Scientific Computing: An Introductory Survey

Chapter 8 - Numerical Integration and Differentiation

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Integration

• For $f: \mathbb{R} \to \mathbb{R}$, definite integral over interval [a,b]

$$I(f) = \int_{a}^{b} f(x) \, dx$$

is defined by limit of Riemann sums

$$R_n = \sum_{i=1}^{n} (x_{i+1} - x_i) f(\xi_i)$$

- ullet Riemann integral exists provided integrand f is bounded and continuous almost everywhere
- Absolute condition number of integration with respect to perturbations in integrand is b-a
- Integration is inherently well-conditioned because of its smoothing effect

Quadrature Rules

An n-point quadrature rule has form

$$Q_n(f) = \sum_{i=1}^n w_i f(x_i)$$

- Points x_i are called *nodes* or *abscissas*
- Multipliers w_i are called weights
- Quadrature rule is
 - open if $a < x_1$ and $x_n < b$
 - closed if $a = x_1$ and $x_n = b$

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Method of Undetermined Coefficients

- Alternative derivation of quadrature rule uses method of undetermined coefficients
- To derive n-point rule on interval [a, b], take nodes x_1, \ldots, x_n as given and consider weights w_1, \ldots, w_n as coefficients to be determined
- ullet Force quadrature rule to integrate first n polynomial basis functions exactly, and by linearity, it will then integrate any polynomial of degree n-1 exactly
- Thus we obtain system of moment equations that determines weights for quadrature rule

Outline

- Numerical Integration
- **Numerical Differentiation**
- Richardson Extrapolation

Numerical Quadrature

- Quadrature rule is weighted sum of finite number of sample values of integrand function
- To obtain desired level of accuracy at low cost,
 - How should sample points be chosen?
 - How should their contributions be weighted?
- Computational work is measured by number of evaluations of integrand function required

Quadrature Rules, continued

- Quadrature rules are based on polynomial interpolation
- Integrand function f is sampled at finite set of points
- Polynomial interpolating those points is determined
- Integral of interpolant is taken as estimate for integral of original function
- In practice, interpolating polynomial is not determined explicitly but used to determine weights corresponding to nodes
- If Lagrange is interpolation used, then weights are given by

 $w_i = \int_a^b \ell_i(x), \quad i = 1, \dots, n$ Michael T. Heath Scientific Computing

Example: Undetermined Coefficients

- Derive 3-point rule $Q_3(f) = w_1 f(x_1) + w_2 f(x_2) + w_3 f(x_3)$ on interval [a,b] using monomial basis
- Take $x_1 = a$, $x_2 = (a + b)/2$, and $x_3 = b$ as nodes
- First three monomials are 1, x, and x^2
- Resulting system of moment equations is

$$\begin{aligned} w_1 \cdot 1 + w_2 \cdot 1 + w_3 \cdot 1 &= \int_a^b 1 \, dx = x|_a^b = b - a \\ w_1 \cdot a + w_2 \cdot (a+b)/2 + w_3 \cdot b &= \int_a^b x \, dx = (x^2/2)|_a^b = (b^2 - a^2)/2 \\ w_1 \cdot a^2 + w_2 \cdot ((a+b)/2)^2 + w_3 \cdot b^2 &= \int_a^b x^2 \, dx = (x^3/3)|_a^b = (b^3 - a^3)/3 \end{aligned}$$

$$\begin{bmatrix} 1 & 1 & 1 \\ a & (a+b)/2 & b \\ a^2 & ((a+b)/2)^2 & b^2 \end{bmatrix} \begin{bmatrix} w_1 \\ w_2 \\ w_3 \end{bmatrix} = \begin{bmatrix} b-a \\ (b^2-a^2)/2 \\ (b^3-a^3)/3 \end{bmatrix}$$

Solving system by Gaussian elimination, we obtain weights

$$w_1 = \frac{b-a}{6}$$
, $w_2 = \frac{2(b-a)}{3}$, $w_3 = \frac{b-a}{6}$

which is known as Simpson's rule

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Accuracy of Quadrature Rules

- Quadrature rule is of degree d if it is exact for every polynomial of degree d, but not exact for some polynomial of degree d+1
- By construction, n-point interpolatory quadrature rule is of degree at least n-1
- Rough error bound

$$|I(f) - Q_n(f)| \le \frac{1}{4} h^{n+1} ||f^{(n)}||_{\infty}$$

where $h=\max\{x_{i+1}-x_i:\ i=1,\dots,n-1\}$, shows that $Q_n(f)\to I(f)$ as $n\to\infty$, provided $f^{(n)}$ remains well behaved

