MTH3045: Statistical Computing

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Challenges I

 Go to Challenges I of the week 6 lecture 1 challenges at https://byoungman.github.io/MTH3045/challenges

Sherman-Morrison formula / Woodbury matrix identity

- In general, unless we actually need the inverse of a matrix (such as for standard errors of regression coefficients in a linear model), we should solve systems of linear equations
- Sometimes, though, we do need or might just have an inverse, and want to calculate something related to it
- The following are a set of formulae, which go by various names, that can be useful in this situation

Woodbury's formula

- Consider an $m \times m$ matrix **A**, with m large, and for which we have the inverse, \mathbf{A}^{-1}
- Suppose **A** receives a small update of the form $\mathbf{A} + \mathbf{U}\mathbf{V}^\mathsf{T}$, for $m \times n$ matrices \mathbf{U} and \mathbf{V} where $n \ll m$
- Then, by Woodbury's formula,

$$(A + UV^{T})^{-1} = A^{-1} - A^{-1}U(I_{n} + V^{T}A^{-1}U)^{-1}V^{T}A^{-1}$$

- Remark: What's important to note here is that we're looking to calculate an $m \times m$ inverse with m large, and so in general this will be an $O(m^3)$ calculation based on the LHS
- However, in the above, the RHS only requires that we to invert an $n \times n$ matrix, at cost $O(n^3)$, which is much less that of the LHS, if we already have \mathbf{A}^{-1} .

Sherman-Morrison-Woodbury formula

• Woodbury's formula above generalises to the so-called **Sherman-Morrison-Woodbury formula** by introducing the $n \times n$ matrix **C**, so that

$$(\mathbf{A} + \mathbf{U}\mathbf{C}\mathbf{V}^\mathsf{T})^{-1} = \mathbf{A}^{-1} - \mathbf{A}^{-1}\mathbf{U}(\mathbf{C}^{-1} + \mathbf{V}^\mathsf{T}\mathbf{A}^{-1}\mathbf{U})^{-1}\mathbf{V}^\mathsf{T}\mathbf{A}^{-1}$$

Challenges II

 Go to Challenges II of the week 6 lecture 1 challenges at https://byoungman.github.io/MTH3045/challenges

Sherman-Morrison formula

 The Sherman-Morrison formula is the special case of Woodbury's formula (and hence the Woodbury-Sherman-Morrison formula) in which the update to A can be considered in terms of m-vectors u and v, so that

$$(\mathbf{A} + \mathbf{u}\mathbf{v}^\mathsf{T})^{-1} = \mathbf{A}^{-1} - \frac{\mathbf{A}^{-1}\mathbf{u}\mathbf{v}^\mathsf{T}\mathbf{A}^{-1}}{1 + \mathbf{v}^\mathsf{T}\mathbf{A}^{-1}\mathbf{u}}$$

 Remark: The Sherman-Morrison formula is particularly useful because it requires no matrix inversion

Example: Bayesian linear regression I

Recall from MTH2006 the normal linear model where

$$Y_i \sim N(\mathbf{x}_i^\mathsf{T} \boldsymbol{\beta}, \sigma^2)$$

with independent errors $\varepsilon_i = Y_i - \mathbf{x}_i^\mathsf{T} \boldsymbol{\beta} \mathbf{s}$, for $i = 1, \dots, n$, where $\mathbf{x}_i^\mathsf{T} = (1, x_{i1}, \dots, x_{ip})$ is the *i*th row of design matrix \mathbf{X} and where $\boldsymbol{\beta} = (\beta_0, \beta_1, \dots, \beta_p)^\mathsf{T}$

Hence

$$\mathbf{Y} \mid \mathbf{X}\boldsymbol{\beta} \sim MVN_n(\mathbf{X}\boldsymbol{\beta}, \sigma^2\mathbf{I}_n)$$

- In Bayesian linear regression, the elements of β and σ^2 are not fixed, unknown parameters: they are random variables, and we must declare a priori our beliefs about their distribution
- The conjugate prior is that

$$eta \sim extit{MVN}_{p+1}(oldsymbol{\mu}_{oldsymbol{eta}}, oldsymbol{\Sigma}_{oldsymbol{eta}}^{-1})$$

• Integrating out eta gives

$$\mathbf{Y} \sim MVN_n(\mathbf{X}\boldsymbol{\mu}_{oldsymbol{eta}}, \sigma^2\mathbf{I}_n + \mathbf{X}\mathbf{\Sigma}_{oldsymbol{eta}}^{-1}\mathbf{X}^{\mathsf{T}})$$

