Abstract. An object-oriented MATLAB system is described that extends the capabilities of Chebfun to smooth functions of two variables defined on rectangles. Functions are approximated to essentially machine precision by using iterative Gaussian elimination with complete pivoting to form “chebfun2” objects representing low rank approximations. Operations such as integration, differentiation, function evaluation, and transforms are particularly efficient. Global optimization, the singular value decomposition, and rootfinding are also extended to chebfun2 objects. Numerical applications are presented.

Key words. MATLAB, Chebfun, Chebyshev polynomials, low rank approximation

AMS subject classifications. 40-04, 65D05

1. Introduction. In scientific computing it is common that models are continuous even though discretizations are used in their solution. Abstraction, in the object-oriented programming sense of the term, is a powerful idea to bridge the gap between these models and the numerical algorithms that solve them. Chebfun2 is a software system written in MATLAB that exploits abstraction to compute with bivariate functions and is the first extension of Chebfun to two dimensions.

Chebfun was first released in 2004 [1] and has become a well-established software system for computing in one dimension [37]. For smooth functions \( f : [-1, 1] \to \mathbb{R} \), a “chebfun” is a polynomial interpolant of \( f(x) \) through \( n + 1 \) Chebyshev points, given by

\[
x_j = \cos \left( \frac{j\pi}{n} \right), \quad 0 \leq j \leq n.
\]  

(1.1)

The polynomial degree is adaptively chosen so that the chebfun approximates \( f \), on the whole interval, to machine precision. A chebfun object stores these \( n + 1 \) values \( (f(x_j))_{0 \leq j \leq n} \) as a vector, which are the coefficients in the Lagrange basis,

\[
f(x) \approx \sum_{j=0}^{n} f(x_j) \ell_j(x), \quad \ell_j(x) = \frac{\prod_{i=0, i \neq j}^{n} (x - x_i)}{\prod_{i=0, i \neq j}^{n} (x_j - x_i)}.
\]

Chebfun allows numerical computing with functions, and commands such as norm and diff return the 2-norm and derivative, respectively [35]. For convenience, we have summarized a selection of Chebfun commands in Table 1.1. A chebfun is more specifically, a column chebfun when size(f) returns \( \text{inf} \times 1 \), where \( \text{inf} \) represents the continuous variable. Alternatively, if size(f) is \( 1 \times \text{inf} \) then it is a row chebfun. We can also horizontally concatenate column chebfuns together, as in \([f \ g \ h] \), to form a column quasimatrix of size \( \text{inf} \times 3 \), and row quasimatrices can be manipulated similarly. As we shall describe, a column quasimatrix, a row quasimatrix and a diagonal matrix can be combined to approximate functions of two variables and are the main ingredients of a chebfun2.

Developing Chebfun2 has brought up many questions, and our answers have guided this process:

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Table 1.1: A selection of Chebfun commands, their corresponding operations and underlying algorithms. In addition to the references cited, most of these algorithms are discussed in [38].

1. **What should we represent?** We have chosen to represent (a) scalar valued functions and (b) vector valued functions with two components, both on rectangles. The reason for giving (a) and (b) primary status is that they both have innumerable applications, but are mathematically quite distinct. For example, the notion of rootfinding has one natural meaning for scalar functions and quite a different one for vector functions.

2. **What type of approximations should we use?** We represent scalar valued functions by low rank approximants, i.e., sums of functions of the form $u(y)v(x)$, where $u(y)$ and $v(x)$ are univariate functions (which are represented as chebfuns). Low rank approximations allow us to build on 1D Chebyshev technology in a powerful way. A vector valued function is represented by a low rank approximant for each component.

3. **How shall we construct them?** We use an algorithm that is mathematically equivalent to Gaussian elimination (GE) with complete pivoting to construct low rank approximations. This is an unusual application of GE in two respects: (1) We use it as an iterative, rather than a direct, algorithm; (2) In principle, we apply GE to functions rather than matrices. We decide when a low rank approximant achieves machine precision by employing an interesting mix of 1D and 2D resolution tests (see section 2.1).

4. **What are the important operations?** We have taken MATLAB and Chebfun as our guide on this and have overloaded over 130 commands for chebfun2 and chebfun2v objects, which represent scalar and vector functions, respectively. For instance, if $f$ is a chebfun2 on the domain $[-1, 1]$, then we have

$$
\text{trace}(f) = \int_{-1}^{1} f(x, x) \, dx, \quad \text{flipud}(f) = f(x, -y).
$$

The meaning of some operations is obvious, like $\text{diff}$ for computing derivatives in the $x$ or $y$ variable. For others, like $\text{svd}$, a clear choice is suggested by the mathematics, and Chebfun2 operations are designed whenever possible to be the continuous analogues of their MATLAB or Chebfun predecessors.

5. **What reliable and efficient algorithms are there?** Often our algorithms are motivated by algorithms for discrete objects, like matrices, and we hop back and forth between a discrete and continuous mode. The recurring theme is the use of low rank approximants, which consist of sums of products of univariate func-
tions. Every algorithm attempts to exploit this low rank structure and build on well-established 1D technology. A full description of the 1D technology is given in [38].

Chebfun2 is publicly available under an open source license accompanied with examples and all the MATLAB code [32]. [Note to referees: this will be true by 4th of March 2013.]

Throughout this article we restrict our attention to scalar and vector valued functions defined on the default unit square, i.e. \([-1,1]^2\), though the software permits easy treatment of general rectangular domains.

In the next section we describe how GE with complete pivoting can be used as a practical algorithm for approximating a function of two variables. In section 3 we describe the Chebfun2 algorithms for integration, differentiation, function evaluation and transforms, which are based on 1D technology. Section 4 presents the vector form of Chebfun2 for operations to implement “div, grad, curl and all that” for vector valued functions. In section 5 we introduce an algorithm for global optimization and, as an example, use it solve a problem from the SIAM 100-Digit Challenge [6, 34]. In section 6 we describe the singular value decomposition of a chebfun2 and numerically demonstrate that GE with complete pivoting can compute near-optimal low rank approximations. In section 7 we discuss the current rootfinding capabilities of Chebfun2. Finally, in section 8 we describe two prospects for further work.

We have announced the release of Chebfun2 earlier in an article in SIAM News [33].

2. Low rank function approximation. Given a continuous bivariate function \(f : [-1,1]^2 \to \mathbb{R}\), the optimal rank \(k\) approximation in the \(L^2\)-norm is given by the singular value decomposition (SVD). The SVD (or Karhunen–Loève expansion) is defined by [29]

\[
f(x, y) = \sum_{j=1}^{\infty} \sigma_j \phi_j(y) \psi_j(x),
\]

(2.1)

where \(\sigma_1, \sigma_2, \ldots,\) is a non-increasing real sequence of singular values, and the sets \(\{\phi_1(x), \phi_2(x), \ldots,\}\) and \(\{\psi_1(y), \psi_2(y), \ldots,\}\) are orthonormal functions in \(L^2([-1,1])\).

Each term in (2.1) is an “outer product” of two univariate functions, called a rank 1 function. The optimal rank \(k\) approximation to \(f\) in the \(L^2\)-norm can be found by truncating (2.1) after \(k\) terms,

\[
f(x, y) \approx f_k(x, y) = \sum_{j=1}^{k} \sigma_j \phi_j(y) \psi_j(x), \quad \|f - f_k\|_{L^2([-1,1]^2)} = \left( \sum_{j=k+1}^{\infty} \sigma_j^2 \right)^{\frac{1}{2}}.
\]

Fortunately, the sequence \(\sigma_1, \sigma_2, \ldots,\) decays rapidly for smooth functions. The convergence rates for the singular values have a similar flavor to univariate approximation theory: the smoother the function, the faster the singular values decay. Some results to this effect can be found in [17, 23, 41], and we hope to discuss the convergence question further in a separate publication.

