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# DECAY BOUNDS AND LINEAR SCALING ALGORITHMS FOR APPROXIMATING FUNCTIONS OF BAND MATRICES\*

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*Dedicated to Gene Golub on the occasion of his 75th birthday*

**Abstract.** We establish decay bounds for the entries of  $f(A)$  where  $A$  is a banded (more generally, sparse)  $n \times n$  diagonalizable matrix and  $f$  is smooth on a subset of the complex plane containing the spectrum of  $A$ . Combined with techniques from approximation theory, the bounds are used to compute banded approximations to  $f(A)$ , resulting in algorithms that under appropriate conditions have linear complexity in the matrix size  $n$ . Applications to various types of problems are discussed and illustrated by numerical examples.

**Key words.** Band matrices, matrix functions, decay rates, linear scaling algorithms, Chebyshev polynomials, Faber polynomials, density matrix, trace, determinant

**AMS subject classifications.** Primary 65F10, 65F30. Secondary 15A.

**1. Introduction.** This paper is devoted to the problem of computing approximations to matrix functions  $f(A)$  where  $A$  is a large, banded or sparse matrix of order  $n$  and  $f$  is smooth on a region containing the eigenvalues of  $A$ . We are motivated by the current need for fast algorithms for approximating a wide class of matrix functions that occur in many areas of computational science and engineering, ranging from quantum chemistry to nanoscience to the numerical solution of evolutionary PDEs by time-implicit methods. Our approach is based on the study and careful exploitation of certain decay properties exhibited by the entries of  $f(A)$ . We show that in many important cases the entries of  $f(A)$  decay very quickly in magnitude outside some band or sparsity pattern. In other words, the entries of  $f(A)$  tend to be strongly localized, an observation that opens the door to the possibility of approximating relevant matrix functions in  $O(n)$  time. Localization phenomena of this kind are well-known to physicists and chemists, especially those working on electronic structure calculations. With few exceptions, numerical analysts have so far largely neglected this property and its impact on the design of fast approximation algorithms. There is also a need for numerical analysts to analyze existing  $O(n)$  algorithms developed by physicists and to develop new, more efficient ones. This paper intends to be a first step towards the realization of such program.

The paper is organized as follows. In section 2 we provide some background on matrix functions and make a few observations on related work. In section 3, after providing some background on approximation theory, we study localization phenomena in matrix functions. In particular, we prove very general decay bounds for the entries of matrix functions. Our results generalize and unify several previous results known in the literature. This theory can be used, in principle, to determine the bandwidth or sparsity pattern outside of which the entries of  $f(A)$  are guaranteed to be so small that they can be neglected without exceeding a prescribed error tolerance. In section 4 we discuss approximation methods based on classical approximation theory,

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including Chebyshev and Faber polynomials, and related approaches. Together with truncation strategies for neglecting small elements, these techniques form the basis for the approximation algorithms we use. Section 5 discusses selected applications, while preliminary numerical experiments illustrating the promising behavior of the approximation schemes are presented in section 6. Finally, in section 7 we give our conclusions and discuss future work.

**2. Functions of matrices.** Unless otherwise specified, all matrices considered in this paper are square and complex. Given a matrix  $A$  and a function  $f$  defined on a subset of the complex plane containing the spectrum of  $A$ , it is possible to give several (equivalent) definitions of the matrix function  $f(A)$ ; see, e.g., [41, 51, 58, 61, 86]. For instance, when  $f$  is analytic,  $f(A)$  can be defined by a power series expansion, or by a contour integral (using Cauchy's Theorem). Other definitions of  $f(A)$  can be given in terms of canonical forms, like the Jordan canonical form or the Schur triangular form. Another useful definition is given by  $f(A) = r(A)$  where  $r$  denotes the Lagrange–Hermite interpolation polynomial that interpolates  $f$  (and, in case of multiple eigenvalues, certain of its derivatives) at the eigenvalues of  $A$ . This last definition shows that *every matrix function of  $A$  is a polynomial in  $A$* ; the coefficients of the polynomial, of course depending on  $A$  (and not just on  $f$ ). For the purpose of the theoretical part of this paper, the most useful definitions are those in terms of spectral decompositions but we also make occasional use of other definitions. We briefly review these important notions next.