Example: Bayesian linear regression II

- Now suppose that we want to evaluate the marginal likelihood for an observation, y, say. Recall the MVN pdf
- The Mahalanobis distance then involves the term

$$(\mathbf{y} - \mathbf{X}\boldsymbol{\mu}_{\boldsymbol{\beta}})^{\mathsf{T}}(\sigma^{2}\mathbf{I}_{n} + \mathbf{X}\boldsymbol{\Sigma}_{\boldsymbol{\beta}}^{-1}\mathbf{X}^{\mathsf{T}})^{-1}(\mathbf{y} - \mathbf{X}\boldsymbol{\mu}_{\boldsymbol{\beta}})$$

- The covariance of the marginal distribution, $\sigma^2 \mathbf{I}_n + \mathbf{X} \mathbf{\Sigma}_{\beta}^{-1} \mathbf{X}^{\mathsf{T}}$, is typically dense, expensive to form, and leads to expensive solutions to systems of linear equations
- Its inverse, however, can be computed through the Sherman-Morrison-Woodbury formula with $\mathbf{A}^{-1}=\sigma^{-2}\mathbf{I}_n$, $\mathbf{U}=\mathbf{V}=\mathbf{X}$, and $\mathbf{C}=\mathbf{\Sigma}_{\boldsymbol{\beta}}^{-1}$

Bibliographic notes

- For more details on matrix decompositions, consider Wood (2015, Appendix B) for a concise overview
- For fuller details consider Monahan (2011, chaps. 3, 4 and 6) or Press et al. (2007, chap. 2 and 11)

Chapter 4: Numerical Calculus

Numerical Calculus: Motivation

In statistics, we often rely on the Normal distribution with pdf

$$\phi(x; \mu, \sigma^2) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{(x-\mu)^2}{2\sigma^2}\right]$$

and cdf

$$\Phi(x;\mu,\sigma^2) = \int_{-\infty}^{x} \phi(u;\mu,\sigma^2) du.$$

- Unfortunately no closed form exists for $\Phi(x; \mu, \sigma^2)$
- However, if x, μ and σ are stored in R as x, mu and sigma, we can still evaluate $\Phi(x; \mu, \sigma^2)$ with pnorm(x, mu, sigma)
- This is one example of a frequently-occurring situation in which we somehow want to evaluate an intractable integral
- This raises the question: are there generic methods that let us evaluate intractable integrals?
- The answer is often numerical integration

Numerical Calculus: Motivation

- Remark: In this chapter, we'll consider generic methods for integration, i.e. that work in many scenarios
- Sometimes, such as evaluating $\Phi(x; \mu, \sigma^2)$, specific algorithms will give more accurate results
- This is what R does for pnorm()

Numerical Differentiation

- In Chapter 5 we will cover optimisation of functions, such as numerically finding maximum likelihood estimates when analytical solutions aren't available
- We'll see that supplying derivatives can considerably improve estimation, typically in terms of reducing computation time
- Here we'll cover some useful results in terms of differentiation of matrices, which will later prove useful
- No knowledge of analytical matrix calculus beyond these results will be needed for MTH3045
- The matrix cookbook (Petersen and Pedersen 2012), however, can provide you with a much more thorough set of differentiation rules, should you ever need them

Differentiation definitions I

- **Definition**: Consider $f : \mathbb{R}^n \to \mathbb{R}$, which we'll consider a function of vector $\mathbf{x} = (x_1, \dots, x_n)^\mathsf{T}$
- The gradient operator, ∇ , is defined as

$$\nabla f(\mathbf{x}) = \begin{pmatrix} \frac{\partial f(\mathbf{x})}{\partial x_1} \\ \frac{\partial f(\mathbf{x})}{\partial x_2} \\ \vdots \\ \frac{\partial f(\mathbf{x})}{\partial x_n} \end{pmatrix}$$

• Note that in MTH3045 we will have no cause to consider multivariate functions, i.e. to consider $\mathbf{f}: \mathbb{R}^n \to \mathbb{R}^m$, for m > 1

