project_report

June 2, 2023

1 Monte Carlo Methods Project

1.1 Getting started

To run the code, first install Julia:

1. Visit https://julialang.org/downloads/ and select the installer for your system (I used 1.8.3, but probably 1.9.0 will work just as well). If you're on Linux, run:

```
wget https://julialang-s3.julialang.org/bin/linux/x64/1.8/julia-1.8.3-linux-x86_64.tar.gz tar zxvf julia-1.8.3-linux-x86_64.tar.gz
```

2. On Linux, that's all you have to do. You can choose to make the julia executable more accessible by, e.g., opening ~/.bashrc and adding the line:

```
export PATH="$PATH:/path/to/<Julia directory>/bin"
```

3. Make sure to start a new terminal or run source ~/.bashrc afterwards.

Consequently, you want to install the relevant packages:

1. Move one level above the current directory (the current directory is the one that contains this file) and activate Julia:

```
cd ..
julia
```

2. Inside Julia, change to the Pkg mode by typing]. In there, you want to activate the project toml file and instantiate the project. This will install all necessary packages and precompile them.

```
]
activate mcm_project
instantiate
```

1.2 Running the code

If you have Visual Studio Code installed, you could open the project.ipynb file, select Julia as your notebook kernel, and then hopefully be able to run everything cell by cell. Slightly simpler would be to run the more barebones project.jl, which contains some additional code to create an images directory and save the generated images as png files. To use the last option, run:

```
cd mcm_project
julia --project=. project.jl
```

1.3 Dependencies

```
[]: using LinearAlgebra using Distributions using Plots
```

General note: It was not quite clear to me whether the starting values should be returned by the sampling algorithms, or discarded. Either way, I don't think it matters much, but I decided to be prudent and discard the starting values.

1.4 Exercise (a)

```
[]: function quadraticpotential_grad(q)
         q
     end
     function leapfrog(q0, p0, tau, 1, mass, potential_grad::Function)
         # Determine dimensionality
         d = length(q0)
         # Also accomodate initial values
         qs = [Vector{Float64}(undef, d) for _ = 1:(l+1)]
         ps = [Vector{Float64}(undef, d) for _ = 1:(1+1)]
         qs[1] = q0
         ps[1] = p0
         for i = 2:(1+1)
             p_half = ps[i - 1] .- tau / 2 .* potential_grad(qs[i - 1])
             qs[i] = qs[i - 1] .+ tau * inv(mass) * p_half
             ps[i] = p_half .- tau / 2 .* potential_grad(qs[i])
         end
         # Discard initial value
         (qs[2:(1+1)], ps[2:(1+1)])
     end
```

leapfrog (generic function with 1 method)

1.5 Exercise (b)

```
[]: q0 = [0.]
  p0 = [1.]
  tau = 1.2
  1 = 20
  mass = [1.;;]

(ps1, qs1) = leapfrog(q0, p0, tau, l, mass, quadraticpotential_grad)
```

```
tau = 0.3
1 = 80

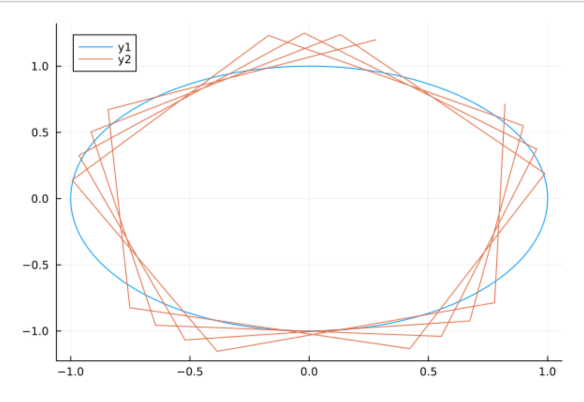
(ps2, qs2) = leapfrog(q0, p0, tau, 1, mass, quadraticpotential_grad);
```

```
[]: qs1_flat = [v[1] for v in qs1]
ps1_flat = [v[1] for v in ps1]

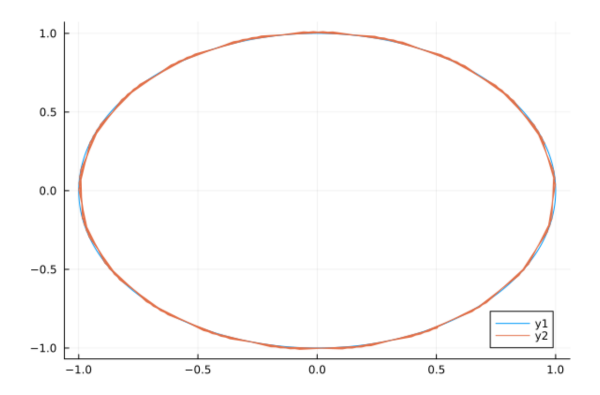
qs2_flat = [v[1] for v in qs2]
ps2_flat = [v[1] for v in ps2];
```

```
[]: t = collect(0:0.01:2)
   q_exact = -cos.(t .+ /2)
   p_exact = sin.(t .+ /2)

plot(q_exact, p_exact)
   plot!(qs1_flat, ps1_flat)
```



```
[]: plot(q_exact, p_exact)
plot!(qs2_flat, ps2_flat)
```