Numerically, a rank \(k\) approximant to \(f\) can be computed by sampling it on a \(n \times n\) Chebyshev tensor grid, taking the matrix of sampled values, and computing its matrix singular value decomposition. The first \(k\) singular values and left and right singular vectors form the optimal rank \(k\) approximation to the sampled matrix in
Algorithm: GE with complete pivoting for functions

Input: A function \( f = f(x, y) \) on \([-1, 1]^2\) and a tolerance \( \text{tol} \)

Output: A low rank approximation \( f_k(x, y) \) satisfying \(|f - f_k| < \text{tol}\)

\[ e_0(x, y) = f(x, y), \quad f_0(x, y) = 0 \]

for \( k = 0, 1, 2, \ldots, \)

\[ |e_k(x_k, y_k)| = \max(|e_k(x, y)|), \quad (x, y) \in [-1, 1]^2 \]

if \( |e_k(x_k, y_k)| < \text{tol}, \text{ stop} \)

\[ e_{k+1}(x, y) = e_k(x, y) - e_k(x_k, y_k)e_k(x_k, y_k)/e_k(x_k, y_k) \]

\[ f_{k+1}(x, y) = f_k(x, y) + e_k(x_k, y_k)e_k(x_k, y_k)/e_k(x_k, y_k) \]

end

Fig. 2.1: Iterative GE with complete pivoting for approximation of functions of two variables. The first \( k \) steps construct a rank \( k \) approximation to \( f \). Our numerical algorithm is based on a discretization of this continuous idealization.

the discrete 2-norm. Functions can be constructed from each left (right) singular vector by polynomial interpolation, and provided \( n \) is sufficiently large, these will be good approximations to the first \( k \) singular functions. This process constructs an approximate singular value decomposition of a function and requires \( O(n^3) \) operations and \( n^2 \) evaluations of \( f \). Instead, we use GE with complete pivoting to compute a near-optimal rank \( k \) approximation in \( O(k^2n + k^3) \) operations.

Here is our algorithm of GE with complete pivoting applied to a bivariate function \( f \). First, we define \( e_0 = f \) and find an approximation to the location of \( \max |e_0(x, y)| \) over \([-1, 1]^2\), say \((x_0, y_0)\). Then we construct a rank 1 function

\[ f_1(x, y) = \frac{e_0(x_0, y_0)e_0(x, y)}{e_0(x_0, y_0)} = d_1c_1(y)r_1(x), \quad d_1 = \frac{1}{e_0(x_0, y_0)}, \]

which interpolates \( f \) along the two lines \( y = y_0 \) and \( x = x_0 \). We calculate the residual \( e_1 = f - f_1 \) and repeat the same procedure to form a rank 2 function

\[ f_2(x, y) = f_1(x, y) + \frac{e_1(x_1, y_1)e_1(x, y_1)}{e_1(x_1, y_1)} = f_1(x, y) + d_2c_2(y)r_2(x), \]

where \((x_1, y_1)\) is an approximate location to the maximum of \( |e_1(x, y)| \). The function \( f_2 \) interpolates \( f \) along \( x = x_0, x = x_1, y = y_0, \) and \( y = y_1 \). We continue constructing successive approximations \( f_1, f_2, \ldots, f_k \), where \( f_k \) interpolates \( f \) along \( 2k \) lines, until an approximate maximum of \( |e_k| = |f - f_k| \) falls below relative machine precision. Figure 2.1 gives the pseudocode for this algorithm. Note that each rank 1 function is the product of two univariate functions, which are represented very efficiently by Chebfun.

The number of steps for convergence of this algorithm is function dependent, and in practice, we usually observe that the approximate maxima \( d_1, \ldots, d_k \) decay at the same rate as the singular values. Surprisingly often, only a few steps of GE are required to approximate a function to machine precision.
We call \((x_0, y_0), \ldots, (x_{k-1}, y_{k-1})\) the pivot locations and \(d_1, \ldots, d_k\) the pivot values. We also refer to the functions \(c_1(y), \ldots, c_k(y)\) and \(r_1(x), \ldots, r_k(x)\) as the column slices and row slices, respectively. After constructing \(k\) successive approximations \(f_1, \ldots, f_k\), we say we have performed \(k\) steps of GE.

2.1. Algorithmic details. So far we have described GE at the continuous level, in terms of the manipulation of functions. However, to achieve machine precision with the speed of numerical linear algebra we actually work with a discretized version of the algorithm of Figure 2.1. The discretized algorithm splits into two stages, with the first designed to find candidate pivot locations, i.e., good approximate locations of the absolute maxima, and the second to ensure we have sufficiently sampled the column and row slices.

**Stage 1: Finding candidate pivot locations and rank \(k\).** First we sample \(f\) on a \(9 \times 9\) Chebyshev tensor grid and perform at most 3 steps of GE. If we find the sampled matrix can be approximated to machine precision by a rank 1, 2 or 3 matrix, then we move on to stage 2; otherwise, we sample on a \(17 \times 17\) Chebyshev tensor grid and perform at most 5 steps of GE. We proceed to stage 2 if a matrix of rank 5, or less, is sufficient. We continue sampling on nested Chebyshev grids of size 9, 17, 33, 65, and so on until we discover the sampled matrix can be approximated to machine precision by a matrix of rank 3, 5, 9, 17, and so on.

Thus, stage 1 approximates a \(2^{j+2} + 1\) \times \(2^{j+2} + 1\) matrix by a matrix of rank \(2^j + 1\), or less, for \(j \geq 1\). Generically, if \(f : [-1, 1]^2 \rightarrow \mathbb{R}\) can be approximated to essentially machine precision by a rank \(k\) function, then stage 1 samples \(f\) on a \((2^{j+2} + 1) \times (2^{j+2} + 1)\) tensor grid, where \(j = \lceil \log_2(k - 1) \rceil\), and \(k\) steps of GE are required. Since

\[
2^{\lceil \log_2(k-1) \rceil + 2} + 1 \leq 8k + 1 = O(k),
\]

stage 1 requires \(O(k^3)\) operations. We store the pivot locations used in the \(k\) successful steps of GE and go to stage 2.

**Stage 2: Resolving column and row slices.** Stage 1 has determined a candidate set of pivot locations required to approximate \(f\), and stage 2 is designed to ensure that the column and row slices are sufficiently sampled. For instance, \(f(x, y) = x \cos(100y)\) is a rank 1 function, so stage 1 completes after sampling on a \(9 \times 9\) Chebyshev tensor grid even though 147 Chebyshev samples are needed to resolve the oscillations in the \(y\)-direction. For efficiency we only sample \(f\) on a \(k\)-skeleton of a tensor grid, i.e., a subset consisting of \(k\) columns and rows of the grid, and perform GE on that skeleton. For example, Figure 2.2 shows the 4-skeleton used when approximating Franke’s function [13],

\[
f(x, y) = \frac{3}{4} e^{-((9x-2)^2 + (9y-2)^2)/4} + \frac{3}{4} e^{-((9x-1)^2 / 49 - (9y+1)/10)} + \frac{1}{2} e^{-((9x-7)^2 + (9y-3)^2)/4} - \frac{1}{5} e^{-((9x-4)^2 + (9y-7)^2)}.
\]  

(2.2)

After \(k\) steps of GE we have sampled the column and row slices at Chebyshev points. Following the procedure used by Chebfun, we convert each column and row slice to coefficients using the fast Fourier transform to ensure that their tails have decayed to essentially machine precision. Figure 2.2 shows the Chebyshev coefficients for the column slices used to approximate (2.2). For instance, if the column slices are not resolved with 33 points, then we sample \(f\) at 65 points along each column and repeat
Fig. 2.2: Left: The skeleton used in stage 2 of the construction algorithm for approximating Franke’s function (2.2). Stage 2 only samples \( f \) on the skeleton, i.e., along the black lines. Right: The Chebyshev coefficients of the four column slices. The coefficients decay to machine precision, indicating that the column slices have been sufficiently sampled.

