbb{P}(\boldsymbol{\theta})$ (or $f(\mathcal{D}|\boldsymbol{\theta})f(\boldsymbol{\theta})$)
- We don't need to know $\mathbb{P}(\mathcal{D})$ which we generally don't know

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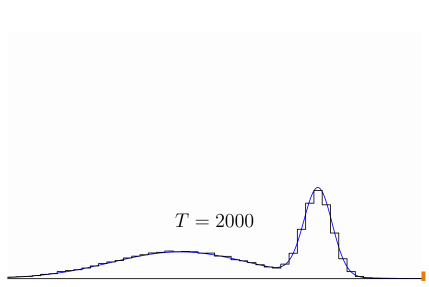
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Burn-In



• Even when we have reached this *equilibration time* each sample

 π (i.e. we have forgotten our initial state)

• We don't even know how long we have to wait!

- is correlated with the previous sample
- To get a good approximation to the posterior expectation requires running for many times the equilibration time
- Note, if we are just finding sample averages then we can use all samples after equilibrating even if they are not independent

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Proposals and Metropolis-Hastings

- We have some freedom in choosing a new proposal θ' from our current position heta—a good choice can increase the acceptance rate making the MCMC more efficient
- We define the proposal distribution $p(\theta'|\theta)$
- For the standard Metropolis algorithm to work we require $p(\boldsymbol{\theta}'|\boldsymbol{\theta}) = p(\boldsymbol{\theta}|\boldsymbol{\theta}')$
- In some cases (e.g when $\theta_i > 0$) this can be hard to achieve
- We can modify our update rule to accept a move with probability

$$\min\left(1, \frac{p(\boldsymbol{\theta}|\boldsymbol{\theta}')f(\mathcal{D}|\boldsymbol{\theta}')f(\boldsymbol{\theta}')}{p(\boldsymbol{\theta}'|\boldsymbol{\theta})f(\mathcal{D}|\boldsymbol{\theta})f(\boldsymbol{\theta})}\right) \blacksquare$$

Traffic Rate

• Consider monitoring the flow of traffic where we have data

$$\mathcal{D} = (N_1, N_2, \dots, N_n)$$

where N_i is the number of car that past on day i

- We assume $N_i \sim \text{Poi}(\mu)$ and want to infer μ
- The Poisson distribution has a beta conjugate prior
- We don't have any prior knowledge on μ so we use a non-informative prior $Gam(\mu|0,0) = 1/\mu$
- Note that we can solve this problem exactly—however, lets compare with MCMC

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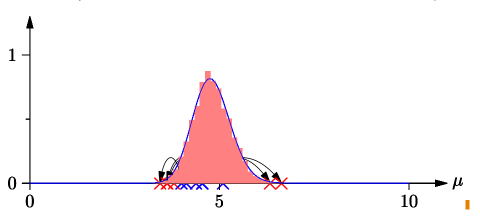
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MCMC in Practice

 $\mathcal{D} = \{4, 4, 6, 4, 2, 2, 5, 9, 5, 4, 3, 2, 5, 4, 4, 11, 6, 2, 3, 11\}$



Proposal Distribution

- If we can choose our proposal distribution $p(\mu'|\mu)$ to be close to the posterior distribution then our acceptance rate would be close to 1
- We choose $p(\mu'|\mu) = \operatorname{Gam}(\mu'|\mu,\mu^2)$ which has $\mathbb{E}[\mu'] = \mu$ and variance 1
- We update with probability min(1,r) where

$$r = \frac{\operatorname{Gam}(\mu|\mu'^{2}, \mu') \frac{1}{\mu'} \prod_{i=1}^{n} \operatorname{Poi}(N_{i}|\mu')}{\operatorname{Gam}(\mu'|\mu^{2}, \mu) \frac{1}{\mu} \prod_{i=1}^{n} \operatorname{Poi}(N_{i}|\mu)}$$

$$= \frac{\mu \operatorname{Gam}(\mu|\mu'^{2}, \mu')}{\mu' \operatorname{Gam}(\mu'|\mu^{2}, \mu)} e^{-n(\mu' - \mu) + \sum_{i=1}^{n} N_{i} \log\left(\frac{\mu'}{\mu}\right)}$$

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MCMC Details

- To compute correct histograms you need to count samples where no move is made multiple times
- On modern computers its quite quick to compute millions of samples
- The code is not very difficult to write (although care is need to get everything correct)
- This can be used on complicated problems such as topic models (LDA) with thousands of parameters
- The accuracy of MCMC is slow if it takes a long time to sample the posterior distribution

The MCMC Industry

- MCMC provides a means to accurately sample from very complex models
- There have been many advanced techniques developed to improve MCMC performance
- E.g. hybrid MCMC simulates a dynamics to find good proposals with similar probability far from the starting point
- Often it seems that MCMC is complicated because there are so many optimisations, but often simple implementations are sufficient

Conclusions

- As soon as we use complex models we are no longer able to compute the posterior in closed form
- Monte Carlo techniques and particularly MCMC are a very general method for computing samples from the posterior
- These techniques have been highly developed, but very frequently even simple implementations are sufficient to do good inference.
- Variational methods provide an approximate closed form solution to problems with complex likelihoods
- Variational methods are mathematically challenging, but are potentially far faster to compute than MCMCI

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