Suppose  $A$  is diagonalizable:  $A = XDX^{-1}$  where  $D$  is a diagonal matrix with the eigenvalues  $\lambda_1, \lambda_2, \dots, \lambda_n$  on the main diagonal and  $X = [x_1 \ x_2 \ \dots \ x_n]$  is a nonsingular matrix whose columns  $x_i$  are eigenvectors of  $A$  associated with the corresponding eigenvalues  $\lambda_i$ . Then one defines  $f(A) = Xf(D)X^{-1}$  where  $f(D)$  is the diagonal matrix with diagonal entries given by  $f(\lambda_1), f(\lambda_2), \dots, f(\lambda_n)$ . It can be shown that this definition does not depend on the particular choice of the eigenvector matrix  $X$ . If  $A$  is normal, then  $X$  is unitary:  $X^{-1} = X^H$ , so that

$$(2.1) \quad f(A) = Xf(D)X^H = \sum_{i=1}^n f(\lambda_i)x_i x_i^H, \quad \text{where } x_i^H x_j = \delta_{ij}.$$

While it is possible to extend this definition to non-diagonalizable matrices using the Jordan canonical form, in this paper we mainly consider diagonalizable matrices.

The definition in terms of contour integrals is as follows: If  $f$  is analytic on a region  $\Omega \subseteq \mathbb{C}$  containing the eigenvalues of  $A$  and if  $\gamma$  is a closed Jordan curve (oriented counterclockwise) entirely contained within  $\Omega$  such that all the eigenvalues of  $A$  lie in its interior, one can define

$$(2.2) \quad f(A) = \frac{1}{2\pi i} \int_{\gamma} f(z)(zI - A)^{-1} dz.$$

This (classical) definition is popular because it can also be used to define analytic functions of bounded linear operators on infinite-dimensional Banach spaces.

The definition in terms of power series expansion is also classical. Let  $z_0 \in \mathbb{C}$  be the center of a disk of radius  $\rho > 0$  containing all the eigenvalues of  $A$  in its interior, and let  $f$  be analytic on this disk. Then  $f(A)$  can be expressed as a power series:

$$(2.3) \quad f(A) = \sum_{n=0}^{\infty} a_n (A - z_0 I)^n.$$

Other series expansions are also possible, including approximations in terms of orthogonal polynomials and rational functions.

We are concerned with the fundamental problem of computing approximations to matrix functions  $f(A)$  where  $A$  is a large, banded (more generally, sparse)  $n \times n$  matrix. This problem is of basic importance in various areas of science and engineering. Although the problem of computing matrix functions has received considerable attention in the literature on numerical linear algebra and differential equations, we intend to address problems and issues that so far have been largely neglected in the numerical analysis literature and yet are of considerable importance for applications and also in their own right. To explain how our approach differs from the large body of existing work, we point out that much of the literature on the approximation of matrix functions falls into two broad classes:

1. Algorithms designed for the “exact” (i.e., high-accuracy) computation of  $f(A)$  where  $A$  is a dense matrix of moderate size; here  $f$  is often some elementary function such as the exponential, the sine (or cosine), the  $p$ -th root, or the sign function. See, e.g., [17, 34, 51, 54, 58, 68, 79].
2. Algorithms designed for the computation of the vector  $f(A)v$  where  $v$  is a given vector and  $A$  is a large, sparse matrix. This problem is especially important in the numerical integration of large systems of time-dependent differential equations, parabolic PDEs, etc.; see, e.g., [14, 15, 18, 23, 29, 30, 31, 36, 37, 38, 45, 59, 60, 66, 72, 87]. Most of these papers are concerned with the computation of the action of the exponential operator  $\exp(-tA)$  on a vector  $v$ , where  $t > 0$  and the eigenvalues of  $A$  have positive real part.