• Higher accuracy can be obtained by increasing n or by decreasing h

Stability of Quadrature Rules

 Absolute condition number of quadrature rule is sum of magnitudes of weights,

$$\sum^n |w_i\>$$

- If weights are all nonnegative, then absolute condition number of quadrature rule is b - a, same as that of underlying integral, so rule is stable
- If any weights are negative, then absolute condition number can be much larger, and rule can be unstable

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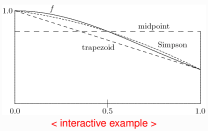
Example: Newton-Cotes Quadrature

Approximate integral $I(f) = \int_0^1 \exp(-x^2) dx \approx 0.746824$

$$M(f) = (1-0)\exp(-1/4) \approx 0.778801$$

$$T(f) = (1/2)[\exp(0) + \exp(-1)] \approx 0.683940$$

$$S(f) = (1/6)[\exp(0) + 4\exp(-1/4) + \exp(-1)] \approx 0.747180$$



Method of Undetermined Coefficients

• More generally, for any n and choice of nodes x_1, \ldots, x_n ,

$$\begin{bmatrix} 1 & 1 & \cdots & 1 \\ x_1 & x_2 & \cdots & x_n \\ \vdots & \vdots & \ddots & \vdots \\ x_1^{n-1} & x_2^{n-1} & \cdots & x_n^{n-1} \end{bmatrix} \begin{bmatrix} w_1 \\ w_2 \\ \vdots \\ w_n \end{bmatrix} = \begin{bmatrix} b-a \\ (b^2-a^2)/2 \\ \vdots \\ (b^n-a^n)/n \end{bmatrix}$$

determines weights w_1, \ldots, w_n

Progressive Quadrature Rules

- Sequence of quadrature rules is progressive if nodes of Q_{n_1} are subset of nodes of Q_{n_2} for $n_2>n_1$
- For progressive rules, function evaluations used in one rule can be reused in another, reducing overall cost
- To attain higher accuracy, we can increase number of points n or subdivide interval into smaller subintervals
- In either case, efficiency is enhanced if successive rules are progressive so that fewer new evaluations of integrand are required

Newton-Cotes Quadrature

Newton-Cotes quadrature rules use equally spaced nodes in interval [a, b]

Midpoint rule

$$M(f) = (b-a)f\left(\frac{a+b}{2}\right)$$

Trapezoid rule

$$T(f) = \frac{b-a}{2}(f(a) + f(b))$$

Simpson's rule

$$S(f) = \frac{b-a}{6} \left(f(a) + 4f\left(\frac{a+b}{2}\right) + f(b) \right)$$

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Error Estimation

ullet Expanding integrand f in Taylor series about midpoint m = (a+b)/2 of interval [a,b],

$$f(x) = f(m) + f'(m)(x - m) + \frac{f''(m)}{2}(x - m)^{2} + \frac{f'''(m)}{6}(x - m)^{3} + \frac{f^{(4)}(m)}{24}(x - m)^{4} + \cdots$$

Integrating from a to b, odd-order terms drop out, yielding

$$I(f) = f(m)(b-a) + \frac{f''(m)}{24}(b-a)^3 + \frac{f^{(4)}(m)}{1920}(b-a)^5 + \cdots$$
$$= M(f) + E(f) + F(f) + \cdots$$

where E(f) and F(f) represent first two terms in error expansion for midpoint rule

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$$I(f) = T(f) - 2E(f) - 4F(f) - \cdots$$

- Thus, provided length of interval is sufficiently small and $f^{(4)}$ is well behaved, midpoint rule is about twice as accurate as trapezoid rule
- Halving length of interval decreases error in either rule by factor of about 1/8



Example: Error Estimation

• We illustrate error estimation by computing approximate value for integral $\int_0^1 x^2 dx = 1/3$

$$\begin{split} M(f) &= (1-0)(1/2)^2 = 1/4 \\ T(f) &= \frac{1-0}{2}(0^2+1^2) = 1/2 \\ E(f) &\approx (T(f)-M(f))/3 = (1/4)/3 = 1/12 \end{split}$$

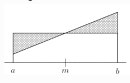
- Error in M(f) is about 1/12, error in T(f) is about -1/6
- Also.