\( k \) steps of GE. We continue increasing the sampling along columns and rows until we have resolved them. Since the sets of 9, 17, 33, \ldots, Chebyshev points are nested, we can always pivot at the same locations as determined in stage 1. If the column slices require degree \( m - 1 \) polynomial interpolants and the row slices require degree \( n - 1 \) polynomial interpolants, then this stage samples \( f \) at, at most,

\[
k \left( 2^{\left\lceil \log_2(n) \right\rceil} + 2^{\left\lceil \log_2(m) \right\rceil} \right) \leq 2k(n + m)
\]

points. We then perform \( k \) steps of GE on the selected rows and columns, requiring \( \mathcal{O} \left( k^2(n + m) \right) \) operations.

At the end of the construction we have approximated a function \( f \) as

\[
f(x, y) \approx f_k(x, y) = \sum_{j=1}^{k} d_j c_j(y) r_j(x).
\]

We store the column slices in a column quasimatrix \( C(y) = [c_1(y), \ldots, c_k(y)] \), where \( c_j(y) \) is a column chebfun. Similarly, the row slices are stored as chebfuns in a row quasimatrix \( R(x) \), and the pivot values \( d_1, \ldots, d_k \) are stored as a vector. We can also write the approximant as

\[
f_k(x, y) = C(y)DR(x), \quad D = \text{diag} \left( d_1, \ldots, d_k \right).
\]

This representation highlights how GE decouples the \( x \) and \( y \) variables, which will be very useful for tensor product operations (see section 3).

Figure 2.3 shows contour plots with pivot locations of six chebfun2 objects that approximate bivariate functions to machine precision. For example, the following Chebfun2 code produces the top-left plot:

```matlab
f = chebfun2(@(x,y) 1./(1+100*(x.^2+y.^2).^2))
contour(f), hold on, plot(f,'.k')
```
Fig. 2.3: Contour plots for six functions $f$ on $[-1,1]^2$, with the pivot locations marked by dots: (a) $1/(1 + 100(x^2 + y^2)^2)$; (b) $1/(1 + 100(\frac{x}{2} - x^2 - y^2)^2)$; (c) $1/(1 + 1000((x - \frac{1}{2})^2(y + \frac{1}{2})(x + \frac{1}{2})(y - \frac{1}{2}))$; (d) $\cos(10(x^2 + y)) + \sin(10(x + y^2))$; (e) $\exp(-5(x^2 + y^2))\exp(-5(x^2 + y^2))$; (f) $\tanh(10x)\tanh(10y)/\tanh(10y^2) + \cos(5x)$.

**Remark.** We deliberately do not represent each column and row slice as an independent chebfun with the minimal degree. Instead we ensure all the column slices have the same degree. Similarly, all the row slices have the same degree (which can be different than the degree of the column slices). This is important for efficiency since subsequent operations can then be vectorized.

### 2.2. Object composition

One or more chebfun2 objects can be combined together to make new ones; the object-oriented programming term for this is *object composition*. Table 2.1 summarizes some composition operations that are possible with chebfun2 objects. For example, if $f$ is a chebfun2, then $\cos(f)$, $\exp(f)$ and $f.^2$ return chebfun2 objects corresponding to the cosine, exponential and square of $f$, respectively. To carry out these operations the Chebfun2 constructor is called, so that the chebfun2 for $\exp(f)$, for example, is constructed by applying the iterative GE algorithm in the usual manner to samples of $\exp(f)$ on a sequence of grids.

### 2.3. Related approaches

Ideas related to Gaussian elimination for functions have been developed by various authors under various names, though the connection with GE is usually not mentioned. We now briefly summarize some of the ideas of *pseudoskeleton approximation* [14], *Adaptive Cross Approximation* (ACA) [3], *Geddes–Newton series* [8], and *rank-revealing decompositions* [20, 27].
Table 2.1: A selection of object composition commands in Chebfun2. In each case
the result is computed by calling the Chebfun2 constructor to generate a low rank
approximation to machine precision.

<table>
<thead>
<tr>
<th>Chebfun2 command</th>
<th>Operation</th>
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<td>+, -</td>
<td>addition, subtraction</td>
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<tr>
<td>.*, ./</td>
<td>multiplication, division</td>
</tr>
<tr>
<td>cos, sin, tan</td>
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<tr>
<td>cosh, sinh, tanh</td>
<td>hyperbolic functions</td>
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<td>exp</td>
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<tr>
<td>power</td>
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</table>

**Pseudoskeletons and Adaptive Cross Approximation.** Pseudoskeletons were developed by Goreinov, Tyrtshnikov, and Zamarashkin [14] and approximate
a matrix $A \in \mathbb{R}^{n \times n}$ by a matrix of low rank by computing the CUR decomposition\(^1\) $A \approx CUR$, where $C \in \mathbb{R}^{n \times k}$ and $R \in \mathbb{R}^{k \times n}$ are subsets of the columns and rows of $A$, respectively, and $U \in \mathbb{R}^{k \times k}$. Selecting good columns and rows of $A$ is of paramount importance, and this can be achieved via maximizing volumes [15], randomized techniques [12], or ACA. ACA was mainly developed by Bebendorf and constructs a skeleton approximation with columns and rows selected adaptively [2]. The selection of a column and row corresponds to choosing a pivot location in GE, where the pivoting entry is the element belonging to both the column and row. If the first $k$ columns and rows are selected, then

$$
\begin{pmatrix}
A_{11} & A_{12} \\
A_{21} & A_{22}
\end{pmatrix} - \begin{pmatrix}
A_{11} \\
A_{21}
\end{pmatrix} A_{11}^{-1} \begin{pmatrix}
A_{11} & A_{12}
\end{pmatrix} = \begin{pmatrix}
0 & 0 \\
0 & S
\end{pmatrix}, \quad A_{11} \in \mathbb{R}^{k \times k}, \quad (2.4)
$$

where $S = A_{22} - A_{21} A_{11}^{-1} A_{12}$ is the Schur complement of $A_{22}$ in $A$. The relation (2.4) is also found in [3, p. 128] and can be compared with Theorem 1.4 of [30] to show that ACA and GE are equivalent. This connection remains even when any $k$ columns and rows are selected [30, Theorem 1.8]. The use of the continuous analogue of GE with complete pivoting by Chebfun2 is equivalent to the continuous analogue of ACA with column and row selection via complete pivoting.

In practice, ACA is used to compute low rank approximation of matrices that are derived by sampling functions [3], and it has been used extensively by Hackbusch and others for constructing hierarchical representations of matrices [18, 19].