Differentiation definitions II

- **Definition**: Consider again $f: \mathbb{R}^n \to \mathbb{R}$
- The Hessian matrix is the matrix of second derivatives of f, whereas the gradient operator considered first derivatives, and is given by

$$\nabla^2 f(\mathbf{x}) = \begin{pmatrix} \frac{\partial^2 f}{\partial x_1^2} & \frac{\partial^2 f}{\partial x_1 \partial x_2} & \cdots & \frac{\partial^2 f}{\partial x_1 \partial x_n} \\ \frac{\partial^2 f}{\partial x_2 \partial x_1} & \frac{\partial^2 f}{\partial x_2^2} & \cdots & \frac{\partial^2 f}{\partial x_2 \partial x_n} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{\partial^2 f}{\partial x_n \partial x_1} & \frac{\partial^2 f}{\partial x_n \partial x_2} & \cdots & \frac{\partial^2 f}{\partial x_n^2} \end{pmatrix}$$

- The Hessian matrix plays an important role in statistics, in particular for estimating parameter uncertainty via the Fisher information, which is covered in MTH3028
- In the next chapter, we'll also see that it's important for optimisation

Differentiation rules I

- Now consider $g: \mathbb{R}^n \to \mathbb{R}$ a function of **x** that, for fixed **A**, takes the quadratic form $g(\mathbf{x}) = \mathbf{x}^\mathsf{T} \mathbf{A} \mathbf{x}$ for $n \times n$ matrix **A** and n-vector **x**
- Then

$$\nabla g(\mathbf{x}) = (\mathbf{A} + \mathbf{A}^\mathsf{T})\mathbf{x}$$
 and $\nabla^2 g(\mathbf{x}) = \mathbf{A} + \mathbf{A}^\mathsf{T}$

• Note that in the case of symmetric \mathbf{A} , $\nabla g(\mathbf{x}) = 2\mathbf{A}\mathbf{x}$ and $\nabla^2 g(\mathbf{x}) = 2\mathbf{A}$

Differentiation rules II

- Next consider $h: \mathbb{R}^n \to \mathbb{R}$ a function of n-vector \mathbf{x} and p-vector \mathbf{y} that, for fixed $n \times n$ matrix \mathbf{A} and $n \times p$ matrix \mathbf{B} , takes the quadratic form $h(\mathbf{x}, \mathbf{y}) = (\mathbf{x} + \mathbf{B}\mathbf{y})^\mathsf{T} \mathbf{A} (\mathbf{x} + \mathbf{B}\mathbf{y})$
- Then

$$\frac{\partial \textit{h}(\textbf{x},\textbf{y})}{\partial \textbf{x}} = (\textbf{A} + \textbf{A}^{\mathsf{T}})(\textbf{x} + \textbf{B}\textbf{y}) \quad \text{and} \quad \frac{\partial \textit{h}(\textbf{x},\textbf{y})}{\partial \textbf{y}} = \textbf{B}^{\mathsf{T}}(\textbf{A} + \textbf{A}^{\mathsf{T}})(\textbf{x} + \textbf{B}\textbf{y}),$$

and also

$$\frac{\partial^2 h(\mathbf{x}, \mathbf{y})}{\partial \mathbf{x} \partial \mathbf{x}^\mathsf{T}} = \mathbf{A} + \mathbf{A}^\mathsf{T} \quad \text{and} \quad \frac{\partial^2 h(\mathbf{x}, \mathbf{y})}{\partial \mathbf{y} \partial \mathbf{y}^\mathsf{T}} = \mathbf{B}^\mathsf{T} (\mathbf{A} + \mathbf{A}^\mathsf{T}) \mathbf{B}$$

• Note that above we use partial derivative notation, i.e. ∂ , as opposed to gradient operator notation, i.e. ∇ , as the derivatives are not w.r.t. all variables.

Challenges III

 Go to Challenges III of the week 6 lecture 1 challenges at https://byoungman.github.io/MTH3045/challenges Week 6 lecture 2

Example: Maximum likelihood estimates of regression coefficients in the normal linear model via matrix calculus I

- ullet Consider the normal linear model ${f Y}={f X}eta+arepsilon$
 - $\mathbf{Y} = (Y_1, \dots, Y_n)^T$
 - **X** is an $n \times (p+1)$ design matrix
 - β is a (p+1)-vector of regression coefficients
 - $\varepsilon = (\varepsilon_1, \dots, \varepsilon_n)^T$ with independent $\varepsilon_i \sim N(0, \sigma^2)$
- The maximum likelihood estimate of β , denoted $\hat{\beta}$, minimises the RSS, i.e. minimises

$$(\mathbf{y} - \mathbf{X}\boldsymbol{\beta})^{\mathsf{T}}(\mathbf{y} - \mathbf{X}\boldsymbol{\beta})$$

if we observe $\mathbf{y} = (y_1, \dots, y_n)^\mathsf{T}$

• Differentiating w.r.t. β gives

$$-\mathbf{X}^{\mathsf{T}}(\mathbf{y} - \mathbf{X}\boldsymbol{\beta}) - (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})^{\mathsf{T}}\mathbf{X}$$

