integration

February 24, 2017

1 Numerical Differentiation and Integration

ScPo Computational Economics 2017

1.1 Derivatives

- 1. Finite Differencing: a numerical approximation
 - Based on Taylor's Theorem
 - Observe variation in function values from evaluating it at "close" points.
 - Forward Differencing and Central Differencing

2. Automatic Differentiation

- Breaks down the actual code that defines a function and performs elementary differentiation rules, after disecting expressions via the chain rule.
- This produces **analytic** derivatives, i.e. there is **no** approximation error.
- This is the future.

3. Symbolic Differentiation

- Some languages (most notably Mathematica) support symbolic algebra. Very useful sometimes if one needs to work through complicated expressions.
- Not very useful for high computational demands, i.e. repeated computation of derivatives in an optimization routine.

1.2 Finite Differences

• Consider the definition of the derivative of *f* at point *x*:

$$f'(x) = \lim_{h \to 0} \frac{f(x+h) - f(x)}{h}$$

• The simplest way to calculate a numerical derivative is to replicate this computation for small *h* with:

$$f'(x) \approx \frac{f(x+h) - f(x)}{h}$$
, h small.

- This is known as the Forward Difference approach.
- There are different approaches, e.g. the central difference approach does

$$f'(x) \approx \frac{f(x+h) - f(x-h)}{2h}$$
, h small.

How does this perform?

```
In [1]: using Plots
    pyplot()
    f(x) = 2 - x^2
    c = -0.75
    sec_line(h) = x -> f(c) + (f(c + h) - f(c))/h * (x - c)
    plot([f, sec_line(1), sec_line(.5), sec_line(.25), sec_line(.05)], -1, 1)
```

/Users/florian.oswald/.julia/v0.5/Conda/deps/usr/lib/python2.7/site-packages/matplotlib/font_man warnings.warn('Matplotlib is building the font cache using fc-list. This may take a moment.')

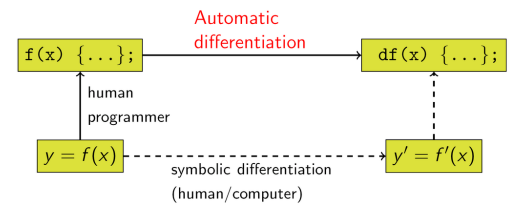
• What's the problem? Well, what is *small*?

1.2.1 Finite Differences: what's the right step size h?

- Theoretically, we would like to have *h* as small as possible, since we want to approximate the limit at zero.
- In practice, on a computer, there is a limit to this. There is a smallest representable number, as we know.
- eps().
- One can show that the optimal step size is $h = \sqrt{\text{eps}()}$

1.3 Automatic Differentiation (AD)

- 2 modes: Forward and Reverse Mode.
- The basic idea is that the derivative of any function can be decomposed into some basic algebraic operations.
- The wikipedia page is informative



By Berland at en.wikipedia [Public domain], from Wikimedia Commons

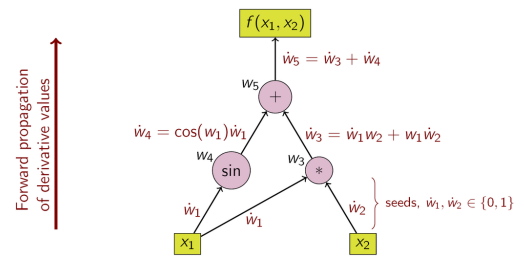
1.4 Example

- Suppose we want to differentiate $f(x_1, x_2) = x_1x_2 + \sin x_1$
- We label subexpressions by w_i as follows:

$$f(x_1, x_2) = x_1 x_2 + \sin x_1$$

= $w_1 w_2 + \sin w_1$
= $w_3 + w_4$
= w_5

- Computation of the partial derivative starts with the seed value, i.e. $\dot{w}_1 = \frac{\partial x_1}{\partial x_1} = 1$.
- We store for each subexpression both the value and the derivative, i.e. (w_i, w_i)
- We then sweep through the expression tree as in this picture:



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1.5 AD in Julia

- The organisation here is http://www.juliadiff.org
- There are many packages to perform differentiation with Julia here.
- Many packages rely on the machinery here.
- Let's quickly look at https://github.com/JuliaDiff/ForwardDiff.jl

```
0.382642 -0.323699
                    0.397432 4.04971
                                        0.667225
                                        0.68702
0.394007
          0.397432 -0.288224 4.16933
4.01658
          4.04971
                    4.16933
                              1.8859
                                        7.00117
0.661422
          0.667225
                    0.68702
                              7.00117 -0.00985418
```

