# Lecture 21: Hamiltonian Monte Carlo

STA702

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#### 2

# Gibbs sampling

Consider model

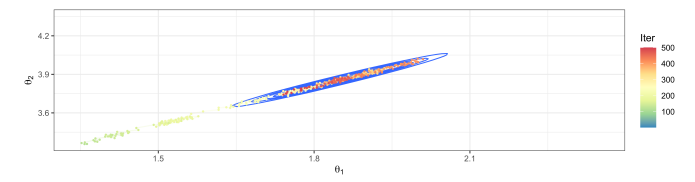
$$egin{aligned} oldsymbol{Y}_1,\ldots,oldsymbol{Y}_n &\sim \mathsf{N}_2\left(oldsymbol{ heta},oldsymbol{\Sigma}
ight);\ heta_j &\sim \mathsf{N}(0,1) \qquad j=1,2. \end{aligned}$$

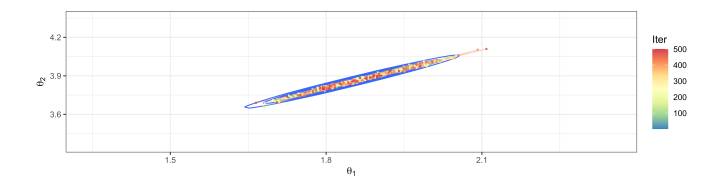
• Suppose that the covariance matrix  $\Sigma$  is known and has the form

$$oldsymbol{\Sigma} = egin{bmatrix} 1 & 
ho \ 
ho & 1 \end{bmatrix}$$

ullet What happens when ho=0.995 for sampling from full conditionsals for  $heta_1$  and  $heta_2$ ?

# Gibbs vs Stan samples

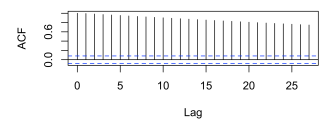




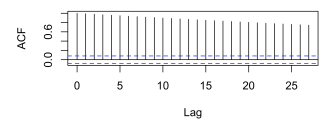
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#### **ACF**

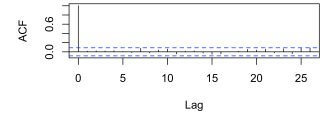
Series norm\_gibbs\_samps[, 1]



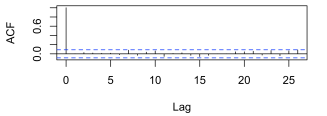
Series norm\_gibbs\_samps[, 2]



Series stan\_res\$theta[, 1]



Series stan\_res\$theta[, 2]



#### Hamiltonian Monte Carlo (HMC)

- HMC creates transitions that *efficiently explore the parameter space* by using concepts from Hamiltonian mechanics.
- In Hamiltonian mechanics, a physical system is specified by positions  ${\bf q}$  and momenta  ${\bf p}$ .
- A space defined by these coordinates is called a phase space
- If the parameters of interest in a typical MCMC method are denoted as  $q_1, \ldots, q_K$ , then HMC introduces auxiliary **momentum** parameters  $p_1, \ldots, p_K$  such that the algorithm produces draws from the joint density:

$$\pi(\mathbf{q}, \mathbf{p}) = \pi(\mathbf{p}|\mathbf{q})\pi(\mathbf{q})$$

• marginalizing over the  $p_k$ 's, we recover the marginal distribution of the  $q_k$ 's Therefore, if we create a Markov Chain that converges to  $\pi(\mathbf{q}, \mathbf{p})$ , we have immediate access to samples from  $\pi(\mathbf{q})$ , which is our target distribution.