Example: Maximum likelihood estimates of regression coefficients in the normal linear model via matrix calculus II

Then

$$-\mathbf{X}^{\mathsf{T}}(\mathbf{y} - \mathbf{X}\boldsymbol{\beta}) - (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})^{\mathsf{T}}\mathbf{X}$$

simplifies to

$$-2\mathbf{X}^{\mathsf{T}}(\mathbf{y}-\mathbf{X}\boldsymbol{\beta})$$

as $\mathbf{X}^\mathsf{T}(\mathbf{y} - \mathbf{X}eta)$ and $(\mathbf{y} - \mathbf{X}eta)^\mathsf{T}\mathbf{X}$ are both *n*-vectors

ullet The derivative of the RSS is zero at \hat{eta} and so

$$-2\mathbf{X}^{\mathsf{T}}(\mathbf{y}-\mathbf{X}\hat{\boldsymbol{\beta}})=0$$

• Therefore $\hat{\beta}$ is the solution of

$$\mathbf{X}^\mathsf{T}\mathbf{X}\hat{oldsymbol{eta}} = \mathbf{X}^\mathsf{T}\mathbf{y}$$

Alternatively

$$\hat{\boldsymbol{\beta}} = (\mathbf{X}^\mathsf{T}\mathbf{X})^{-1}\mathbf{X}^\mathsf{T}\mathbf{y}$$

as we were given in MTH2006

Finite-differencing

- Consider again $f(\mathbf{x})$, a function of vector \mathbf{x}
- **Definition**: Consider $f: \mathbb{R}^n \to \mathbb{R}$ for *n*-vector **x**
- Let, e_i be the n-vector comprising entirely zeros, except for its ith element, which is one
- Then the **partial derivative** of $f(\mathbf{x})$ w.r.t. x_i , the *i*th element of \mathbf{x} , is

$$\frac{\partial f(\mathbf{x})}{\partial x_i} = \lim_{h \to 0} \frac{f(\mathbf{x} + h\mathbf{e}_i) - f(\mathbf{x})}{h}$$

 The preceding definition leads us to the finite-difference partial derivative approximation

$$\frac{\partial f(\mathbf{x})}{\partial x_i} \simeq \frac{f(\mathbf{x} + \delta \mathbf{e}_i) - f(\mathbf{x})}{\delta},$$

where δ is small

Example: Finite-differencing of sin(x) I

- Use finite-differencing to approximate the derivative of $f(x) = \sin(x)$ for $x \in [0, 2\pi]$, and compare its accuracy to the true derivative
- First, let's calculate f and its *true* derivative, i.e. $f'(x) = \cos(x)$, and store this as partial0 for $\{x_i\}$, $i = 1, \ldots, 100$, a set of equally-spaced points on $[0, 2\pi]$

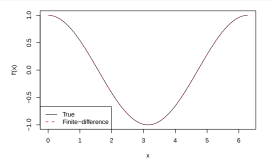
```
n <- 100
x <- seq(0, 2 * pi, 1 = 100)
f <- sin(x)
partial0 <- cos(x)</pre>
```

• Now we'll set $\delta=10^{-6}$ and calculate $x_i+\delta$, for each x_i , i.e. each element of x, so that the finite-difference approximation to the derivative is given by $[\sin(x_i+10^{-6})-\sin(x_i)]/10^{-6}$, which is calculated below and stored as partial1

```
delta1 <- 1e-6
x1 <- x + delta1
f1 <- sin(x1)
partial1 <- (f1 - f) / delta1</pre>
```

Example: Finite-differencing of sin(x) II

• We'll then plot the f'(x) against it finite-difference approximation matplot(x, cbind(partial0, partial1), type = 'l', xlab = 'x', ylab = "f'(x)") legend('bottomleft', lty = 1:2, col = 1:2, lwd = 1:2, legend = c('True', 'Finite-difference'))



 In fact, the true derivative and its finite-difference approximation are so similar that's it's difficult to distinguish the two, but they're both there in the plot!