**Geddes–Newton series.** Independently of ACA, Chapman in his PhD thesis [10] in 2003 developed a theoretical framework for what he calls the *Geddes–Newton series*. For a function $f$ and a splitting point $(a, b) \in [-1, 1]$ such that $f(a, b) \neq 0$, the splitting operator $\Upsilon_{(a, b)}$ is defined as

$$
\Upsilon_{(a, b)} f(x, y) = \frac{f(x, b) f(a, y)}{f(a, b)},
$$

---

\(^1\)The pseudoskeleton literature writes $A \approx CGR$. More recently, it has become popular to follow the nomenclature of [12] and replace $G$ by $U$. 
and this is the rank 1 approximation to $f$ obtained after one step of GE with pivot location $(a, b)$. The splitting operator is now applied to the function

$$f(x, y) - \Upsilon_{(a,b)} f(x, y)$$

with a different splitting point. Repeating this process $k$ times is equivalent to applying $k$ steps of GE on $f$.

Carvajal, Chapman and Geddes used GE for the quadrature of symmetric functions [8], and it was their work that initially led us to develop the algorithm for Chebfun2. Their integration algorithm first maps a function to $[0, 1]^2$ and then decomposes it into symmetric and anti-symmetric parts, ignoring the anti-symmetric part since it integrates to zero. However, Carvajal et al. employed a specialized pivoting strategy designed to preserve symmetry, whereas we use complete pivoting. Geddes–Newton series stand out for being the only case of an iterative GE type of algorithm that we know of in the literature introduced mainly to approximate functions rather than matrices [10].

**Rank-revealing decomposition.** The SVD computes the optimal rank $k$ approximation to a matrix in the 2-norm, though it is relatively expensive. The QR, UTV$^T$ and LU decompositions can also be used for constructing low rank matrix approximations that can be near-optimal [21, 27, 31]. The LU decomposition computed via GE with complete pivoting can be an excellent rank-revealing decomposition, and hence, the rank of a chebfun2 approximant to a function is usually close to the minimal rank required to achieve machine precision.

3. **Quadrature and other tensor product operations.** If $f$ is a matrix, then the MATLAB command \texttt{sum(f)}, or \texttt{sum(f,1)}, returns a row vector representing the sum of each column, and \texttt{sum(f,2)} returns a column vector after summing up the rows. If $f$ is a chebfun2, then we have

$$\texttt{sum(f,1)} = \int_{-1}^{1} f(x,y)dy, \quad \texttt{sum(f,2)} = \int_{-1}^{1} f(x,y)dx,$$

represented as row and column chebfuns, respectively. Thus in the language of probability, the sum command computes marginal distributions. The algorithm underlying these operations exemplifies how Chebfun2 takes advantage of its low rank representations, together with the established 1D Chebfun technology, to carry out substantial computations much faster than one might expect.

For example, consider the computation of \texttt{sum(f,2)}, whose result is a chebfun in the $y$ variable. If $f$ is of rank $k$, the necessary formula is

$$\int_{-1}^{1} f(x,y)dx = \int_{-1}^{1} \sum_{j=1}^{k} d_j c_j(y) r_j(x) dx = \sum_{j=1}^{k} d_j c_j(y) \left( \int_{-1}^{1} r_j(x)dx \right). \quad (3.1)$$

It follows that the evaluation of \texttt{sum(f,2)} reduces to the integration of $k$ chebfuns, in parentheses in (3.1), and the addition of the corresponding multiples of $c_j(y)$ in Chebfun. The integration is done by calling Chebfun’s \texttt{sum} command, which utilizes the Clenshaw–Curtis quadrature rule

$$\int_{-1}^{1} r_j(x)dx = \sum_{i=0}^{n} w_i r_j(x_i), \quad 1 \leq j \leq k,$$
where $x_i$ are the Chebyshev nodes (1.1) and $w_i$ are the quadrature weights.

Following familiar MATLAB syntax, the command `sum(sum(f))` delivers the definite integral over $[-1, 1]^2$ by integrating over the $y$-variable, constructing an intermediary row chebfun, and then integrating this. The command `sum2(f)` computes the same quantity more efficiently because an intermediary chebfun is not constructed. The necessary result corresponds to the integration of $2k$ chebfuns in which all the column (row) slices are of the same degree. Thus, the Clenshaw–Curtis quadrature can be vectorized, and only two sets of quadrature weights are computed using Chebfun’s algorithm based on [39].

Thus, integration is very efficient for a chebfun2, because it is carried out via one-dimensional quadrature rules. A similar surprising efficiency extends to other tensor product operators, which represent a significant part of Chebfun2 functionality.

**Definition 3.1.** A tensor product operator $L$ is a linear operator on functions of $x$ and $y$ with the property that if $f(x, y) = c(y)r(x)$, then $L = L_y(c) L_x(r)$, for some operators $L_y$ and $L_x$. Thus, if $f$ has rank $k$, then

$$L \left( \sum_{j=1}^{k} d_j c_j(y)r_j(x) \right) = \sum_{j=1}^{k} d_j (L_y c_j(y))(L_x r_j(x)).$$

A tensor product operation can be computed with $O(k)$ calls to well-established Chebfun algorithms since $L_y$ and $L_x$ act on functions of one variable. Four important examples of tensor product operators are integration (described above), differentiation, evaluation, and the fast Fourier transform.

**Differentiation.** If $f$ is an $n \times n$ matrix, then the MATLAB command `diff(f,N)`, or `diff(f,N,1)`, returns an $(n - N) \times n$ matrix of $N$th order differences along the columns, while `diff(f,N,2)` calculates the differences along the rows. For a chebfun2 the same syntax represents differentiation:

$$\text{diff}(f,N,1) = \frac{\partial^N f}{\partial y^N}, \quad \text{diff}(f,N,2) = \frac{\partial^N f}{\partial x^N}.$$  

Differentiation is the tensor product of univariate differentiation since

$$\frac{\partial^N f}{\partial y^N} = \sum_{j=1}^{k} d_j \frac{\partial^N c_j(y)}{\partial y^N} r_j(x), \quad \frac{\partial^N f}{\partial x^N} = \sum_{j=1}^{k} d_j c_j(y) \frac{\partial^N r_j(x)}{\partial y^N}.$$  

These functions are computed with $k$ calls to the Chebfun `diff` command.

These differentiation and integration algorithms mean that vector calculus on chebfun2 and chebfun2v objects is very efficient (see section 4).

**Function evaluation.** In MATLAB, indexing into a matrix selects a subset of its elements. In Chebfun2, the analogous operation is evaluation of $f$ at various points, with $f(x,y)$ returning the value of $f$ at $(x,y)$. Evaluation is a tensor product operation since

$$f(x,y) = \sum_{j=1}^{k} d_j c_j(y)r_j(x),$$

which involves the evaluation of $2k$ univariate functions, and this computation is carried out by using the barycentric formula [5, 28]. This formula has been vectorized so that multiple column and row slices of the same degree are evaluated efficiently.
**Computation of Chebyshev coefficients.** If \( f \) is a chebfun, \( \text{chebpoly}(f) \) computes the Chebyshev expansion coefficients of a Chebfun, i.e., the coefficients in the expansion

\[
f(x) = \sum_{j=0}^{n} \alpha_j T_j(x),
\]

in \( O(n \log n) \) operations using the fast Fourier transform. This is a very important operation and the main reason why Chebfun uses a Chebyshev basis instead of, for example, a basis of Legendre polynomials.