In contrast, we focus on the problem of finding approximations (to within a prescribed tolerance and for an appropriate choice of the error metric, usually dictated by the underlying application) to the actual entries of  $f(A)$  when  $A$  is large and banded or sparse. The ultimate goal is to develop reliable algorithms with guaranteed accuracy and optimal complexity. For a sparse or banded  $n \times n$  matrix  $A$ , a lower bound for the best possible complexity is  $O(n)$  arithmetic work and storage: i.e., the cost of the approximation, in terms of time and storage, scales at least linearly in the size of the problem. Generally speaking, attainment of this lower bound rules out the computation of  $f(A)$  by reduction to diagonal or triangular (Schur) form. Also, the straightforward use of algorithms for computing  $f(A)v$  applied to the case where  $v = e_i$  (the  $i$ th column of the identity matrix, with  $i = 1 : n$ ) is likely to result in  $O(n^2)$  complexity, although careful implementations of inexact (“sparse-sparse”) variants of this approach may work in some cases.

One approach that has been used in the literature to approximate selected entries of matrix functions is the method based on Gaussian quadrature rules developed by Golub and collaborators over the years; see, e.g., [50]. The idea behind this approach is to express the bilinear form  $h(u, v) = u^T f(A)v$  as a Riemann-Stieltjes integral with respect to a suitable measure (the spectral measure), and to approximate this integral by Gauss-type quadrature rules. These in turn can be computed by means of the Lanczos process, using only matrix–vector products with  $A$ . The choice  $u = e_i$  and  $v = e_j$  yields approximations to the  $(i, j)$  entry of  $f(A)$ . See [4, 5, 11] for further details and some applications. While the original method was developed for real symmetric matrices, nonsymmetric extensions have been since developed; see, e.g., [22] for an approach based on the Arnoldi process. This methodology is quite different from that used in the present paper. We plan to perform a comparison in the near future.

Clearly,  $O(n)$  algorithms are feasible only if  $f(A)$  can be well-approximated, in some representation, by a banded or sparse matrix, with an error that is independent of  $n$  (see section 3). It is also worth noting that in many applications not all the entries of  $f(A)$  are needed, but only selected ones. For instance, it may happen that only an estimate of the trace (sum of the diagonal elements) of  $f(A)$  is required. Another scalar function that is sometimes necessary to estimate is the determinant  $\det[f(A)]$ , and in some cases it is possible to approximate this quantity without computing all the entries of  $f(A)$ . Also, in many applications very high accuracy is not required or warranted.

The key idea of the paper is to exploit the fact that for large classes of matrices  $A$  and functions  $f$ , the “important” entries of  $f(A)$  are highly localized; for instance, the entries of  $f(A)$  exhibit exponentially fast off-diagonal decay. We emphasize that the vast majority of the algorithms currently in vogue among numerical analysts for the computation of  $f(A)$  do not exploit this property. Among the few exceptions, see [4, 6, 31, 62, 85]; however, these papers all address some very special problems, and a systematic treatment of sparsity in matrix functions is lacking. We also mention that chemists and physicists who work in the important field of electronic structure calculations have observed the exponential decay property in the density matrix (a function  $P = f(H)$  of the Hamiltonian  $H$ ) for non-metallic systems, see [3, 48, 57, 67, 76]. (For metallic systems, the decay is algebraic rather than exponential, see [47].) Physicists have exploited this property by developing “sparsified” iterative algorithms for the approximation of density matrices. Partial results and a few algorithms tailored to specific matrix functions can be found scattered in the literature, but no general theory or rigorous foundations for many of the algorithms developed by physicists exist. There is, in other words, little available theory for such approximation algorithms; moreover, error estimates and safe stopping criteria have yet to be developed, and optimality results are also missing. Finally, the existing  $O(n)$  algorithms are applicable to the Hermitian case only, and there is a strong need for methods that can handle non-Hermitian matrices as well. In this paper we begin to address some of these questions.

**3. Decay bounds.** Many physical phenomena are characterized by strong localization, that is, rapid decay outside of a small spatial or temporal region. Frequently, such localization can be expressed as decay in the entries of a function  $f(A)$  of an appropriate sparse or banded matrix  $A$  that encodes a description of the system under study. In this section we survey the current state of knowledge on this topic and contribute some new results that extend and unify previously known ones. We begin with a simple graph-theoretical result on the sparsity pattern of  $f(A)$ , according to which such matrix is generally full. Then we consider banded approximations to matrices that exhibit exponential off-diagonal decay of the entries, showing that approximations that scale linearly in the size of the problem are feasible. Next, we provide some needed background on approximation theory and finally we prove new exponential decay bounds for a wide class of functions  $f$  and matrices  $A$ .