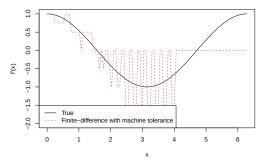
Example: Finite-differencing of sin(x) III

- ullet We might be tempted to choose δ as small as possible
- Suppose we were to repeat Example 4.3 with $\delta=\epsilon_{\it m}$, i.e. R's machine tolerance
- The following calculates the finite-difference approximation as partial2

```
delta2 <- .Machine$double.eps
x2 <- x + delta2
f2 <- sin(x2)
partial2 <- (f2 - f) / delta2</pre>
```

Example: Finite-differencing of sin(x) III

• The the following plots this against the true value of f'(x)



- Using $\delta = \epsilon_m$ gives a terrible approximation to f'(x), which gets worse as x increases
- We've actually encountered an example calculation error, which was introduced in Chapter 2

Challenges I

 Go to Challenges I of the week 6 lecture 2 challenges at https://byoungman.github.io/MTH3045/challenges

Example: Finite differencing of the multivariate Normal log-likelihood I

- Find $\partial \log f(\mathbf{y} \mid \boldsymbol{\mu}, \boldsymbol{\Sigma})/\partial y_i$ analytically, for $i = 1, \dots, p$, where $f(\mathbf{y} \mid \boldsymbol{\mu}, \boldsymbol{\Sigma})$ is the $MVN_p(\boldsymbol{\mu}, \boldsymbol{\Sigma})$ pdf, for arbitrary \mathbf{y} , $\boldsymbol{\mu}$ and $\boldsymbol{\Sigma}$
- Evaluate this for \mathbf{y} , μ and Σ as in Example 3.2
- Then approximate the same derivative using finite-differencing
- The logarithm of the $MVN_p(\mu, \Sigma)$ pdf is given by

$$\log f(\mathbf{y} \mid \boldsymbol{\mu}, \boldsymbol{\Sigma}) = -\frac{1}{2} \left[p \log(2\pi) + \log(|\boldsymbol{\Sigma}|) + (\mathbf{y} - \boldsymbol{\mu})^\mathsf{T} \boldsymbol{\Sigma}^{-1} (\mathbf{y} - \boldsymbol{\mu}) \right]$$

which we know, from Example 3.13, we can evaluate with dmvn3()

Using the above properties

$$rac{\partial \log f(\mathbf{y} \mid \mu, \mathbf{\Sigma})}{\partial \mathbf{y}} = -\mathbf{\Sigma}^{-1}(\mathbf{y} - \mu)$$

since Σ is symmetric

Example: Finite differencing of the multivariate Normal log-likelihood II

• We can evaluate this for ${f y},~\mu$ and ${f \Sigma}$ as in Example 3.2 with the following

```
y <- c(.7, 1.3, 2.6)

mu <- 1:3

Sigma <- matrix(c(4, 2, 1, 2, 3, 2, 1, 2, 2), 3, 3)

deriv1 <- -solve(Sigma, y - mu)
```

which gives

```
as.vector(deriv1)
```

```
## [1] -0.08 0.38 -0.14
```

Example: Finite differencing of the multivariate Normal log-likelihood III

 To approximate the derivative by finite-differencing, it makes sense to write a multi-purpose function for finite differencing, which we'll call fd()

```
fd \leftarrow function(x, f, delta = 1e-6, ...) {
  # Function to evaluate derivative w.r.t. vector by finite-differencing
  # x is a p-vector
  # fn is the function for which the derivative is being calculated
  # delta is the finite-differencing step, which defaults to 10^{-6}
  # returns a vector of length x
  f0 \leftarrow f(x, ...)
  p <- length(x)
  f1 <- numeric(p)
  for (i in 1:p) {
    ei <- replace(numeric(p), i, 1)
    f1[i] \leftarrow f(x + delta * ei, ...)
  (f1 - f0) / delta
```

Example: Finite differencing of the multivariate Normal log-likelihood IV

Then we can use this with dmvn3() with the following

```
deriv2 <- fd(y, dmvn3, mu = mu, Sigma = Sigma)
```

```
which gives
deriv2
```

```
## [1] -0.0800002 0.3799993 -0.1400008

and is the same as the analytical result

all.equal(deriv1, deriv2)
```

```
## [1] "Mean relative difference: 2.832689e-06"
once we allow for error in the finite-difference approximation
```

Challenges II

 Go to Challenges II of the week 6 lecture 2 challenges at https://byoungman.github.io/MTH3045/challenges Week 6 lecture 3

