5.13 Rational Chebyshev Approximation

In §5.8 and §5.10 we learned how to find good polynomial approximations to a given function $f(x)$ in a given interval $a \leq x \leq b$. Here, we want to generalize the task to find good approximations that are rational functions (see §5.3). The reason for doing so is that, for some functions and some intervals, the optimal rational function approximation is able to achieve substantially higher accuracy than the optimal polynomial approximation with the same number of coefficients. This must be weighed against the fact that finding a rational function approximation is not as straightforward as finding a polynomial approximation, which, as we saw, could be done elegantly via Chebyshev polynomials.

Let the desired rational function $R(x)$ have numerator of degree $m$ and denominator of degree $k$. Then we have

$$R(x) \equiv \frac{p_0 + p_1 x + \cdots + p_m x^m}{1 + q_1 x + \cdots + q_k x^k} \approx f(x) \quad \text{for } a \leq x \leq b \quad (5.13.1)$$

The unknown quantities that we need to find are $p_0, \ldots, p_m$ and $q_1, \ldots, q_k$, that is, $m+k+1$ quantities in all. Let $r(x)$ denote the deviation of $R(x)$ from $f(x)$, and let $r$ denote its maximum absolute value,

$$r(x) \equiv R(x) - f(x) \quad r \equiv \max_{a \leq x \leq b} |r(x)| \quad (5.13.2)$$

The ideal minimax solution would be that choice of $p$'s and $q$'s that minimizes $r$. Obviously there is some minimax solution, since $r$ is bounded below by zero. How can we find it, or a reasonable approximation to it?

A first hint is furnished by the following fundamental theorem: If $R(x)$ is nondegenerate (has no common polynomial factors in numerator and denominator), then there is a unique choice of $p$'s and $q$'s that minimizes $r$; for this choice, $r(x)$ has $m+k+2$ extrema in $a \leq x \leq b$, all of magnitude $r$ and with alternating sign. (We have omitted some technical assumptions in this theorem. See Ralston [1] for a precise statement.) We thus learn that the situation with rational functions is quite analogous to that for minimax polynomials: In §5.8 we saw that the error term of an $n$th order approximation, with $n+1$ Chebychev coefficients, was generally dominated by the first neglected Chebychev term, namely $T_{n+1}$, which itself has $n+2$ extrema of equal magnitude and alternating sign. So, here, the number of rational coefficients, $m+k+1$, plays the same role of the number of polynomial coefficients, $n+1$.

A different way to see why $r(x)$ should have $m+k+2$ extrema is to note that $R(x)$ can be made exactly equal to $f(x)$ at any $m+k+1$ points $x_i$. Multiplying equation (5.13.1) by its denominator gives the equations

$$p_0 + p_1 x_i + \cdots + p_m x_i^m = f(x_i)(1 + q_1 x_i + \cdots + q_k x_i^k) \quad i = 1, 2, \ldots, m+k+1 \quad (5.13.3)$$

This is a set of $m+k+1$ linear equations for the unknown $p$'s and $q$'s, which can be solved by standard methods (e.g., $LU$ decomposition). If we choose the $x_i$'s to all be in the interval $(a, b)$, then there will generically be an extremum between each chosen $x_i$ and $x_{i+1}$, plus also extrema where the function goes out of the interval at $a$ and $b$, for a total of $m+k+2$ extrema. For arbitrary $x_i$'s, the extrema will not have the same magnitude. The theorem says that, for one particular choice of $x_i$'s, the magnitudes can be beaten down to the identical, minimal, value of $r$.

Instead of making $f(x_i)$ and $R(x_i)$ equal at the points $x_i$, one can instead force the residual $r(x_i)$ to any desired values $y_i$ by solving the linear equations

$$p_0 + p_1 x_i + \cdots + p_m x_i^m = [f(x_i) - y_i](1 + q_1 x_i + \cdots + q_k x_i^k) \quad i = 1, 2, \ldots, m+k+1 \quad (5.13.4)$$
In fact, if the \( x_i \)'s are chosen to be the extrema (not the zeros) of the minimax solution, then the equations satisfied will be

\[
p_0 + p_1 x_i + \cdots + p_m x_i^m = [f(x_i) \pm r](1 + q_1 x_i + \cdots + q_r x_i^r)
\]

\[ i = 1, 2, \ldots, m + k + 2 \]

(5.13.5)

where the \( \pm \) alternates for the alternating extrema. Notice that equation (5.13.5) is satisfied at \( m + k + 2 \) extrema, while equation (5.13.4) was satisfied only at \( m + k + 1 \) arbitrary points. How can this be? The answer is that \( r \) in equation (5.13.5) is an additional unknown, so that the number of both equations and unknowns is \( m + k + 2 \). True, the set is mildly nonlinear (in \( r \)), but in general it is still perfectly soluble by methods that we will develop in Chapter 9.