• The authors provide some benchmarks. Let's run those:

include(joinpath(Pkg.dir("ForwardDiff"), "benchmark", "ForwardDiffBenchmarks.jl"))

1.6 Numerical Approximation of Integrals

- We will focus on methods that represent integrals as weighted sums.
- The typical representation will look like:

$$E[G(\epsilon)] = \int_{\mathbb{R}^N} G(\epsilon) w(\epsilon) d\epsilon \approx \sum_{j=1}^J \omega_j G(\epsilon_j)$$

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- *N* is the dimensionality of the integration problem.
- $G: \mathbb{R}^N \mapsto \mathbb{R}$ is the function we want to integrate wrt $\epsilon \in \mathbb{R}^N$.
- w is a density function s.t. $\int_{\mathbb{R}^n} w(\epsilon) d\epsilon = 1$.
- ω are weights such that (most of the time) $\sum_{j=1}^{J} \omega_j = 1$. We will look at normal shocks $\epsilon \sim N(0_N, I_N)$ in that case, $w(\epsilon) = (2\pi)^{-N/2} \exp\left(-\frac{1}{2}\epsilon^T\epsilon\right)$

- I_N is the n by n identity matrix, i.e. there is no correlation among the shocks for now.
- Other random processes will require different weighting functions, but the principle is identical.
- For now, let's say that N = 1

1.7 Quadrature Rules

- We focus exclusively on those and leave Simpson and Newton Cowtes formulas out.
 - This is because Quadrature is the method that in many situations gives highes accuracy with lowest computational cost.
- Quadrature provides a rule to compute weights w_i and nodes ϵ_i .
- There are many different quadrature rules.
- They differ in their domain and weighting function.
- https://en.wikipedia.org/wiki/Gaussian_quadrature
- In general, we can convert our function domain to a rule-specific domain with change of variables.

1.8 Gauss-Hermite: Expectation of a Normally Distributed Variable

- There are many different rules, all specific to a certain random process.
- Gauss-Hermite is designed for an integral of the form

$$\int_{-\infty}^{+\infty} e^{-x^2} G(x) dx$$

and where we would approximate

$$\int_{-\infty}^{+\infty} e^{-x^2} f(x) dx \approx \sum_{i=1}^{n} \omega_i G(x_i)$$

• Now, let's say we want to approximate the expected value of function f when it's argument $z \sim N(u, \sigma^2)$:

$$E[f(z)] = \int_{-\infty}^{+\infty} \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(z-\mu)^2}{2\sigma^2}\right) f(z) dz$$

1.9 Gauss-Hermite: Expectation of a Normally Distributed Variable

• The rule is defined for *x* however. We need to transform *z*:

$$x = \frac{(z-\mu)^2}{2\sigma^2} \Rightarrow z = \sqrt{2}\sigma x + \mu$$

This gives us now (just plug in for z)

$$E[f(z)] = \int_{-\infty}^{+\infty} \frac{1}{\sqrt{\pi}} \exp(-x^2) f(\sqrt{2}\sigma x + \mu) dx$$

• And thus, our approximation to this, using weights ω_i and nodes x_i is

$$E[f(z)] \approx \sum_{j=1}^{J} \frac{1}{\sqrt{\pi}} \omega_j f(\sqrt{2}\sigma x_j + \mu)$$

1.10 Using Quadrature in Julia

• https://github.com/ajt60gaibb/FastGaussQuadrature.jl

In [3]: #Pkg.add("FastGaussQuadrature")

using FastGaussQuadrature

using DataFrames

integ = DataFrame(Rule=Symbol[Symbol(x) for x in keys(rules)],nodes=[x[1] for x in value

```
Out[3]: 4E3 DataFrames.DataFrame
         Row
              Rule
                          nodes
                          [-1.0,0.0,1.0]
         1
              lobatto
         2
                          [-1.22474, -8.88178e-16, 1.22474]
              hermite
         3
                          [-0.774597,0.0,0.774597]
              legendre
         4
              chebyshev
                         [-0.866025,6.12323e-17,0.866025]
         Row
              weights
         1
              [0.333333,1.33333,0.333333]
         2
              [0.295409,1.18164,0.295409]
         3
              [0.555556,0.888889,0.555556]
         4
              [1.0472,1.0472,1.0472]
```