Quadrature

- Another common requirement in statistics is that some integral needs to be evaluated
- To start, let's consider a simple integral of the form

$$I = \int_{a}^{b} f(x) \mathrm{d}x$$

- We'll first take a look at some deterministic approaches to numerically evaluating integrals
- In fact, these all boil down to assuming that

$$I \simeq \sum_{i=1}^{N} w_i f(x_i^*)$$

for some weights w_i and nodes x_i^* , i = 1, ..., N

 Note that here we're considering the so-called composite approach to approximating an interval, i.e. in which a rule is applied over a collection of sub-intervals

Midpoint rule

- Perhaps the first numerical integration scheme we come across is the midpoint rule
- Put simply, we divide [a, b] into N equally-sized intervals, and use the midpoints of these as the nodes, x_i^*
- This gives

$$\int_a^b f(x) dx \simeq h \sum_{i=1}^N f(x_i^*),$$

where $x_i^* = a + (i - 0.5)(b - a)/N$ and h = (b - a)/N

- The error in the approximation is $O(h^2)$
- Thus more intervals reduces h and gives a more accurate approximation
- We can measure accuracy through relative absolute error
- Definition: The relative absolute error, or sometimes just relative error, of an estimate of some true value is given by

Example: Midpoint rule I

- Consider the integral $\int_0^1 \exp(x) dx = \exp(1) 1 \simeq 1.7182818$
- ullet Use R and the midpoint rule to estimate the integral with ${\it N}=10$, 100 and 1000
- Then compare the relative absolute error of each
- We'll start by calculating the true value of the integral, which we'll store as true

```
true <- exp(1) - 1
```

Then we'll store the values of N that we're testing as N_vals
 N_vals <- 10^c(1:3)

Example: Midpoint rule II

- The following then creates a vector, midpoint, in which to store the integral approximations, and calculates the approximations with a for loop
- Inside the loop the integration nodes (i.e. the midpoints) and *h* are calculated

```
midpoint <- numeric(length(N_vals))
for (i in 1:length(N_vals)) {
   N <- N_vals[i]
   nodes <- (1:N - .5) / N
   h <- 1 / N
   midpoint[i] <- h * sum(exp(nodes))
}
midpoint</pre>
```

```
## [1] 1.717566 1.718275 1.718282
```

Example: Midpoint rule III

 The relative absolute error for each is then given in the vector rel_err_mp below

```
rel_err_mp <- abs((true - midpoint) / true)
rel_err_mp</pre>
```

```
## [1] 4.165452e-04 4.166655e-06 4.166667e-08
```

• We clearly see that the absolute error reduces by two factors of ten for each factor of ten increase in N, which is consistent with the above comment of $O(h^2)$ error, where here h=1/N

Challenges III

 Go to Challenges III of the week 6 lecture 2 challenges at https://byoungman.github.io/MTH3045/challenges

Simpson's rule

- The midpoint rule works simply by approximating f(x) over a sub-interval of [a, b] by a horizontal line
- The trapezium rule (which we'll overlook) assumes a straight line
- Simpson's rule is derived from a quadratic approximation and given by

$$\int_{a}^{b} f(x) dx \simeq \frac{h}{6} \left(f(a) + 4 \sum_{i=1}^{N} f(x_{1i}^{*}) + 2 \sum_{i=1}^{N-1} f(x_{2i}^{*}) + f(b) \right),$$

where
$$x_{1i}^* = a + h(2i - 1)/2$$
, $x_{2i}^* = a + ih$ and $h = (b - a)/N$

- Note that Simpson's rule requires ${\it N}+1$ more evaluations of ${\it f}$ than the midpoint rule
 - however, a benefit of those extra evaluations is that its error reduces to $O(h^4)$

Example: Simpson's rule I

- Now use R and Simpson's rule to approximate the integral $\int_0^1 \exp(x) \mathrm{d}x = \exp(1) 1$ with N = 10, 100 and 1000, compare the relative absolute error for each, and against those of the midpoint rule in Example 4.5
- We already have true and N_vals from Example 4.5, and we can use a similar for loop to approximate the integral using Simpson's rule
- The main difference is that we create two sets of nodes, nodes1 and nodes2, which correspond to the x_{1i} s and x_{2i} s in Equation (4.1), respectively