We thus see that, given only the locations of the extrema of the minimax rational function, we can solve for its coefficients and maximum deviation. Additional theorems, leading up to the so-called Remes algorithms [1], tell how to converge to these locations by an iterative process. For example, here is a (slightly simplified) statement of Remes’ Second Algorithm: (1) Find an initial rational function with \( m + k + 2 \) extrema \( x_i \) (not having equal deviation). (2) Solve equation (5.13.5) for new rational coefficients and \( r \). (3) Evaluate the resulting \( R(x) \) to find its actual extrema (which will not be the same as the guessed values). (4) Replace each guessed value with the nearest actual extremum of the same sign. (5) Go back to step 2 and iterate to convergence. Under a broad set of assumptions, this method will converge.Ralston [1] fills in the necessary details, including how to find the initial set of \( x_i \)'s.

Up to this point, our discussion has been textbook-standard. We now reveal ourselves as heretics. We don’t much like the elegant Remes algorithm. Its two nested iterations (on \( r \) in the nonlinear set 5.13.5, and on the new sets of \( x_i \)'s) are finicky and require a lot of special logic for degenerate cases. Even more heretical, we doubt that compulsive searching for the exactly best, equal deviation, approximation is worth the effort — except perhaps for those few people in the world whose business it is to find optimal approximations that get built into compilers and microchips.

When we use rational function approximation, the goal is usually much more pragmatic: Inside some inner loop we are evaluating some function a zillion times, and we want to speed up its evaluation. Almost never do we need this function to the last bit of machine accuracy. Suppose (heresy!) we use an approximation whose error has \( m + k + 2 \) extrema whose deviations differ by a factor of 2. The theorems on which the Remes algorithms are based guarantee that the perfect minimax solution will have extrema somewhere within this factor of 2 range — forcing down the higher extrema will cause the lower ones to rise, until all are equal. So our “sloppy” approximation is in fact within a fraction of a least significant bit of the minimax one.

That is good enough for us, especially when we have available a very robust method for finding the so-called “sloppy” approximation. Such a method is the least-squares solution of overdetermined linear equations by singular value decomposition (§2.6 and §15.4). We proceed as follows: First, solve (in the least-squares sense) equation (5.13.3), not just for \( m + k + 1 \) values of \( x_i \), but for a significantly larger number of \( x_i \)'s, spaced approximately like the zeros of a high-order Chebyshev polynomial. This gives an initial guess for \( R(x) \). Second, tabulate the resulting deviations, find the mean absolute deviation, call it \( r \), and then solve (again in the least-squares sense) equation (5.13.5) with \( r \) fixed and the \( \pm \) chosen to be the sign of the observed deviation at each point \( x_i \). Third, repeat the second step a few times.

You can spot some Remes orthodoxy lurking in our algorithm: The equations we solve are trying to bring the deviations not to zero, but rather to plus-or-minus some consistent value. However, we dispense with keeping track of actual extrema; and we solve only linear equations at each stage. One additional trick is to solve a weighted least-squares problem, where the weights are chosen to beat down the largest deviations fastest.

Here is a program implementing these ideas. Notice that the only calls to the function \( f \) occur in the initial filling of the table \( x \). You could easily modify the code to do this filling outside of the routine. It is not even necessary that your abscissas \( x \) be exactly the ones that we use; though the quality of the fit will deteriorate if you do not have several abscissas between each extremum of the (underlying) minimax solution. Notice that the rational coefficients are output in a format suitable for evaluation by the routine \texttt{ratval} in §5.3.
Chapter 5. Evaluation of Functions

Figure 5.13.1. Solid curves show deviations $r(x)$ for five successive iterations of the routine `ratlsq` for an arbitrary test problem. The algorithm does not converge to exactly the minimax solution (shown as the dotted curve). But, after one iteration, the discrepancy is a small fraction of the last significant bit of accuracy.

```c
#include <stdio.h>
#include <math.h>
#include "nrutil.h"
#define NPFAC 8
#define MAXIT 5
#define PIO2 (3.141592653589793/2.0)
#define BIG 1.0e30

void ratlsq(double (*fn)(double), double a, double b, int mm, int kk, double cof[], double *dev)
{
    double ratval(double x, double cof[], int mm, int kk);
    void dsvbksb(double **u, double w[], double **v, int m, int n, double b[],
        double x[]);
    void dsvdcmp(double **a, int m, int n, double w[], double **v);
    These are double versions of svdcmp, svbksb.
    int i,it,j,ncof,npt;
    double devmax,e,hth,power,sum,*bb,*coff,*ee,*fs,**u,**v,*w,*wt,*xs;
    ncof=mm+kk+1;
    npt=NPFAC*ncof;
    bb=dvector(1,npt);
    coff=dvector(0,ncof-1);
    ee=dvector(1,npt);
    fs=dvector(1,npt);
    u=dmatrix(1,npt,1,ncof);
    v=dmatrix(1,ncof,1,ncof);
    wt=dvector(1,npt);
```