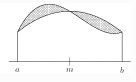
$$S(f) = (2/3)M(f) + (1/3)T(f) = (2/3)(1/4) + (1/3)(1/2) = 1/3$$

which is exact for this integral, as expected

Accuracy of Newton-Cotes Quadrature

- In general, odd-order Newton-Cotes rule gains extra degree beyond that of polynomial interpolant on which it is
- n-point Newton-Cotes rule is of degree n-1 if n is even, but of degree n if n is odd
- This phenomenon is due to cancellation of positive and negative errors





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Clenshaw-Curtis Quadrature

- As with polynomial interpolation, use of Chebyshev points produces better results
- Improved accuracy results from good approximation properties of interpolation at Chebyshev points
- Weights are always positive and approximate integral always converges to exact integral as $n \to \infty$
- Quadrature rules using Chebyshev points are known as Clenshaw-Curtis quadrature, which can be implemented very efficiently
- Clenshaw-Curtis quadrature has many attractive features, but still does not have maximum possible degree for number of nodes used

Error Estimation, continued

 Difference between midpoint and trapezoid rules provides estimate for error in either of them

$$T(f) - M(f) = 3E(f) + 5F(f) + \cdots$$

so

$$E(f) \approx \frac{T(f) - M(f)}{3}$$

 $E(f) \approx \frac{T(f) - M(f)}{3}$ • Weighted combination of midpoint and trapezoid rules eliminates E(f) term from error expansion

$$\begin{split} I(f) &= \frac{2}{3}M(f) + \frac{1}{3}T(f) - \frac{2}{3}F(f) + \cdots \\ &= S(f) - \frac{2}{3}F(f) + \cdots \end{split}$$

which gives alternate derivation for Simpson's rule and estimate for its error

Accuracy of Newton-Cotes Quadrature

- Since n-point Newton-Cotes rule is based on polynomial interpolant of degree n-1, we expect rule to have degree
- Thus, we expect midpoint rule to have degree 0, trapezoid rule degree 1, Simpson's rule degree 2, etc.
- From Taylor series expansion, error for midpoint rule depends on second and higher derivatives of integrand, which vanish for linear as well as constant polynomials
- So midpoint rule integrates linear polynomials exactly, hence its degree is $1\ \mathrm{rather}\ \mathrm{than}\ 0$
- Similarly, error for Simpson's rule depends on fourth and higher derivatives, which vanish for cubics as well as quadratic polynomials, so Simpson's rule is of degree 3

Drawbacks of Newton-Cotes Rules

- Newton-Cotes guadrature rules are simple and often effective, but they have drawbacks
- Using large number of equally spaced nodes may incur erratic behavior associated with high-degree polynomial interpolation (e.g., weights may be negative)
- Indeed, every n-point Newton-Cotes rule with $n \ge 11$ has at least one negative weight, and $\sum_{i=1}^{n} |w_i| \to \infty$ as $n \to \infty$, so Newton-Cotes rules become arbitrarily ill-conditioned
- Newton-Cotes rules are not of highest degree possible for number of nodes used

Gaussian Quadrature

- Gaussian quadrature rules are based on polynomial interpolation, but nodes as well as weights are chosen to maximize degree of resulting rule
- ullet With 2n parameters, we can attain degree of 2n-1
- Gaussian quadrature rules can be derived by method of undetermined coefficients, but resulting system of moment equations that determines nodes and weights is nonlinear
- Also, nodes are usually irrational, even if endpoints of interval are rational
- Although inconvenient for hand computation, nodes and weights are tabulated in advance and stored in subroutine for use on computer

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Example: Gaussian Quadrature Rule

• Derive two-point Gaussian rule on [-1, 1],

$$G_2(f) = w_1 f(x_1) + w_2 f(x_2)$$

where nodes x_i and weights w_i are chosen to maximize degree of resulting rule

- We use method of undetermined coefficients, but now nodes as well as weights are unknown parameters to be determined
- Four parameters are to be determined, so we expect to be able to integrate cubic polynomials exactly, since cubics depend on four parameters



Example, continued

• One solution of this system of four nonlinear equations in four unknowns is given by

$$x_1 = -1/\sqrt{3}$$
, $x_2 = 1/\sqrt{3}$, $w_1 = 1$, $w_2 = 1$

- Another solution reverses signs of x_1 and x_2
- Resulting two-point Gaussian rule has form

$$G_2(f) = f(-1/\sqrt{3}) + f(1/\sqrt{3})$$

and by construction it has degree three

- In general, for each n there is unique n-point Gaussian rule, and it is of degree 2n-1
- Gaussian quadrature rules can also be derived using orthogonal polynomials