1.6 Exercise (c)

Substituting Equation (4) into (7), and then using Equation (8), we get:

$$\begin{split} \pi(\mathbf{q}, \mathbf{p}) &\propto \exp\left(-H(\mathbf{q}, \mathbf{p})\right) \\ &\propto e^{-U(\mathbf{q}) - K(\mathbf{p})} \\ &\propto e^{--ln(\pi(\mathbf{q}))} e^{-\frac{1}{2}\mathbf{p}^T M^{-1}\mathbf{p}} \\ &\propto \pi(\mathbf{q}) e^{-\frac{1}{2}\mathbf{p}^T M^{-1}\mathbf{p}}. \end{split}$$

If we fix \mathbf{p} , we can treat it as a constant:

$$\pi(\mathbf{q} \mid \mathbf{p}) \propto \pi(\mathbf{q}) e^{-\frac{1}{2}\mathbf{p}^T M^{-1}\mathbf{p}} \\ \propto \pi(\mathbf{q}).$$

This shows that sampling from $\pi(\mathbf{q}, \mathbf{p})$ is equivalent to sampling from $\pi(\mathbf{q})$ if \mathbf{p} is fixed.

1.7 Exercise (d)

Fixing \mathbf{q} to specify a pdf for \mathbf{p} yields:

$$\begin{split} \pi(\mathbf{p} \mid \mathbf{q}) &\propto \pi(\mathbf{q}) e^{-\frac{1}{2}\mathbf{p}^T M^{-1}\mathbf{p}} \\ &\propto e^{-\frac{1}{2}\mathbf{p}^T M^{-1}\mathbf{p}} \\ &\propto (2\pi)^{-d/2} \det\left(M\right)^{-1/2} e^{-\frac{1}{2}\mathbf{p}^T M^{-1}\mathbf{p}}. \end{split}$$

In other words, $\pi(\mathbf{p} \mid \mathbf{q})$ is equal to a 0-mean multivariate Gaussian pdf up to a normalising constant. M can be interpreted as the covariance matrix of that Gaussian.

1.8 Exercise (e)

 $-H(\mathbf{q}^*, \mathbf{p}^*) + H(\mathbf{q}, \mathbf{p})$ can be interpreted as $\frac{\Delta H}{\Delta t}$. We have seen above that with an exact trajectory, the Hamiltonian is conserved: $\frac{dH}{dt} = 0$. Under this condition, the acceptance probability becomes:

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

= $\min \{1, \exp(0)\}$
= 1.

This means that using an approach that approximates the trajectory well would lead to acceptance probabilities close to 1.

1.9 Exercise (f)

```
[]: function quadraticpotential(q)
         0.5 * q' * q
     end
     function hamiltonian(q, p, mass, potential::Function)
         potential(q) + 0.5 * p' * inv(mass) * p
     end
     function hmc(n, q0, tau_dist, 1, potential::Function, potential_grad::Function)
         # Determine dimensionality
         d = length(q0)
         # Also accomodate initial values
         qs = [Vector{Float64}(undef, d) for _ = 1:(n+1)]
         ps = [Vector{Float64}(undef, d) for _ = 1:(n+1)]
         # Julia can infer the size of I from the mathematical context
         mass = I
         norm = MvNormal(zeros(Float64, d), mass)
         qs[1] = q0
```

```
for t = 2:(n+1)
        ps[t - 1] = vec(rand(norm, 1))
        tau = rand(tau_dist, 1)[1]
        # Note that qs star has a different dimensionality (1) than qs (n)
        (qs_star, ps_star) = leapfrog(qs[t - 1], ps[t - 1], tau, 1, mass, 
 →potential_grad)
        q_star = qs_star[1]
        p_star = ps_star[1]
        h_prev = hamiltonian(qs[t - 1], ps[t - 1], mass, potential)
        h_star = hamiltonian(q_star, p_star, mass, potential)
        alpha = min(1, exp(h_prev - h_star))
        if rand() < alpha</pre>
            qs[t] = q_star
            ps[t] = p_star
        else
            qs[t] = qs[t - 1]
            ps[t] = ps[t - 1]
        end
    end
    (qs[2:(n+1)], ps[2:(n+1)])
end
```

hmc (generic function with 1 method)