Analogously, the Chebfun2 command \( X=\text{chebpoly2}(f) \) computes the expansion coefficients of a chebfun2, i.e., the coefficients in the expansion

\[
f(x, y) = \sum_{i=0}^{m-1} \sum_{j=0}^{n-1} \alpha_{ij} T_i(y) T_j(x),
\]

and the \( m \times n \) matrix returned represents these coefficients. If \( f \) is of rank \( k \), then the coefficient matrix is also of rank \( k \), and the command \( [A \; D \; B]=\text{chebpoly2}(f) \) requires only \( O(k(n \log n + m \log m)) \) operations since it returns the coefficient matrix in its low rank form, i.e., \( X = ADB^T \), where \( A, B \in \mathbb{R}^{n \times k} \) and \( D \in \mathbb{R}^{k \times k} \) is diagonal. This operation is computed with \( 2k \) calls to the Chebfun command \( \text{chebpoly} \) since

\[
\text{chebpoly2}(f) = \sum_{j=1}^{k} d_j \text{chebpoly}(c_j)^T \text{chebpoly}(r_j),
\]

where each \( c_j \) is evaluated at \( m \) Chebyshev points and each \( r_j \) at \( n \) Chebyshev points using the fast Fourier transform. For evaluation at points that are a subset of a Chebyshev grid, this approach can be significantly faster than use of the barycentric formula.

Tensor product operators represent the ideal situation where well-established Chebfun technology can be exploited for computing in two dimensions. Table 3.1 shows a selection of Chebfun2 commands that are tensor product operations. If a prescribed function happens to be univariate, then \( k = 1 \), and these Chebfun2 commands are essentially the same as their univariate counterparts.

**4. Chebfun2v objects and vector calculus.** As mentioned in the Introduction, Chebfun2 is also designed to work with vector valued functions defined on rectangles, as well as scalar valued ones. Our convention is to use a lower case letter for a scalar function, \( f \), and an upper case letter for a vector function, \( F = (f_1, f_2)^T \).

From the point of view of approximation, the vector aspect of Chebfun2 introduces no new complications. A vector function \( F \) is represented as two independent scalar functions, and such a representation is constructed by computing a low rank approximation to each component.

The more interesting side of vector Chebfun2 is the set of operations that can be implemented — algorithmically similar to scalar functions, but potentially very useful for applications.

\[\text{chebpoly2}(f) = \sum_{j=1}^{k} d_j \text{chebpoly}(c_j)^T \text{chebpoly}(r_j),\]

where each \( c_j \) is evaluated at \( m \) Chebyshev points and each \( r_j \) at \( n \) Chebyshev points using the fast Fourier transform. For evaluation at points that are a subset of a Chebyshev grid, this approach can be significantly faster than use of the barycentric formula.

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The Chebfun command \( \text{chebpoly} \) always returns a row vector of coefficients as of version 4.2.2194, though this may change in a future release.
### Table 3.1
A selection of scalar Chebfun2 commands corresponding to tensor product operations. In each case the result is computed with much greater speed than one might expect because the algorithms can take advantage of the rank \( k \) structure.

<table>
<thead>
<tr>
<th>Chebfun2 command</th>
<th>Operation</th>
</tr>
</thead>
<tbody>
<tr>
<td>sum, sum2</td>
<td>integration</td>
</tr>
<tr>
<td>cumsum</td>
<td>cumulative integration</td>
</tr>
<tr>
<td>prod</td>
<td>product integration</td>
</tr>
<tr>
<td>cumprod</td>
<td>cumulative product integration</td>
</tr>
<tr>
<td>norm</td>
<td>( L^2 )-norm</td>
</tr>
<tr>
<td>diff</td>
<td>differentiation</td>
</tr>
<tr>
<td>chebpoly2</td>
<td>fast Fourier transform</td>
</tr>
<tr>
<td>( f(x,y) )</td>
<td>evaluation</td>
</tr>
<tr>
<td>plot, surf, contour</td>
<td>plotting</td>
</tr>
<tr>
<td>diag</td>
<td>univariate function along a diagonal</td>
</tr>
<tr>
<td>trace</td>
<td>integration along ( y = x )</td>
</tr>
<tr>
<td>flipud, fliplr</td>
<td>reverse direction of coordinates</td>
</tr>
<tr>
<td>mean2, std2</td>
<td>mean, standard deviation</td>
</tr>
</tbody>
</table>

### Algebraic operations.
The basic non-differential operations of vector calculus are scalar multiplication \( fG \), vector addition \( F + G \), dot product \( F \cdot G \), and cross product \( F \times G \). Unfortunately, there are two notational inconveniences in MATLAB: (1) The inability to use the ‘.’ symbol for the dot product, and (2) The lack of a ‘x’ symbol for the cross product. The ‘.’ symbol is already used as a special character for referencing, and ‘x’ is usually used as a variable name, preventing it from being an operator. Accordingly, in Chebfun2 we have chosen \( f .* G \), \( F + G \), \( \text{dot}(F,G) \), and \( \text{cross}(F,G) \) for the four operations. We explain these operations in turn.

Scalar multiplication is the product of a scalar function with a vector function, denoted by \( f .* G \), and algorithmically, it is achieved by two scalar function multiplications, one for each component.

Vector addition, denoted by \( F + G \), yields another chebfun2v and is computed by adding the two scalar components together. It satisfies the parallelogram law

\[
2\|F\|_2^2 + 2\|G\|_2^2 = \|F + G\|_2^2 + \|F - G\|_2^2,
\]

which can be verified numerically, as in this example:

\[
F = \text{chebfun2v}(@(x,y)\cos(x.*y),@(x,y)\sin(x.*y));
G = \text{chebfun2v}(@(x,y)x+y, @(x,y)1+x+y);
abs((2*norm(F)^2 + 2*norm(G)^2) - (norm(F+G)^2 + norm(F-G)^2))
\]

\[\text{ans} = 3.5527e-15\]

The dot product, denoted by \( \text{dot}(F,G) \), takes two vector functions and returns a scalar function. Algebraically, it is the inner product, i.e., the sum of the products of the two components. If the dot product of two chebfun2v objects takes the value zero at \((x_0, y_0)\), then the vectors \( F(x_0, y_0) \) and \( G(x_0, y_0) \) are orthogonal.

The cross product, denoted by \( \text{cross}(F,G) \), is well known as an operation on vector valued functions with three components, and it can also be defined for 2D vector fields by

\[
\text{cross}(F,G) = f_1g_2 - f_2g_1, \quad F = \begin{pmatrix} f_1 \\ f_2 \end{pmatrix}, \quad G = \begin{pmatrix} g_1 \\ g_2 \end{pmatrix}.
\]
Note that the cross product of two vector valued functions with two components is a scalar function, which can be represented by a chebfun2.

**Differential operations.** Vector calculus also involves various differential operators defined on scalar or vector valued functions such as gradient $\nabla f$, curl $\nabla \times F$, divergence $\nabla \cdot F$, and Laplacian $\nabla^2 f$.

The gradient of a chebfun2 is represented by a chebfun2v such that

$$\text{grad}(f) = \left( \frac{\partial f}{\partial x}, \frac{\partial f}{\partial y} \right),$$

which points in the direction of steepest ascent of $f$. The gradient theorem says that the integral of $\text{grad}(f)$ over a curve only depends on the values of $f$ at the endpoints of that curve. We can check this numerically by using the Chebfun2v command `integral`. This command computes the line integral of a vector valued function

$$\text{integral}(F,C) = \int_C F(r) \cdot dr,$$

where $C$ is a smooth curve represented by a complex valued chebfun (see Note below) and $\cdot$ is the dot product. For example,

$$f = \text{chebfun2}(\theta(x,y)\sin(2x)+x.*y.^2); \quad \% \text{chebfun2 object}$$

$$F = \text{grad}(f); \quad \% \text{gradient (chebfun2v)}$$

$$C = \text{chebfun}(@(t) t.*\exp(100i*t),[0 \ \pi/10]); \quad \% \text{spiral curve}$$

$$v = \text{integral}(F,C); \quad \text{ends} = f(pi/10,0)-f(0,0); \quad \% \text{line integral}$$

$$\text{abs}(v-\text{ends}) \quad \% \text{gradient theorem}$$

$$\text{ans} = 5.5511e-16$$

The curl of 2D vector function is a scalar function defined by

$$\text{curl}(F) = \frac{\partial f_2}{\partial x} - \frac{\partial f_1}{\partial y}, \quad F = \left( \begin{array}{c} f_1 \\ f_2 \end{array} \right).$$

If the chebfun2v $F$ describes a vector velocity field of fluid flow, for example, then $\text{curl}(F)$ is a scalar function equal to twice the angular speed of a particle in the flow at each point. A particle moving in a gradient field has zero angular speed and hence, $\text{curl}(\text{grad}(f)) = 0$, a well known identity that can also be checked numerically.