Change of Interval, continued

 Many transformations are possible, but simple linear transformation

$$t = \frac{(b-a)x + a\beta - b\alpha}{\beta - \alpha}$$

has advantage of preserving degree of quadrature rule

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Progressive Gaussian Quadrature

- Avoiding this additional work is motivation for Kronrod quadrature rules
- Such rules come in pairs, n-point Gaussian rule G_n , and (2n+1)-point Kronrod rule K_{2n+1} , whose nodes are optimally chosen subject to constraint that all nodes of G_n are reused in K_{2n+1}
- ullet (2n+1)-point Kronrod rule is of degree 3n+1, whereas true (2n+1)-point Gaussian rule would be of degree
- ullet In using Gauss-Kronrod pair, value of K_{2n+1} is taken as approximation to integral, and error estimate is given by

$$(200|G_n - K_{2n+1}|)^{1.5}$$

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Example, continued

 Requiring rule to integrate first four monomials exactly gives moment equations

$$\begin{array}{rcl} w_1+w_2&=&\int_{-1}^1 1\,dx=x|_{-1}^1=2\\ \\ w_1x_1+w_2x_2&=&\int_{-1}^1 x\,dx=(x^2/2)|_{-1}^1=0\\ \\ w_1x_1^2+w_2x_2^2&=&\int_{-1}^1 x^2\,dx=(x^3/3)|_{-1}^1=2/3\\ \\ w_1x_1^3+w_2x_2^3&=&\int_{-1}^1 x^3\,dx=(x^4/4)|_{-1}^1=0 \end{array}$$

Change of Interval

- Gaussian rules are somewhat more difficult to apply than Newton-Cotes rules because weights and nodes are usually derived for some specific interval, such as [-1,1]
- Given interval of integration [a, b] must be transformed into standard interval for which nodes and weights have been tabulated
- To use quadrature rule tabulated on interval $[\alpha, \beta]$,

$$\int_{\alpha}^{\beta} f(x) dx \approx \sum_{i=1}^{n} w_i f(x_i)$$

to approximate integral on interval [a, b],

$$I(g) = \int_{a}^{b} g(t) dt$$

we must change variable from x in $[\alpha, \beta]$ to t in [a, b]

Gaussian Quadrature

- Gaussian quadrature rules have maximal degree and optimal accuracy for number of nodes used
- Weights are always positive and approximate integral always converges to exact integral as $n \to \infty$
- Unfortunately, Gaussian rules of different orders have no nodes in common (except possibly midpoint), so Gaussian rules are *not* progressive
- Thus, estimating error using Gaussian rules of different order requires evaluating integrand function at full set of nodes of both rules

Progressive Gaussian Quadrature, continued

- Because they efficiently provide high accuracy and reliable error estimate, Gauss-Kronrod rules are among most effective methods for numerical quadrature
- They form basis for many quadrature routines available in major software libraries
- Pair (G_7, K_{15}) is commonly used standard
- Patterson quadrature rules further extend this idea by adding 2n+2 optimally chosen nodes to 2n+1 nodes of Kronrod rule K_{2n+1} , yielding progressive rule of degree
- Gauss-Radau and Gauss-Lobatto rules specify one or both endpoints, respectively, as nodes and then choose remaining nodes and all weights to maximize degree

simple quadrature rule in each subinterval

 This approach is equivalent to using piecewise interpolation to derive composite quadrature rule

Composite Quadrature

overall integral

stable

Examples: Composite Quadrature

- Subdivide interval [a, b] into k subintervals of length h = (b - a)/k, letting $x_j = a + jh$, j = 0, ..., k
- Composite midpoint rule

$$M_k(f) = \sum_{j=1}^k (x_j - x_{j-1}) \, f\left(\frac{x_{j-1} + x_j}{2}\right) = h \, \sum_{j=1}^k f\left(\frac{x_{j-1} + x_j}{2}\right)$$