Testing with the quadratic potential and its gradient. Since we are not applying a -log to to the potential, we are essentially sampling from:

$$\pi(\mathbf{q} \mid \mathbf{p}) \propto e^{-U(\mathbf{q}) - K(\mathbf{p})}$$

$$\propto e^{-U(\mathbf{q})}$$

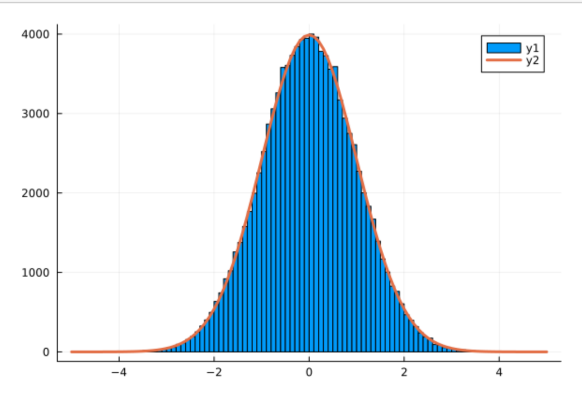
$$\propto e^{-\frac{1}{2}\mathbf{q}^T\mathbf{q}},$$

which is proportional to a multivariate normal pdf with mean 0 and covariance matrix I. In other words, we should get a normal distribution if we set d = 1, which the histogram below shows to be the case.

```
[]: tau_dist = Uniform(0.5, 1.5)
samples, _ = hmc(100000, q0, tau_dist, l, quadraticpotential,
quadraticpotential_grad);
```

```
[]: samples_flat = [s[1] for s in samples]
histogram(samples_flat)
```

```
x = -5:0.1:5
plot!(x, pdf.(Normal(0, 1), x) .* 1e4, linewidth = 3)
```



1.10 Exercise (g)

```
function rw_mh(n, start, prop_dist, target::Function)
    d = length(start)

samples = [Vector{Float64}(undef, d) for _ = 1:(n+1)]
    samples[1] = start

for i = 2:(n+1)
    sample_prop = vec(rand(prop_dist, 1)) + samples[i - 1]

dens_prev = target(samples[i - 1])
    dens_prop = target(sample_prop)
    alpha = min(1, dens_prop / dens_prev)

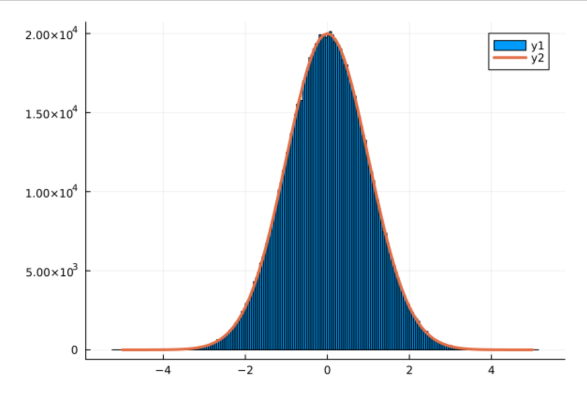
if rand() < alpha
    samples[i] = sample_prop
    else
    samples[i] = samples[i - 1]
    end</pre>
```

```
end
samples[2:(n+1)]
end
```

rw_mh (generic function with 1 method)

```
[]: target(x) = exp(-quadraticpotential(x))
samples = rw_mh(1000000, [0.], MvNormal(zeros(1), I), target)
histogram([s[1] for s in samples])

x = -5:0.1:5
plot!(x, target.(x) .* 2e4, linewidth = 3)
```



1.11 Exercise (h)

```
[]: function rand_sigma_mh(n, start, sigma_dist, target::Function)
    mh_samples = [Vector{Float64}(undef, d) for _ = 1:(n+1)]
    mh_samples[1] = start

mu = zeros(size(start))

for i = 2:(n+1)
```

```
sigma = rand(sigma_dist, 1)[1]
prop_dist = MvNormal(mu, sigma * I)

mh_samples[i] = rw_mh(1, mh_samples[i - 1], prop_dist, target)[1]
end

mh_samples[2:(n+1)]
end
```

rand_sigma_mh (generic function with 1 method)