The divergence of a chebfun2v is defined as

$$\text{div}(F) = \frac{\partial f_1}{\partial x} + \frac{\partial f_2}{\partial y}, \quad F = \left( \begin{array}{c} f_1 \\ f_2 \end{array} \right).$$

This measures a vector field’s distribution of sources or sinks. The Laplacian is closely related and is the divergence of the gradient $\text{lap}(f) = \text{div}(\text{grad}(f))$.

Table 4.1 summarizes the vector calculus commands available in Chebfun2.

**Phase portraits.** A phase portrait is a graphical representation of a system of trajectories for a two-variable autonomous dynamical system. In Chebfun2 we can plot phase portraits by using the `quiver` command, which has been overloaded to plot the vector field (see Figure 4.1).

In addition, Chebfun2 makes it easy to compute and plot individual trajectories of a vector field. If $F$ is a chebfun2v, then `ode45(F,tspan,y0)` solves the autonomous system $dx/dt = f_1(x,y)$, $dy/dt = f_2(x,y)$, where $f_1$ and $f_2$ are the first and second components of $F$, respectively, with the prescribed time interval and initial conditions. This command returns a complex valued chebfun representing the trajectory in the form $x(t) + iy(t)$. 

13
<table>
<thead>
<tr>
<th>Command</th>
<th>Operation</th>
</tr>
</thead>
<tbody>
<tr>
<td>+, -</td>
<td>addition, subtraction</td>
</tr>
<tr>
<td>dot</td>
<td>dot product</td>
</tr>
<tr>
<td>cross</td>
<td>cross product</td>
</tr>
<tr>
<td>grad</td>
<td>gradient</td>
</tr>
<tr>
<td>curl</td>
<td>curl</td>
</tr>
<tr>
<td>div</td>
<td>divergence</td>
</tr>
<tr>
<td>lap</td>
<td>Laplacian</td>
</tr>
<tr>
<td>quiver</td>
<td>phase portrait</td>
</tr>
</tbody>
</table>

Table 4.1: Vector calculus commands in Chebfun2. In each case the result is computed with greater speed than one might expect because of the exploitation of the tensor product structure (see section 3).

**Note.** As is well known, a pair of real scalar functions \( f \) and \( g \) can be encoded as a complex function \( f + ig \). In the design of Chebfun2 we have debated how much use to make of this trick, which can simplify many operations, but at the same time may be confusing. For example, every (real) chebfun2v might have been realized as a complex chebfun2. We decided not to go that far, and the complex output of \texttt{ode45} and \texttt{roots} (see section 7), as well as the second input argument to the command \texttt{integral} (above) are exceptions. The complex output of \texttt{ode45} allows the trajectory’s length to be computed with the Chebfun command \texttt{arclength}.

The following code produces the phase portrait for a dynamical system corresponding to a nonlinear pendulum, \( \dot{x} = y, \dot{y} = -\sin(x)/4 \), together with some sample trajectories:

```matlab
phasedom=[-4 4 -2 2]; % phase plane domain
F = chebfun2v(@(x,y)y,@(x,y)-sin(x)/4,phasedom);
for ic = .5:.5:3 % initial conditions
    [t y]=ode45(F,[0 40],[ic,0]); % solve autonomous system
    plot(y,'r'), hold on, % plot trajectory
end
quiver(F) % vector field
arclength(y) % length of a trajectory
ans = 14.0386
```

Figure 4.1 shows the resulting image, as well as a corresponding picture for the Duffing oscillator

\[
\dot{x} = y, \quad \dot{y} = \frac{4}{100}y - \frac{3}{4}x + x^3,
\]

where the time interval is \([0, 50]\) and the initial condition is \(x(0) = 0, y(0) = \frac{1}{2}\).

**5. Global Optimization.** If \( f \) is a matrix, then the MATLAB command \texttt{max(f)}, or \texttt{max(f,[],1)}, returns a row vector of maximum values along each column of \( f \). Similarly, \texttt{max(f,[],2)} returns a column vector of maximum values along each row. If \( f \) is a chebfun2, then the same syntax yields analogous results,

\[
\max(f,[],1) = \max_{y \in [-1,1]} f(x,y), \quad \max(f,[],2) = \max_{x \in [-1,1]} f(x,y)
\]
Fig. 4.1: Phase plane diagrams and example trajectories for the nonlinear pendulum (left) and the Duffing oscillator (right). Red lines are trajectories in the phase plane computed using the Chebfun2 command `ode45`.

represented as row and column chebfuns, respectively. Often these chebfuns are only piecewise smooth, and Chebfun’s edge detection algorithm is used to decide the locations of the breakpoints [26]. If \( f \) and \( g \) are chebfun2 objects, then we do not currently allow \( \max(f, g) \), because this 2D function will generally have discontinuities in its first derivative lying along curves, and Chebfun2 cannot represent such nonsmooth functions.

The global 2D maximum of a chebfun2 can be computed by \( \max(\max(f)) \), but the same result can be obtained more efficiently by avoiding an intermediary non-smooth chebfun with \( \max2(f) \). If a chebfun2 is of rank 1, i.e., \( f(x, y) = d_1 c_1(y) r_1(x) \), then the global maximum is computed by noting that

\[
\max2(f) = d_1 \max \{ \max c_1 \cdot \max r_1, \min c_1 \cdot \max r_1, \max c_1 \cdot \min r_1, \min c_1 \cdot \min r_1 \},
\]

where each \( \max \) and \( \min \) involves only univariate functions and existing Chebfun algorithms are used [38, Chap. 18]. For \( k \geq 2 \), however, Chebfun2 uses other algorithms for global optimization, and Table 5.1 summarizes how the command and rank of a chebfun2 determine the algorithm employed. All the algorithms, for \( k \geq 2 \), are based on obtaining initial guesses by one method and then switching to a superlinearly convergent constrained trust-region method\(^3\), based on [9], in which the iterates are constrained to the domain of the chebfun2.

5.1. Convex hull algorithm. For low rank chebfun2 objects, currently those with rank \( 2 \leq k \leq 3 \), if the global maximum or minimum is desired, then we employ the following algorithm based on the convex hull. Mathematically, this algorithm is not restricted to \( k = 2, 3 \), but it is inefficient if \( k > 3 \). For any fixed point \( (x, y) \in \)

\(^3\) [Note to referees: The current implementation employs the MATLAB command `fmincon`, which is only available in the Optimization Toolbox. This will be replaced by our own publicly available implementation in early 2013.]
\[ k = 1 \quad 2 \leq k \leq 3 \quad k \geq 4 \]

<table>
<thead>
<tr>
<th></th>
<th>( k \rightarrow )</th>
<th>( 2 \leq k \leq 3 )</th>
<th>( k \geq 4 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>min2, max2 or minandmax2</td>
<td>Chebfun</td>
<td>convex hull algorithm</td>
<td>grid evaluation</td>
</tr>
<tr>
<td>norm(( \cdot, \infty ))</td>
<td>Chebfun</td>
<td>convex hull algorithm</td>
<td>power iteration</td>
</tr>
</tbody>
</table>

Table 5.1: Optimization algorithms in Chebfun2. Various algorithms are used in different situations and the choice depends on the rank \( k \) of the chebfun2 and on the particular Chebfun2 command. For \( k \geq 2 \), the algorithms listed are used to provide initial guesses for a trust region iteration.