Composite trapezoid rule

$$T_k(f) = \sum_{j=1}^k \frac{(x_j - x_{j-1})}{2} (f(x_{j-1}) + f(x_j))$$

= $h(\frac{1}{2}f(a) + f(x_1) + \dots + f(x_{k-1}) + \frac{1}{2}f(b))$

Adaptive Quadrature

Adaptive Quadrature

- Composite quadrature rule with error estimate suggests simple automatic quadrature procedure
- Continue to subdivide all subintervals, say by half, until overall error estimate falls below desired tolerance
- Such uniform subdivision is grossly inefficient for many integrands, however
- More intelligent approach is adaptive quadrature, in which domain of integration is selectively refined to reflect behavior of particular integrand function

Adaptive Quadrature, continued

- Adaptive quadrature tends to be effective in practice, but it can be fooled: both approximate integral and error estimate can be completely wrong
- Integrand function is sampled at only finite number of points, so significant features of integrand may be missed
- For example, interval of integration may be very wide but "interesting" behavior of integrand may be confined to narrow range
- Sampling by automatic routine may miss interesting part of integrand behavior, and resulting value for integral may be completely wrong

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Integrating Tabular Data

- If integrand is defined only by table of its values at discrete points, then reasonable approach is to integrate piecewise
- For example, integrating piecewise linear interpolant to tabular data gives composite trapezoid rule
- Excellent method for integrating tabular data is to use Hermite cubic or cubic spline interpolation
- In effect, overall integral is computed by integrating each of cubic pieces that make up interpolant
- This facility is provided by many spline interpolation packages

simple rule has degree at least zero

• Alternative to using more nodes and higher degree rule is

Summing partial results then yields approximation to

 Approximate integral converges to exact interval as number of subintervals goes to infinity provided underlying

to subdivide original interval into subintervals, then apply

• Composite rule is always stable if underlying simple rule is

Composite Quadrature Rules, continued

- Composite quadrature offers simple means of estimating error by using two different levels of subdivision, which is easily made progressive
- For example, halving interval length reduces error in midpoint or trapezoid rule by factor of about 1/8
- Halving width of each subinterval means twice as many subintervals are required, so overall reduction in error is by factor of about 1/4
- If h denotes subinterval length, then dominant term in error of composite midpoint or trapezoid rules is $\mathcal{O}(h^2)$
- Dominant term in error of composite Simpson's rule is $\mathcal{O}(h^4)$, so halving subinterval length reduces error by factor of about 1/16

Adaptive Quadrature

Adaptive Quadrature, continued

- Start with pair of quadrature rules whose difference gives error estimate
- Apply both rules on initial interval [a, b]
- If difference between rules exceeds error tolerance, subdivide interval and apply rules in each subinterval
- Continue subdividing subintervals, as necessary, until tolerance is met on all subintervals
- Integrand is sampled densely in regions where it is difficult to integrate and sparsely in regions where it is easy
 - < interactive example >

Adaptive Quadrature, continued

- Adaptive quadrature routine may be inefficient in handling discontinuities in integrand
- For example, adaptive routine may use many function evaluations refining region around discontinuity of
- To prevent this, call quadrature routine separately to compute integral on either side of discontinuity, avoiding need to resolve discontinuity

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Improper Integrals

To compute integral over infinite or semi-infinite interval, several approaches are possible

- Replace infinite limits of integration by carefully chosen finite values
- Transform variable of integration so that new interval is finite, but care must be taken not to introduce singularities
- Use quadrature rule designed for infinite interval

Multiple Integrals

- To evaluate multiple integrals in higher dimensions, only generally viable approach is Monte Carlo method
- ullet Function is sampled at n points distributed randomly in domain of integration, and mean of function values is multiplied by area (or volume, etc.) of domain to obtain estimate for integral
- Error in estimate goes to zero as $1/\sqrt{n}$, so to gain one additional decimal digit of accuracy requires increasing nby factor of 100
- For this reason, Monte Carlo calculations of integrals often require millions of evaluations of integrand

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Integral Equations

Typical integral equation has form

$$\int_{a}^{b} K(s,t)u(t) dt = f(s)$$

where kernel K and right-hand side f are known functions, and unknown function u is to be determined

 Solve numerically by discretizing variables and replacing integral by quadrature rule

$$\sum_{i=1}^{n} w_{j} K(s_{i}, t_{j}) u(t_{j}) = f(s_{i}), \quad i = 1, \dots n$$