$R(x) - f(x)$

$m = k = 4$

$f(x) = \cos(x)/(1 + e^x)$

$0 < x < \pi$
xs=dvector(1,npt);
*dev=BIG;
for (i=1;i<npt;i++) {
  if (i < npt/2) {
    hth=PIO2*(i-1)/(npt-1.0);
    xs[i]=a+(b-a)*DSQR(sin(hth));
  } else {
    hth=PIO2*(npt-i)/(npt-1.0);
    xs[i]=b-(b-a)*DSQR(sin(hth));
  }
  fs[i]=(*fn)(xs[i]);
  wt[i]=1.0;
  ee[i]=1.0;
}
for (it=1;it<MAXIT;it++) {
  for (i=1;i<npt;i++) {
    if (i < npt/2) {
      hth=PIO2*(i-1)/(npt-1.0);
      xs[i]=a+(b-a)*DSQR(sin(hth));
    } else {
      hth=PIO2*(npt-i)/(npt-1.0);
      xs[i]=b-(b-a)*DSQR(sin(hth));
    }
    fs[i]=(*fn)(xs[i]);
    wt[i]=1.0;
    ee[i]=1.0;
  }
  e=0.0;
  for (it=1;it<MAXIT;it++) {
    bb[i]=power*(fs[i]+SIGN(e,ee[i]));
    Key idea here: Fit to fn(x)+e where the deviation is positive, to fn(x)−e where
    it is negative. Then e is supposed to become an approximation to the equal-ripple
    deviation.
    for (j=1;j<=mm+1;j++) {
      u[i][j]=power;
      power *= xs[i];
    }
    power = -bb[i];
    for (j=mm+2;j<=ncof;j++) {
      u[i][j]=power;
    }
    dsvdcmp(u,npt,ncof,w,v);
    Singular Value Decomposition.
    In especially singular or difficult cases, one might here edit the singular values w[1..ncof],
    replacing small values by zero. Note that dsvbdcmp works with one-based arrays, so we
    must subtract 1 when we pass it the zero-based array coff.
    dsvbksb(u,w,v,npt,ncof,bb,coff-1);
    devmax=sum=0.0;
    for (j=1;j<npt;j++) {
      ee[j]=ratval(xs[j],coff,mm,kk)-fs[j];
      wt[j]=fabs(ee[j]);
      Use weighting to emphasize most deviant points.
      sum += wt[j];
      if (wt[j] > devmax) devmax=wt[j];
    }
    e=sum/npt;
    Update e to be the mean absolute deviation.
    if (devmax <= *dev) {
      Save only the best coefficient set found.
      for (j=0;j<ncof;j++) coff[j]=coff[j];
      *dev=devmax;
    }
    printf(" ratlsq iteration= %2d max error= %10.3e\n",it,devmax);
  }
  free_dvector(xs,1,npt);
  free_dvector(wt,1,npt);
  free_dmatrix(v,1,ncof,1,ncof);
  free_dmatrix(u,1,npt,1,ncof);
  free_dvector(fs,1,npt);
  free_dvector(ee,1,npt);
  free_dvector(coff,0,ncof-1);
  free_dvector(bb,1,npt);
}
Figure 5.13.1 shows the discrepancies for the first five iterations of \texttt{ratlsq} when it is applied to find the \( m = k = 4 \) rational fit to the function \( f(x) = \cos x / (1 + e^x) \) in the interval \((0, \pi)\). One sees that after the first iteration, the results are virtually as good as the minimax solution. The iterations do not converge in the order that the figure suggests: In fact, it is the second iteration that is best (has smallest maximum deviation). The routine \texttt{ratlsq} accordingly returns the best of its iterations, not necessarily the last one; there is no advantage in doing more than five iterations.

CITED REFERENCES AND FURTHER READING:

5.14 Evaluation of Functions by Path Integration

In computer programming, the technique of choice is not necessarily the most efficient, or elegant, or fastest executing one. Instead, it may be the one that is quick to implement, general, and easy to check.

One sometimes needs only a few, or a few thousand, evaluations of a special function, perhaps a complex valued function of a complex variable, that has many different parameters, or asymptotic regimes, or both. Use of the usual tricks (series, continued fractions, rational function approximations, recurrence relations, and so forth) may result in a patchwork program with tests and branches to different formulas. While such a program may be highly efficient in execution, it is often not the shortest way to the answer from a standing start.

A different technique of considerable generality is direct integration of a function’s defining differential equation—an ab initio integration for each desired function value—along a path in the complex plane if necessary. While this may at first seem like swatting a fly with a golden brick, it turns out that when you already have the brick, and the fly is asleep right under it, all you have to do is let it fall!

As a specific example, let us consider the complex hypergeometric function \( {}_2F_1(a, b; c; z) \), which is defined as the analytic continuation of the so-called hypergeometric series,

\[
{}_2F_1(a, b; c; z) = 1 + \frac{ab}{c} z + \frac{a(a+1)b(b+1)}{c(c+1)} \frac{z^2}{2!} + \cdots \\
+ \frac{a(a+1)\cdots(a+j-1)b(b+1)\cdots(b+j-1)}{c(c+1)\cdots(c+j-1)} \frac{z^j}{j!} + \cdots
\]  

The series converges only within the unit circle \(|z| < 1\) (see [1]), but one’s interest in the function is often not confined to this region.

The hypergeometric function \( {}_2F_1 \) is a solution (in fact the solution that is regular at the origin) of the hypergeometric differential equation, which we can write as

\[
z(1-z)F'' = abF' - [c - (a + b + 1)z]F'
\]