#### Hamiltonian

• Hamilton's equations describe the time evolution of the system in terms of the **Hamiltonian**,  $\mathcal{H}$ , which corresponds to the total energy of the system:

$$\mathcal{H}(\mathbf{p}, \mathbf{q}) = K(\mathbf{q}, \mathbf{p}) + U(\mathbf{q})$$

•  $K(\mathbf{q}, \mathbf{p})$  represents the **kinetic energy** of the system and is equal to the negative logarithm of the momentum distribution, e.g.

$$K(\mathbf{p}) = rac{\mathbf{p}^T \mathbf{M}^{-1} \mathbf{p}}{2} = \sum_i rac{p_i^2}{2m_i}$$

- M is the Mass matrix
- $U(\mathbf{q})$  the **potential energy** of the system; equal to the negative logarithm of the distribution of  $\mathbf{q}$ .
- Joint  $\pi(\mathbf{q},\mathbf{p}) \propto e^{-\mathcal{H}(\mathbf{q},\mathbf{p})} = e^{-K(\mathbf{p})}e^{-U(\mathbf{q})}$

#### **Evolution**

- At each iteration of the sampling algorithm, HMC implementations make draws from some distribution  $\pi(\mathbf{p}|\mathbf{q})$  and then evolves the system  $(\mathbf{q},\mathbf{p})$  to obtain the next sample of  $\mathbf{q}$ .
- To "evolve the system" is to move  $(\mathbf{q}, \mathbf{p})$  forward in "time," i.e. to change the values of  $(\mathbf{q}, \mathbf{p})$  according to Hamilton's differential equations:

$$\frac{d\mathbf{p}}{dt} = -\frac{\partial \mathcal{H}}{\partial \mathbf{q}} = -\frac{\partial K}{\partial \mathbf{q}} - \frac{\partial U}{\partial \mathbf{q}}$$
$$\frac{d\mathbf{q}}{dt} = +\frac{\partial \mathcal{H}}{\partial \mathbf{p}} = +\frac{\partial K}{\partial \mathbf{p}}$$

ullet Defines a mapping  $T_s$  from the state at any time t to the state at t+s

"The differential change in momentum parameters  ${f p}$  over time is governed in part by the differential information of the density over the target parameters."

# **Key Properties**

- 1. Reversibility The mapping of the state at time t ( $\mathbf{p}(t)$ ,  $\mathbf{q}(t)$ ) to the state at t+s ( $\mathbf{p}(t+s)$ ,  $\mathbf{q}(t+s)$ ) is one-to-one and we have an inverse  $T_{-s}$  obtained by negating the derivatives;  $K(\mathbf{p}) = K(-\mathbf{p})$  MCMC updates using the dymamics don't modify invariant distribution!
- 2. **Invariance/Conservation** the dymamics keep the Hamiltonian invariant if we use the dynamics to generate proposals, the acceptance probability of MH is equal to one if  $\mathcal{H}$  is kept invariant!
- 3. Volume Preservation/Symplectiness the mapping  $T_s$  of a region R to  $T_s(R)$  preserves volume means that we do not need to compute Jacobians

in practice we need to use approximations to solve the PDE's so won't have exact invariance etc so acceptance probability is not 1!

# **Approximate Solutions to Differential Eqs**

- Discretize time into steps  $\epsilon$
- Euler's Method for ith coordinate

$$egin{aligned} p_i(t+\epsilon) &= p_i(t) + \epsilon rac{dp_i}{t}(t) = p_i(t) - \epsilon rac{\partial U(q_i(t))}{\partial q_i} \ q_i(t+\epsilon) &= q_i(t) + \epsilon rac{dq_i}{t}(t) = q_i(t) + \epsilon rac{\partial K(p_i(t))}{\partial p_i} = q_i(t) + \epsilon rac{p_i(t)}{m_i} \end{aligned}$$

Modified Euler method

$$p_i(t+\epsilon) = p_i(t) - \epsilon rac{\partial U(q_i(t))}{\partial q_i} \ q_i(t+\epsilon) = q_i(t) + \epsilon rac{p_i(t+\epsilon)}{m_i}$$

# Leapfrog

- Divide into half steps
- apply Modified Euler

$$egin{aligned} p_i(t+\epsilon/2) &= p_i(t) - rac{\epsilon}{2} rac{\partial U(q_i(t))}{\partial q_i} \ q_i(t+\epsilon) &= q_i(t) + \epsilon rac{p_i(t+\epsilon/2)}{m_i} \ p_i(t+\epsilon) &= p_i(t) - rac{\epsilon}{2} rac{\partial U(q_i(t+\epsilon))}{\partial q_i} \end{aligned}$$

- Preserves volume exactly
- Reversible
- We don't get exact invariance (so probability of acceptance is not 1)
- Step size and number of steps is still important!