1.11 Quadrature in more dimensions: Product Rule

- If we have N > 1, we can use the product rule: this just takes the kronecker product of all univariate rules.
- This works well as long as N is not too large. The number of required function evaluations grows exponentially.

$$E[G(\epsilon)] = \int_{\mathbb{R}^N} G(\epsilon) w(\epsilon) d\epsilon \approx \sum_{j_1=1}^{J_1} \cdots \sum_{j_N=1}^{J_N} \omega_{j_1}^1 \cdots \omega_{j_N}^N G(\epsilon_{j_1}^1, \dots, \epsilon_{j_N}^N)$$

where $\omega_{j_1}^1$ stands for weight index j_1 in dimension 1, same for ϵ .

• Total number of nodes: $J = J_1 J_2 \cdots J_N$, and J_i can differ from J_k .

1.11.1 Example for N = 3

- Suppose we have $e^i \sim N(0,1)$, i = 1,2,3 as three uncorrelated random variables.
- Let's take I=3 points in all dimensions, so that in total we have $I^N=27$ points.
- We have the nodes and weights from before in rules ["hermite"].

```
In [4]: nodes = Any[]
        push! (nodes,repeat(rules["hermite"][1],inner=[1],outer=[9]))
        push!(nodes,repeat(rules["hermite"][1],inner=[3],outer=[3]))
        push! (nodes,repeat(rules["hermite"][1],inner=[9],outer=[1]))
        weights = kron(rules["hermite"][2],kron(rules["hermite"][2],rules["hermite"][2]))
        df = hcat(DataFrame(weights=weights),DataFrame(nodes,[:dim1,:dim2,:dim3]))
Out[4]: 27C4 DataFrames.DataFrame
         Row weights
                         dim1
                                       dim2
                                                     dim3
              0.0257793 -1.22474
                                       -1.22474
                                                     -1.22474
         1
         2
              0.103117
                         -8.88178e-16 -1.22474
                                                     -1.22474
                                       -1.22474
         3
              0.0257793 1.22474
                                                     -1.22474
         4
                                       -8.88178e-16 -1.22474
              0.103117
                        -1.22474
```

```
5
    0.412469
                -8.88178e-16 -8.88178e-16 -1.22474
                1.22474
                              -8.88178e-16 -1.22474
6
    0.103117
7
    0.0257793
               -1.22474
                              1.22474
                                            -1.22474
8
                -8.88178e-16 1.22474
                                            -1.22474
    0.103117
9
    0.0257793 1.22474
                              1.22474
                                            -1.22474
    0.103117
                -1.22474
                              -1.22474
                                            -8.88178e-16
10
11
    0.412469
                -8.88178e-16 -1.22474
                                            -8.88178e-16
    0.103117
                -1.22474
                              1.22474
                                            -8.88178e-16
16
17
    0.412469
                -8.88178e-16 1.22474
                                            -8.88178e-16
18
    0.103117
                1.22474
                              1.22474
                                            -8.88178e-16
    0.0257793 -1.22474
19
                              -1.22474
                                            1.22474
20
    0.103117
                -8.88178e-16 -1.22474
                                            1.22474
21
    0.0257793 1.22474
                              -1.22474
                                            1.22474
22
    0.103117
                -1.22474
                              -8.88178e-16
                                            1.22474
23
    0.412469
                -8.88178e-16 -8.88178e-16 1.22474
24
    0.103117
                1.22474
                              -8.88178e-16 1.22474
25
    0.0257793 -1.22474
                              1.22474
                                            1.22474
26
    0.103117
                -8.88178e-16 1.22474
                                            1.22474
27
    0.0257793 1.22474
                              1.22474
                                            1.22474
```

 Imagine you had a function g defined on those 3 dims: in order to approximate the integral, you would have to evaluate g at all combinations of dimx, multiply with the corresponding weight, and sum.

1.11.2 Alternatives to the Product Rule

- Monomial Rules: They grow only linearly.
- Please refer to [@judd-book] for more details.