Example: Simpson's rule II

• The integral approximations are stored as simpson

```
simpson <- numeric(length(N_vals))
N_vals <- 10^c(1:3)
for (i in 1:length(N_vals)) {
    N <- N_vals[i]
    h <- 1 / N
    simpson[i] <- 1 + exp(1)
    nodes1 <- h * (2*c(1:N) - 1) / 2
    simpson[i] <- simpson[i] + 4 * sum(exp(nodes1))
    nodes2 <- h * c(1:(N - 1))
    simpson[i] <- simpson[i] + 2 * sum(exp(nodes2))
    simpson[i] <- h * simpson[i] / 6
}
print(simpson, digits = 12)</pre>
```

```
## [1] 1.71828188810 1.71828182847 1.71828182846
```

 \bullet We print this to 11 decimal places so we can see where the approximations changes with N

Example: Simpson's rule III

• Finally we calculate the relative absolute errors, rel_err_simp,

```
rel_err_simp <- abs((true - simpson) / true)
rel_err_simp</pre>
```

```
## [1] 3.471189e-08 3.472270e-12 6.461239e-16
```

- We see a dramatic improvement in the accuracy of approximation that Simpson's rule brings, with relative absolute errors of the same order of magnitude as those form the midpoint rule using N=1000 achieved with N=10 for Simpson's rule
- Note, though, that for given N, Simpson's rule requires N+1 more evaluations of f()

Challenges I

 Go to Challenges I of the week 6 lecture 3 challenges at https://byoungman.github.io/MTH3045/challenges

Gaussian quadrature I

- We've seen that Simpson's rule can considerably improve on the midpoint rule for approximating integrals
- However, we might still consider both restrictive in that they consider an equally-spaced set of nodes
- **Definition**: Consider g(x), a polynomial of degree 2N-1, and a fixed weight function w(x)
- Then, the Gauss-Legendre quadrature rule states that

$$\int_a^b w(x)g(x)dx = \sum_{i=1}^N w_ig(x_i),$$

where, for i = 1, ..., N, w_i and x_i depend on w(x), a and b, but not g(x)

Gaussian quadrature II

- The Gauss-Legendre quadrature rule is the motivation for Gaussian quadrature, whereby we assume that the integral we're interested in can be well-approximated by a polynomial
- This results in the approximation

$$\int_a^b f(x) dx \simeq \sum_{i=1}^N w_i f(x_i)$$

for a fixed set of x values, x_i with corresponding weights w_i , for i = 1, ..., N

- There are many rules for choosing the weights, w_i , but (perhaps fortunately) we won't go into them in detail in MTH3045
- Instead, we'll just consider the function pracma::gaussLegendre() (for which you'll need to install the pracma package), where pracma::gaussLegendre(N, a, b) produces N nodes and corresponding weights on the interval [a,b], with N = N, a = a and b = b

Gaussian quadrature III

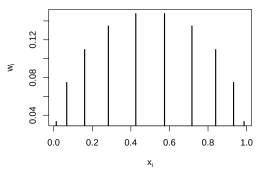
ullet The following produces nodes and weights for ${\it N}=10$ on [0,1]

```
gq <- pracma::gaussLegendre(10, 0, 1)
gq

## $x
## [1] 0.01304674 0.06746832 0.16029522 0.28330230 0.42556283 0.57443717
## [7] 0.71669770 0.83970478 0.93253168 0.98695326
##
## $w
## [1] 0.03333567 0.07472567 0.10954318 0.13463336 0.14776211 0.14776211
## [7] 0.13463336 0.10954318 0.07472567 0.03333567</pre>
```

Gaussian quadrature IV

The plots below shows the nodes and their weights



- ullet The nodes are spread further apart towards the middle of the [0,1] range, but given more weight
- Note that pracma::gaussLegendre() is named so because it implements Gauss-Legendre quadrature, i.e. Gaussian quadrature with Legendre polynomials¹

¹Wikipedia has a useful pages on Gaussian quadrature and on Legendre polynomials, should you want to read more on them

Example: Gaussian quadrature I

- Now use R and Gauss-Legendre quadrature to approximate the integral $\int_0^1 \exp(x) dx$ with N=10
- Explore what value of N gives a comparable estimate to that of the midpoint rule with N=100 based on relative absolute error
- We can re-use true from Example 4.5 and then we'll consider N = 10, 4 and 3, which we'll call N_vals, and store the resulting integral approximations in gauss.

```
N_vals <- c(10, 4, 3)
gauss <- numeric(length(N_vals))
for (i in 1:length(N_vals)) {
  N <- N_vals[i]
  xw <- pracma::gaussLegendre(N, 0, 1)
  gauss[i] <- sum(xw$w * exp(xw$x))
}
gauss</pre>
```

```
## [1] 1.718282 1.718282 1.718281
```