#### MCMC with HMC

Steps: replace  ${f q}$  with  ${m heta}$ 

- 1. sample a new value for the momentum  $\mathbf{p}^{(t)} \sim \mathsf{N}(\mathbf{0}_K, \mathbf{M})$
- 2. Metropolis: from current state  $(\mathbf{q}^{(t-1)}, \mathbf{p}^{(t)})$  simulate proposal  $(\mathbf{q}^*, \mathbf{p}^*)$  using Hamiltonian dynamics by applying Leapfrog with step size  $\epsilon$  for L steps (tuning parameters)
- 3. Accept or reject acceptance probability is

$$\min\{1, \exp(-\mathcal{H}(\mathbf{q}^*, \mathbf{p}^*) + \mathcal{H}(\mathbf{q}^{(t-1)}, \mathbf{p}^{(t)})\}$$

#### **Tuning**

- in addition to tuning  $\epsilon$  and L, we can tune  ${f M}$
- ullet  $\operatorname{Cov}(\mathbf{q}) = \mathbf{V}$  can be highly variable
- Consider reparameterization  ${f A}{f q}={f q}'$  so that  ${
  m Cov}({f A}{f q})={f A}{f V}{f A}^T={f I}_d;$   ${f A}={f V}^{-1/2}$
- eliminates posterior correlation!
- general trick of reparameterizing to reduce posterior correlation is often called **pre-conditioning** improves efficiency!
- use  $\mathbf{M} = \mathbf{I}_d$
- Automatic tuning is achieved by the No-U-Turn-Sampler (NUTS)
- bit complicated, but used by STAN
- other variations Metropolis-Adjusted Langevin Algorithm (MALA)

# **Hybrid Approaches**

Recall mixed effects model

$$Y_{ij} = \mathbf{x}_{ij}^T oldsymbol{eta} + \mathbf{z}_{ij}^T oldsymbol{\gamma}_j + \epsilon_{ij} \qquad \epsilon_{ij} \sim \mathsf{N}(0, \sigma^2)$$

- random effects  $oldsymbol{\gamma}_j \sim \mathsf{N}_d(oldsymbol{0}_d, oldsymbol{\Psi})$  (diagonal  $oldsymbol{\Psi}$ )
- marginalize over the random effects

$$\mathbf{Y}_j = \mathsf{N}(\mathbf{X}_j oldsymbol{eta}, \mathbf{Z}_j oldsymbol{\Psi} \mathbf{Z}_j^T + \sigma^2 \mathbf{I}_{n_j})$$

- we could use Gibbs on the conditional model, but we may get slow mixing (i.e. due to updating variance components)
- run HMC within Gibbs to update the variance components  $\Psi$  and  $\sigma^2$  using the marginal model given  $\pmb{\beta}$
- HMC in its basic form doesn't like constraints so reparameterize to use log transformations

#### **Advantages & Disadvantages**

- HMC can produce samples with low correlation and high acceptance ratio!
- driven by step size (larger time steps mean values are farther away but may lead to lower acceptance- error is  $O(\epsilon^2)$  for the leapfrog method)
- number of steps (more steps reduces correlation; to avoid U turns stan uses NUTS)
- most implementations limited to continuous variables (need gradients of log densities)
- need to calculate gradients (automatic differentiation methods)
- can mix Gibbs (for discrete) and HMC (for continuous)
- Nishimura et al (2020 Biometrika) for HMC with discrete targets
- rates of convergence and other theory