1.12 Monte Carlo Integration

- A widely used method is to just draw N points randomly from the space of the shock ϵ , and to assign equal weights $\omega_j = \frac{1}{N}$ to all of them.
- The expectation is then

$$E[G(\epsilon)] \approx \frac{1}{N} \sum_{j=1}^{N} G(\epsilon_j)$$

- This in general a very inefficient method.
- Particularly in more than 1 dimensions, the number of points needed for good accuracy is very large.

1.13 Quasi Monte Carlo Integration

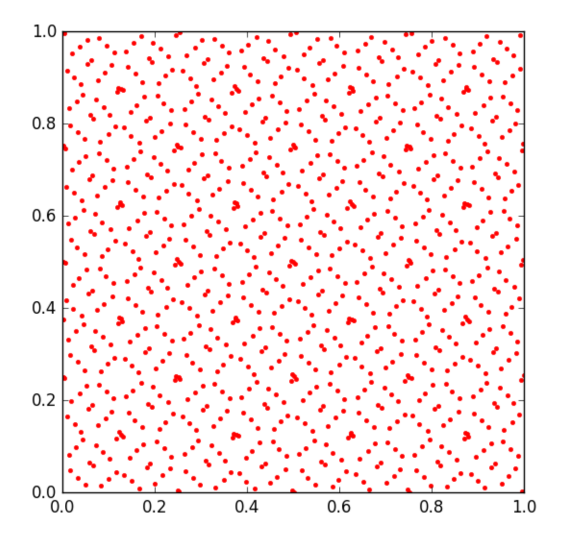
- Uses non-product techniques to construct a grid of uniformly spaced points.
- The researcher controlls the number of points.
- We need to construct equidistributed points.
- Typically one uses a low-discrepancy sequence of points, e.g. the Weyl sequence:

• $x_n = nv$ where v is an irrational number and {} stands for the fractional part of a number. for $v = \sqrt{2}$,

$$x_1 = \{1\sqrt{2}\} = \{1.4142\} = 0.4142, x_2 = \{2\sqrt{2}\} = \{2.8242\} = 0.8242,...$$

• Other low-discrepancy sequences are Niederreiter, Haber, Baker or Sobol.

```
In [2]: # Pkg.add("Sobol")
    using Sobol
    using PyPlot
    s = SobolSeq(2)
    p = hcat([next(s) for i = 1:1024]...)'
    subplot(111, aspect="equal")
    plot(p[:,1], p[:,2], "r.")
```



 ${\tt INFO:}$ No packages to install, update or remove

INFO: Package database updated

INFO: METADATA is out-of-date you may not have the latest version of Sobol

INFO: Use `Pkg.update()` to get the latest versions of your packages

Correlated Shocks

- We often face situations where the shocks are in fact correlated.
 - One very typical case is an AR1 process:

$$z_{t+1} = \rho z_t + \varepsilon_t, \varepsilon \sim N(0, \sigma^2)$$

• The general case is again:

$$E[G(\epsilon)] = \int_{\mathbb{R}^N} G(\epsilon) w(\epsilon) d\epsilon \approx \sum_{j_1=1}^{J_1} \cdots \sum_{j_N=1}^{J_N} \omega_{j_1}^1 \cdots \omega_{j_N}^N G(\epsilon_{j_1}^1, \dots, \epsilon_{j_N}^N)$$

- Now $\epsilon \sim N(\mu, \Sigma)$ where Σ is an N by N variance-covariance matrix.
- The multivariate density is

$$w(\epsilon) = (2\pi)^{-N/2} det(\Sigma)^{-1/2} \exp\left(-\frac{1}{2}(\epsilon - \mu)^T (\epsilon - \mu)\right)$$

- We need to perform a change of variables before we can integrate this.
- Given Σ is symmetric and positive semi-definite, it has a Cholesky decomposition,

$$\Sigma = \Omega \Omega^T$$

where Ω is a lower-triangular with strictly positive entries.

• The linear change of variables is then

$$v = \Omega^{-1}(\epsilon - \mu)$$

Plugging this in gives

$$\sum_{j=1}^{J} \omega_j G(\Omega v_j + \mu) \equiv \sum_{j=1}^{J} \omega_j G(\epsilon_j)$$

where $v \sim N(0, I_N)$.

• So, we can follow the exact same steps as with the uncorrelated shocks, but need to adapt the nodes.

1.14 References

• The Integration part of these slides are based on [@maliar-maliar] chapter 5