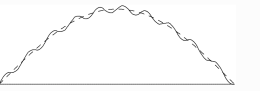
ullet This system of linear algebraic equations Ax=y, where $a_{ij} = w_i K(s_i, t_i), y_i = f(s_i), \text{ and } x_i = u(t_i), \text{ is solved for } \boldsymbol{x}$ to obtain discrete sample of approximate values of u

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Numerical Differentiation

Numerical Differentiation

- Differentiation is inherently sensitive, as small perturbations in data can cause large changes in result
- Differentiation is inverse of integration, which is inherently stable because of its smoothing effect
- For example, two functions shown below have very similar definite integrals but very different derivatives



Double Integrals

Approaches for evaluating double integrals include

- Use automatic one-dimensional quadrature routine for each dimension, one for outer integral and another for inner integral
- Use product quadrature rule resulting from applying one-dimensional rule to successive dimensions
- Use non-product quadrature rule for regions such as triangles

Multiple Integrals, continued

- Monte Carlo method is not competitive for dimensions one or two, but strength of method is that its convergence rate is independent of number of dimensions
- For example, one million points in six dimensions amounts to only ten points per dimension, which is much better than any type of conventional quadrature rule would require for same level of accuracy

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Integral Equations, continued

- Though straightforward to solve formally, many integral equations are extremely sensitive to perturbations in input data, which are often subject to random experimental or measurement errors
- Resulting linear system is highly ill-conditioned
- Techniques for coping with ill-conditioning include
 - Truncated SVD
 - Regularization
 - Constrained optimization

Numerical Differentiation

Numerical Differentiation, continued

- To approximate derivative of function whose values are known only at discrete set of points, good approach is to fit some smooth function to given data and then differentiate approximating function
- If given data are sufficiently smooth, then interpolation may be appropriate, but if data are noisy, then smoothing approximating function, such as least squares spline, is more appropriate

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Finite Difference Approximations

- Given smooth function $f: \mathbb{R} \to \mathbb{R}$, we wish to approximate its first and second derivatives at point x
- Consider Taylor series expansions

$$f(x+h) = f(x) + f'(x)h + \frac{f''(x)}{2}h^2 + \frac{f'''(x)}{6}h^3 + \cdots$$

$$f(x-h) = f(x) - f'(x)h + \frac{f''(x)}{2}h^2 - \frac{f'''(x)}{6}h^3 + \cdots$$

• Solving for f'(x) in first series, obtain *forward difference* approximation

$$f'(x) = \frac{f(x+h) - f(x)}{h} - \frac{f''(x)}{2}h + \dots \approx \frac{f(x+h) - f(x)}{h}$$

which is first-order accurate since dominant term in remainder of series is $\mathcal{O}(h)$

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Numerical Integration
Numerical Differentiation

Numerical Differentiation
Finite Difference Approximations

Finite Difference Approximations, continued

 Adding both series together gives centered difference approximation for second derivative

$$f''(x) = \frac{f(x+h) - 2f(x) + f(x-h)}{h^2} - \frac{f^{(4)}(x)}{12}h^2 + \cdots$$

$$\approx \frac{f(x+h) - 2f(x) + f(x-h)}{h^2}$$

which is also second-order accurate

 Finite difference approximations can also be derived by polynomial interpolation, which is less cumbersome than Taylor series for higher-order accuracy or higher-order derivatives, and is more easily generalized to unequally spaced points

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Richardson Extrapolatio

Richardson Extrapolation

- In many problems, such as numerical integration or differentiation, approximate value for some quantity is computed based on some step size
- Ideally, we would like to obtain limiting value as step size approaches zero, but we cannot take step size arbitrarily small because of excessive cost or rounding error
- Based on values for nonzero step sizes, however, we may be able to estimate value for step size of zero
- One way to do this is called Richardson extrapolation

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Numerical Differentiation Richardson Extrapolation

Richardson Extrapolation, continued

- Suppose we have computed F for two step sizes, say hand h/q for some positive integer q
- Then we have

$$F(h) = a_0 + a_1 h^p + \mathcal{O}(h^r)$$

$$F(h/q) = a_0 + a_1 (h/q)^p + \mathcal{O}(h^r) = a_0 + a_1 q^{-p} h^p + \mathcal{O}(h^r)$$

ullet This system of two linear equations in two unknowns a_0 and a_1 is easily solved to obtain

$$a_0 = F(h) + \frac{F(h) - F(h/q)}{q^{-p} - 1} + \mathcal{O}(h^r)$$

• Accuracy of improved value, a_0 , is $\mathcal{O}(h^r)$

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Numerical Integration
Numerical Differentiation