In \([-1, 1]^2\) we have

\[
f(x, y) = \sum_{j=1}^{k} d_j c_j(y) r_j(x) = \left( \sqrt{|d_1|} c_1(y), \ldots, \sqrt{|d_k|} c_k(y) \right) \left( \pm \sqrt{|d_1|} r_1(x) \\
\vdots \\
\pm \sqrt{|d_k|} r_k(x) \right) = s(y)^T t(x),
\]

where \( s, t : [-1, 1] \to \mathbb{R}^k \) are continuous functions and the signs are chosen appropriately depending on \( d_j, 1 \leq j \leq k \). We can define the following parameterizable curves in \( \mathbb{R}^k \),

\[
S = \{ s(y) \in \mathbb{R}^k : y \in [-1, 1] \}, \quad T = \{ t(x) \in \mathbb{R}^k : x \in [-1, 1] \},
\]

and approximate them by discrete sets \( \tilde{S} \in \mathbb{R}^{m \times k}, \tilde{T} \in \mathbb{R}^{n \times k} \) by evaluating \( s(y) \) and \( t(x) \) at \( m \) and \( n \) Chebyshev points, respectively. As always, \( m \) and \( n \) are the numbers of Chebyshev coefficients required to represent the column and row slices (see section 2.1). We then make the following approximation to the global maximum:

\[
\max \{ f(x, y) : x, y \in [-1, 1] \} = \max \{ s(y)^T t(x) : x, y \in [-1, 1] \} \\
= \max \{ a^T b : a \in S, b \in T \} \\
\approx \max \{ a^T b : a \in \tilde{S}, b \in \tilde{T} \} \\
= \max \{ a^T b : a \in \text{conv}(\tilde{S}), b \in \text{conv}(\tilde{T}) \},
\]

where \( \text{conv}(X) \) is the convex hull of \( X \). The convex hulls of \( \tilde{S} \) and \( \tilde{T} \) are computed using the MATLAB command \texttt{convhull}. We discard any point \( (x, y) \) if either \( t(x) \notin \text{conv}(\tilde{T}) \) or \( s(y) \notin \text{conv}(\tilde{S}) \).

Figure 5.1 shows an example of the convex hull algorithm. Here we take the rank 2 function \( f(x, y) = 2y \cos(5x^2) + x \sin(2y^2) \) and show the convex hull of \( \tilde{S} \) and \( \tilde{T} \) and the corresponding excluded gray region in \([-1, 1]^2\).
The convex hull algorithm allows us to obtain candidate regions in $[-1, 1]^2$ containing the global extrema, and in these regions we sample the chebfun2 on the Chebyshev tensor grid and take the discrete maximum and minimum as initial guesses in a trust-region iteration.

5.2. Grid evaluation. For the commands $\text{max2}$, $\text{min2}$, and $\text{minandmax2}$, if $k > 3$, then we evaluate the chebfun2 on a $m \times n$ Chebyshev grid using the fast Fourier transform, where $m$ and $n$ are the numbers of Chebyshev coefficients required to represent the column and row slices. From this grid we take the discrete maximum and minimum as initial guesses for the trust-region iteration.

5.3. Power iteration. If the global absolute maximum of $|f(x, y)|$ is required with $k > 3$, such as for $\text{norm}(f, \text{inf})$, then we use a fast power iteration as proposed in [4]. The advantage of this method is that it can be much faster when $k \ll \min(m, n)$.

For the power iteration we first sample the column and row slices of a chebfun2 at $m$ and $n$ Chebyshev points, respectively, and obtain a matrix $A \in \mathbb{R}^{m \times n}$ of rank $k$, though we do not form this matrix explicitly. Given a rank $k$ matrix $A \in \mathbb{R}^{m \times n}$,

$$A = \sum_{j=0}^{k} d_j c_j r_j^T, \quad c_j \in \mathbb{R}^{m \times 1}, \quad r_j \in \mathbb{R}^{n \times 1}, \quad d_j \in \mathbb{R},$$

the diagonal matrix $B \in \mathbb{R}^{mn \times mn}$ defined by

$$B = \sum_{j=1}^{k} d_j \text{diag}(c_j) \otimes \text{diag}(r_j)$$

has eigenvalues $A_{ij}$ with eigenvectors $e_i \otimes e_j$, where $e_i$ is the $i$th canonical vector. Hence, the maximum entry of $|A|$ can be computed by performing the power iteration on $B$. The low rank structure of $A$ is exploited so that the $mn$ diagonal entries of $B$ are not formed explicitly. Moreover, the initial vector $x \in \mathbb{R}^{mn \times 1}$ is stored in Kronecker product form and kept in this form throughout the iteration by compression [4]. This power iteration requires $O(k^2(m + n))$ operations per iteration.
Example: Global minimum of a complicated function. In February 2002, an article in SIAM News by the second author set a challenge to solve ten problems each to ten digits of precision (the solution of each problem was a real number) [34]. Figure 5.2 shows one of the problems, which involves finding the global minimum of a function. Since the term \((x^2 + y^2)/4\) grows away from \((0, 0)\) while the other terms remain bounded, it can be shown that the global minimum occurs in \([-1, 1]^2\) [6].

The function is complicated and oscillatory, but of rank 4, as can be seen by rearranging its terms and using the identity \(\sin(a + b) = \sin(a) \cos(b) + \cos(a) \sin(b)\),

\[
f(x, y) = \left(\frac{x^2}{4} + e^{\sin(50x)} + \sin(70\sin(x))\right) + \left(\frac{y^2}{4} + \sin(60e^y) + \sin(80\sin(y))\right)
- \cos(10x) \sin(10y) - \sin(10x) \cos(10y).
\]

The Chebfun2 algorithm as implemented in the command min2 finds the global minimum in 0.18 seconds\(^4\) and achieves 12 correct digits.

\[
f = @(x,y) \exp(\sin(50*x)) + \sin(60*exp(y)) + \sin(70*\sin(x)) + \ldots
- \sin(\sin(80*y)) - \sin(10*(x+y)) + (x.^2+y.^2)/4;
g = chebfun2(f); Y = min2(g); % Global minimum
abs(Y - (-3.306868647475237)) % Compare with exact answer
ans = 4.4098e-13
\]

A surprising number of interesting functions of two variables are of low rank, and many that are not mathematically of low rank still have excellent low rank approximations.

6. Singular value decomposition. If \(A\) is a \(n \times n\) matrix, the MATLAB command \([U, S, V]=\text{svd}(A, 0)\) returns the “economy-sized” SVD, where \(U, V \in \mathbb{R}^{n \times k}\) and \(S \in \mathbb{R}^{k \times k}\) is a diagonal matrix. If \(f\) is a chebfun2, then \([U, S, V]=\text{svd}(f)\), or \(\text{svd}(f, 0)\), returns the SVD of \(f\), where \(U, V\) are column quasimatrices with \(k\) orthonormal columns (in the \(L^2\) sense) and \(S\) is a \(k \times k\) diagonal matrix. Of course this is not the exact SVD, which in general would have infinite rank, but a finite rank approximation accurate to machine precision.