To be able to use Hamiltonian Monte Carlo, we first need to find the gradient $\nabla_{\mathbf{q}}U$. Since

$$\nabla_{\mathbf{q}} U = \begin{pmatrix} \frac{\partial U}{\partial q_1} \\ \vdots \\ \frac{\partial U}{\partial q_d} \end{pmatrix},$$

we need to find $\frac{\partial U}{\partial q_i}$, where $i=1,\ldots,d$. Because we want to sample from a multivariate normal, we get the -log of its pdf:

$$\begin{split} \frac{\partial U}{\partial q_i} &= \frac{\partial}{\partial q_i} \left[-\log \left((2\pi)^{-\frac{d}{2}} \det(\Sigma)^{-\frac{1}{2}} e^{-\frac{1}{2} \mathbf{q}^T \Sigma^{-1} \mathbf{q}} \right) \right] \\ &= \frac{\partial}{\partial q_i} \left[-\left(\log \left((2\pi)^{-\frac{d}{2}} \det(\Sigma)^{-\frac{1}{2}} \right) + \log \left(e^{-\frac{1}{2} \mathbf{q}^T \Sigma^{-1} \mathbf{q}} \right) \right) \right] \\ &= \frac{\partial}{\partial q_i} \left[\frac{1}{2} \mathbf{q}^T \Sigma^{-1} \mathbf{q} \right]. \end{split}$$

 Σ is diagonal, so its inverse can be found by taking the reciprocals of its diagonal values:

$$\begin{split} \frac{\partial U}{\partial q_i} &= \frac{\partial}{\partial q_i} \left[\frac{1}{2} \mathbf{q}^T \begin{pmatrix} \frac{1}{\sigma_1} & \dots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \dots & \frac{1}{\sigma_d} \end{pmatrix} \mathbf{q} \right] \\ &= \frac{\partial}{\partial q_i} \left[\frac{1}{2} \mathbf{q}^T \begin{pmatrix} \frac{1}{\sigma_1} q_1 \\ \vdots \\ \frac{1}{\sigma_d} q_d \end{pmatrix} \right] \\ &= \frac{\partial}{\partial q_i} \left[\frac{1}{2} \left(\frac{1}{\sigma_1} q_1^2 + \dots + \frac{1}{\sigma_d} q_d^2 \right) \right] \\ &= \frac{1}{\sigma_i} q_i. \end{split}$$

```
[]: sigmas = collect(0.01:0.01:1)

Σ = Diagonal(sigmas)

d = size(Σ, 1)

= zeros(d)
```

```
target(x) = pdf(MvNormal(, Σ), x)

n = 1000
l = 150
tau_dist = Uniform(0.0104, 0.0156)
potential(x) = -log(target(x))
potential_grad(x) = @. (1 / sigmas) * x
samples_hmc, _ = hmc(n, , tau_dist, l, potential, potential_grad)

n = 150000
sigma_dist = Uniform(0.0176, 0.0264)
samples_mh = rand_sigma_mh(n, , sigma_dist, target)
samples_mh_thinned = samples_mh[1:150:length(samples_mh)];
```

Plotting the HMC and thinned MH samples (first on top of each other, then separately) shows that the MH chain moves/mixes much slower than the HMC one. In other words, there seems to be more autocorrelation in the MH samples.

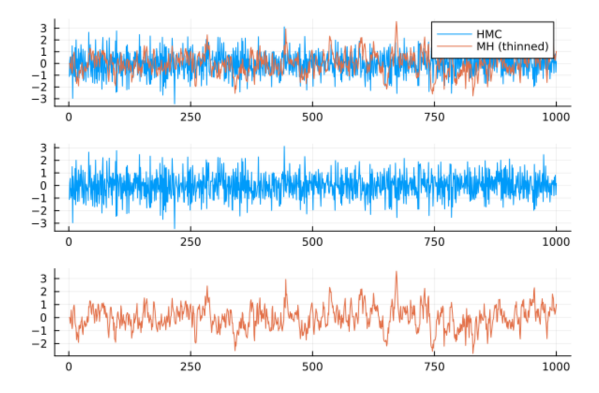
```
[]: last_comp_hmc = [v[d] for v in samples_hmc]
last_comp_mh_th = [v[d] for v in samples_mh_thinned]

plot(last_comp_hmc, labels = "HMC")
p1 = plot!(last_comp_mh_th, labels = "MH (thinned)")

p2 = plot(last_comp_hmc, legend = false)

p3 = plot(last_comp_mh_th, color = 2, legend = false)

plot(p1, p2, p3, layout = (3, 1))
```



Plotting the sample variation over the component standard deviations, we see a fairly constant error for the HMC sampler (except for a little dip at the very end), while for the MH sampler, the error seems to increase with the standard deviation.

```
[]: # Turns the vector of vectors inside-out (components outside, samples inside)
    comps_hmc = [[v[i] for v in samples_hmc] for i = 1:d]
    means_hmc = mean.(comps_hmc)
    plot(sigmas, means_hmc, labels = "HMC")

comps_mh = [[v[i] for v in samples_mh] for i = 1:d]
    means_mh = mean.(comps_mh)
    p1 = plot!(sigmas, means_mh, labels = "MH (thinned)")

p2 = plot(sigmas, means_hmc, legend = false)

p3 = plot(sigmas, means_mh, color = 2, legend = false)

plot(p1, p2, p3, layout = (3, 1))
```