Numerical Differentiation Finite Difference Approximations

Finite Difference Approximations, continued

 Similarly, from second series derive backward difference approximation

$$f'(x) = \frac{f(x) - f(x - h)}{h} + \frac{f''(x)}{2}h + \cdots$$
$$\approx \frac{f(x) - f(x - h)}{h}$$

which is also first-order accurate

 Subtracting second series from first series gives centered difference approximation

$$f'(x) = \frac{f(x+h) - f(x-h)}{2h} - \frac{f'''(x)}{6}h^2 + \cdots$$

 $\approx \frac{f(x+h) - f(x-h)}{2h}$

which is second-order accurate

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Numerical Differentiation

Automatic Differentiation

Automatic Differentiation

- Computer program expressing function is composed of basic arithmetic operations and elementary functions, each of whose derivatives is easily computed
- Derivatives can be propagated through program by repeated use of chain rule, computing derivative of function step by step along with function itself
- Result is true derivative of original function, subject only to rounding error but suffering no discretization error
- Software packages are available implementing this automatic differentiation (AD) approach

Richardson Extrapolation, continued

Numerical Differentiation Richardson Extrapolation

- Let F(h) denote value obtained with step size h
- If we compute value of F for some nonzero step sizes, and if we know theoretical behavior of F(h) as $h \to 0$, then we can extrapolate from known values to obtain approximate value for F(0)
- Suppose that

$$F(h) = a_0 + a_1 h^p + \mathcal{O}(h^r)$$

as $h \to 0$ for some p and r, with r > p

• Assume we know values of p and r, but not a_0 or a_1 (indeed, $F(0) = a_0$ is what we seek)

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Richardson Extrapolation

Richardson Extrapolation, continued

- Extrapolated value, though improved, is still only approximate, not exact, and its accuracy is still limited by step size and arithmetic precision used
- If F(h) is known for several values of h, then extrapolation process can be repeated to produce still more accurate approximations, up to limitations imposed by finite-precision arithmetic

Numerical Differentiation Richardson Extrapolation

Example: Richardson Extrapolation

- Use Richardson extrapolation to improve accuracy of finite difference approximation to derivative of function sin(x) at x = 1
- Using first-order accurate forward difference approximation, we have

$$F(h) = a_0 + a_1 h + \mathcal{O}(h^2)$$

so p=1 and r=2 in this instance

• Using step sizes of h = 0.5 and h/2 = 0.25 (i.e., q = 2), we obtain

$$F(h) = \frac{\sin(1.5) - \sin(1)}{0.5} = 0.312048$$

$$F(h/2) = \frac{\sin(1.25) - \sin(1)}{0.25} = 0.430055$$

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Numerical Differentiation Richardson Extrapolation Example: Romberg Integration

As another example, evaluate

$$\int_0^{\pi/2} \sin(x) \, dx$$

Using composite trapezoid rule, we have

$$F(h) = a_0 + a_1 h^2 + \mathcal{O}(h^4)$$

so p=2 and r=4 in this instance

- With $h = \pi/2$, $F(h) = F(\pi/2) = 0.785398$
- With q = 2, $F(h/2) = F(\pi/4) = 0.948059$

Romberg Integration

- Continued Richardson extrapolations using composite trapezoid rule with successively halved step sizes is called Romberg integration
- It is capable of producing very high accuracy (up to limit imposed by arithmetic precision) for very smooth integrands
- It is often implemented in automatic (though nonadaptive) fashion, with extrapolations continuing until change in successive values falls below specified error tolerance



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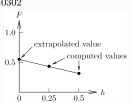
Numerical Differentiation Richardson Extrapolation

Example, continued

Extrapolated value is then given by

$$F(0) = a_0 = F(h) + \frac{F(h) - F(h/2)}{(1/2) - 1} = 2F(h/2) - F(h) = 0.548061$$

• For comparison, correctly rounded result is $\cos(1) = 0.540302$



< interactive example >

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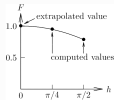
Numerical Differentiation Richardson Extrapolation

Extrapolated value is then given by

Example, continued

$$F(0) = a_0 = F(h) + \frac{F(h) - F(h/2)}{2^{-2} - 1} = \frac{4F(h/2) - F(h)}{3} = 1.002280$$

which is substantially more accurate than values previously computed (exact answer is 1)



< interactive example >

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