If each column of \(U\) and \(V\) is a chebfun of degree at most \(n\), then computing the SVD of a rank \(k\) chebfun2 requires \(O(k^2n)\) operations. Figure 6.1 shows the pseudocode used for the computation. The first step requires the Householder triangularization of column quasimatrices and uses the Chebfun command \(\text{qr}\) described in [36]. An excellent way to construct the SVD of a smooth bivariate function is usually by first constructing a chebfun2 and then orthogonalizing the column and row slices.

---

\(^4\)Timing was done on a 2012 1.6GHz Intel Core i5 MacBook Air with MATLAB 2012a.
Pseudocode: SVD of a chebfun2

**Input:** A chebfun2 \( f(x,y) = C(y)DR(x) \) (see 2.3)

**Output:** Quasimatrices with orthonormal columns \( U_f \) and \( V_f \), and a diagonal matrix \( S_f \)

1. Householder triangularization: \( C(y) = QC(y)RC \), \( R(x)^T = QR(x)R \)
2. Compute: \( A = R_CDR_C^T \)
3. Matrix SVD: \( A = U_A S_A V_A^T \)
4. Compute: \( U_f = QC(y)U_A, V_f = QR(x)V_A \) and \( S_f = S_A \)

Fig. 6.1: Pseudocode for computing the SVD of a chebfun2. This algorithm uses Householder triangularization of a quasimatrix [36].

**Example: Gaussian bumps.** As an example, we compute the singular values of a function composed by adding together 300 Gaussian bumps at arbitrary locations,

\[
f(x,y) = \sum_{j=0}^{300} e^{-\gamma((x-s_j)^2+(y-t_j)^2)}, \quad (s_j, t_j) \in [-1,1]^2, \quad (6.1)
\]

where \( \gamma = 10, 100, 1000 \). A Gaussian bump is a rank 1 function, and hence, mathematically, \( f \) is of rank 300 (almost surely). However, it can be approximated to machine precision by a function of much lower rank. Figure 6.2 displays a contour plot for the case \( \gamma = 1000 \) and shows that the singular values of \( f \) decay supergeometrically. The figure is computed with the following code:

```matlab
s = RandStream('mt19937ar','Seed',1); % for reproducibility
gamma = 1000; f = chebfun2();
for n = 1:300
    x0 = 2*rand(s)-1; y0 = 2*rand(s)-1;
    df = chebfun2(@(x,y)exp(-((x-x0).^2+(y-y0).^2)/gamma));
    f = f + df;
end
subplot(1,2,1), contour(f)
subplot(1,2,2), semilogy(svd(f))
```

**Example: Near-optimality.** The singular value decomposition allows us to explore the near-optimality of GE with complete pivoting for constructing low rank approximations. In Figure 6.3 we take the 2D Runge function given by

\[
f(x,y) = \frac{1}{1 + \gamma (x^2 + y^2)^3}, \quad \gamma = 1, 10, 100,
\]

which is analytic, and Wendland’s compactly supported radial basis functions [40],

\[
\phi_{3,k}(|x-y|) = \phi_{3,k}(r) = \begin{cases}
(1-r) & k = 3 \\
(1-r)^4 & k = 1 \\
(1-r)^2 & k = 0,
\end{cases}
\]

\[
n(x,y) = \frac{1}{1 + \gamma (x^2 + y^2)^3}, \quad \gamma = 1, 10, 100,
\]

which is analytic, and Wendland’s compactly supported radial basis functions [40],
Fig. 6.2: Left: Three hundred arbitrarily centered Gaussian bump functions added together, as in (6.1), with $\gamma = 1000$. Right: Supergeometric decay of the normalized singular values (the first singular value is scaled to be 1) for $\gamma = 10, 100, 1000$. Mathematically, (6.1) is almost surely of rank 300, but it can be approximated to machine precision in these realizations by functions of rank 21, 59 and 176, respectively. The supergeometric decay of the singular values can be explained by theory related to the fast Gauss transform [16].

Fig. 6.3: A comparison of SVD and GE approximations shows the near-optimality of the latter approximations to smooth bivariate functions. Left: 2D Runge functions. Right: Wendland’s compactly supported radial basis functions.

which have $2k$ continuous derivatives and thus belong to $C^{2k}$. Figure 6.3 shows the $L^2$ errors of the optimal rank approximations computed via the expensive singular value decomposition, and the near-optimal rank approximations constructed via the Chebfun2 constructor (GE with complete pivoting). We observe that GE with complete pivoting usually computes near-optimal low rank approximations.

7. Rootfinding. There can be infinitely many complex roots of a scalar valued function of two variables and, generically, they form continuous curves that do not end abruptly$^5$ [25]. A goal appropriate for many applications is to find representations of

$^5$A formal discussion is beyond the scope of this paper.
Fig. 7.1: The critical points of \( f(x, y) = (x^2 - y^3 + 1/8) \sin(10 xy) \) (black dots) lie at the intersections of the zero contours of \( f_x \) (blue curves), and \( f_y \) (red curves). If \( f \) is a chebfun2, then \( \text{roots}(f) \) returns complex valued chebfuns interpolating the zero contours of \( f \), and \( \text{roots}(\text{gradient}(f)) \) returns the critical points of \( f \).

these zero curves. Currently, in Chebfun2 we use the MATLAB command \texttt{contourc} to find points lying on the zero contours of \( f \), and then use a complex valued chebfun to interpolate these points. Accordingly, if \( f \) is a chebfun2, \( \text{roots}(f) \) returns a quasi-matrix, where each column is a complex valued chebfun interpolating a zero contour of \( f \) in \([-1,1]^2\).

Determining the roots of a vector valued bivariate function with two components is an entirely different problem since, generically, the roots are now finite in number and isolated [22]. If \( F \) is a chebfun2v, then \( \text{roots}(F) \) aims to return the isolated roots of the system

\[
F(x, y) = \begin{pmatrix} f_1(x, y) \\ f_2(x, y) \end{pmatrix} = 0, \quad (x, y) \in [-1,1]^2. \tag{7.1}
\]

To achieve this, \( \text{roots}(F) \) approximates the zero contours of \( f_1 \) and \( f_2 \) using \texttt{contourc} and then finds their intersections, which are used as initial guesses in a Newton iteration. For example, Figure 7.1 plots the critical points and the zero contours of \( f_x \) and \( f_y \) for \( f(x, y) = (x^2 - y^3 + 1/8) \sin(10 xy) \) by using the following code:

```matlab
f = chebfun2(@(x,y) (x.^2-y.^3+1/8).*sin(10*x.*y));
r = roots(gradient(f)); % critical points
plot(roots(diff(f,1,2)),’b’), hold on % plot zero contours of f_x
plot(roots(diff(f)),’r’) % plot zero contours of f_y
plot(r(:,1),r(:,2),’k.’,’MarkerSize’,30) % plot extrema
```

Rootfinding by computing zero contours and their intersection works remarkably well in many examples. However, this algorithm is not completely robust and certainly does not offer the same reliably as univariate rootfinding based on the eigenvalues of the colleague matrix [38, Chap. 18]. Occasionally, rootfinding via the intersections of zero contours fails to find all the solutions to (7.1), and we are investigating alternative algorithms to take better advantage of the low rank structure of a chebfun2.
8. Future work. We are entering into an exciting stage of development where computing with scalar and vector valued multivariate functions to machine precision is practical. Chebfun2 has shown that computations can be performed in MATLAB in 2D by building on Chebfun technology. Challenges ahead include the extension of Chebfun2 to functions with singularities, more general domains, and higher dimensions. Mathematically, a key issue is to analyze the approximation and convergence properties of the low rank approximations that form the basis of Chebfun2.

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