Example: Gaussian quadrature II

The relative absolute errors, rel_err_gauss,

```
rel_err_gauss <- abs((true - gauss) / true)
rel_err_gauss</pre>
```

```
## [1] 2.584496e-16 5.429651e-10 4.795992e-07
```

show that, having considered N=3,4,10, choosing N=3 for Gaussian quadrature gives closest relative absolute error to that of the midpoint rule with N=100, which really is quite impressive

- Note, though, that $f(x) = \exp(x)$ is a very smooth function
 - for wiggler functions, larger N is likely to be needed, and improvements in performance, such as Gaussian quadrature over the midpoint rule, might be significantly less

Challenges II

 Go to Challenges II of the week 6 lecture 3 challenges at https://byoungman.github.io/MTH3045/challenges

Example: Poisson marginal approximation using Gaussian quadrature I

- Consider a single random variable $Y \mid \lambda \sim \mathsf{Poisson}(\lambda)$, where we can characterise our prior beliefs about λ as $\lambda \sim \mathsf{N}(\mu, \sigma^2)$
- Use Gaussian quadrature with ${\it N}=7$ to estimate the marginal pdf of ${\it Y}$ if $\mu=10$ and $\sigma=3$
- The marginal pdf of Y is given by

$$f(y) = \int_{-\infty}^{\infty} f(y \mid \lambda) f(\lambda) d\lambda$$

 Remark: The three sigma rule is a heuristic rule of thumb that 99.7% of values lie within three standard deviations of the mean

Example: Poisson marginal approximation using Gaussian quadrature II

- Hence for the N(10, 3²) distribution we should expect 99.7% of values to lie within $10 \pm 3 \times 3$
- Hence we'll take this as our range for the Gaussian quadrature nodes.

```
m_{11} < -10
sigma <- 3
N \leftarrow 7 # no. of nodes
xw <- pracma::gaussLegendre(</pre>
        Ν,
        mu - 3 * sigma, # left-hand end
        mu + 3 * sigma # right-hand end
XW
## $x
## [1]
        1.458029 3.326219 6.347394 10.000000 13.652606 16.673781 18.541971
##
## $w
## [1] 1.165365 2.517349 3.436470 3.761633 3.436470 2.517349 1.165365
```

which are stored as xwx with corresponding weights xwx, w_1, \ldots, w_N

Example: Poisson marginal approximation using Gaussian quadrature III

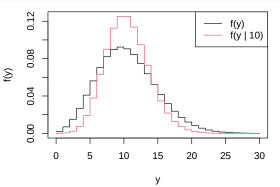
• Next we want a set of values at which to evaluate f(y), and for this we'll choose $0, 1, \ldots, 30$, which we can create in R with $y_vals < 0.30$

• Then we can estimate f(y) as

$$\hat{f}(y) \simeq \sum_{i=1}^{N} w_i f(y \mid \lambda_i^*) f(\lambda_i^*)$$

Example: Poisson marginal approximation using Gaussian quadrature IV

• Finally, we'll plot $\hat{f}(y)$ against the pdf of the Poisson(10) distribution



• f(y) is broader than $f(y \mid 10)$, which is to be expected given that f(y) integrates out the variability in λ given by the N(10, 3²) distribution

One-dimensional numerical integration in R

- \bullet Unsurprisingly, R has a function for one-dimensional numerical integration
- It's called integrate()
- It uses a method that builds on Gaussian quadrature, but we won't go into its details
- Use of integrate(), however, is fairly straightforward

Example: Integration with integrate()

- Evaluate the integral $\int_0^1 \exp(x) dx = \exp(1) 1$ using R's integrate() function with N = 10 and report its relative absolute error
- We can use the following code, where the first argument to integrate() is the function we're integrating, the second and third are the lower and upper ranges of the definite integral, and subdivisions is the maximum number of nodes to use in the approximation, which defaults to 100.

```
true <- \exp(1) - 1
estimate <- integrate(function(x) \exp(x), 0, 1, subdivisions = 10)
estimate
```

```
## 1.718282 with absolute error < 1.9e-14
rel_err <- abs((true - estimate$value) / true)
rel_err</pre>
```

```
## [1] 1.292248e-16
```

- Note above that the absolute error is similarly tiny to that of Gaussian quadrature above
- The values themselves, being so close to the machine tolerance, are incomparable
 - but we can be sure that the approximation is incredibly accurate

References

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