**3.1. A structural result.** Let  $A = [a_{ij}]$  be an  $n \times n$  matrix, not necessarily diagonalizable. We associate to this matrix a digraph  $G(A) = (V, E)$  in the standard way (we refer to [35] for basic background on graph theory). The vertex set  $V$  consists of the integers from 1 to  $n$  and the edge set  $E$  contains all ordered pairs  $(i, j)$  with  $a_{ij} \neq 0$ . For what follows we will assume that the diagonal elements of  $A$  are nonzero and therefore the digraph  $G(A)$  contains loops. Proposition 3.1 states a simple structural result about  $f(A)$ . Recall that the *index* of an eigenvalue is the size of the largest

Jordan block associated with that eigenvalue.

**PROPOSITION 3.1.** *Let  $A$  be an irreducible  $n \times n$  matrix with distinct eigenvalues  $\lambda_1, \dots, \lambda_s$  and let  $n_i$  denote the index of  $\lambda_i$ . Let  $f$  be a function defined on a set containing the eigenvalues of  $A$  that is  $m - 1$  times differentiable where  $m$  is the maximum value of all  $n_i$ 's. In addition, assume that  $f^{(j)}(\lambda_i) \neq 0$  for  $i = 1, \dots, s$  and  $j = 0, \dots, n_i - 1$ . Then the matrix  $f(A)$  is structurally full.*

*Proof.* Let  $r(A)$  be the unique Lagrange–Hermite interpolating polynomial satisfying  $r^{(j)}(\lambda_i) = f^{(j)}(\lambda_i)$  for  $j = 0, \dots, n_i - 1$  and  $i = 1, \dots, s$ , then  $f(A) = r(A)$ . It is easy to check that under our assumptions,  $r(A)$  is of degree  $n - 1$ . We may assume that the matrix  $A$  is a 0-1 matrix since we are only interested in its structure. The digraph  $G(A)$  associated with  $A$  is strongly connected since  $A$  is irreducible. For a positive integer  $l$ , the  $(i, j)$  entry of  $A^l$  is the number of walks of length  $l$  in  $G(A)$  from vertex  $i$  to vertex  $j$ . Let  $k$  denote the length of the longest path in  $G(A)$ ,  $k \leq n - 1$ . Every walk of length less than  $k$  can be extended to a walk of length  $k$  since  $A$  has no zero diagonal elements. Finally, because  $G(A)$  is strongly connected all entries of  $A^k$  must be non. Hence, the matrix  $r(A) = f(A)$  is structurally full.  $\square$

The above result means that barring fortuitous cancellation, all entries of  $f(A)$  are nonzero when  $A$  is irreducible and  $f$  is a “generic” function, by which we mean a function with “sufficiently many” non-vanishing derivatives. We note that this result generalizes a well-known fact about the inverse of an irreducible matrix. Thus, it would seem that the quest for  $O(n)$  algorithms for approximating  $f(A)$  is doomed from the start. However, as we will see, it is often the case that most of the entries in  $f(A)$  are very small in magnitude, thus making banded or sparse approximations possible.

**3.2. Banded approximations to highly localized matrices.** Here we consider approximating a dense matrix with certain decay properties by means of a banded (or sparse) matrix. The problem is important for several reasons. First, there are situations where the matrix  $A$  itself is completely dense, but has rapidly decaying entries; matrices of this kind are sometimes called *pseudospars*. For example, a matrix representing a differential operator with respect to a basis made of globally supported but highly localized functions will typically be pseudospars. It is common practice to replace  $A$  with a banded or sparse matrix  $\tilde{A}$  obtained by “sparsifying” (or “truncating”) the matrix outside a certain band, or by dropping entries whose magnitude falls under a certain (small) threshold. As a preliminary condition for developing linear scaling algorithms, it is essential to show that the *sparsification error*  $\|A - \tilde{A}\|$  may be bounded, for an appropriate norm, by a quantity independent of the order  $n$  of the matrix; here we show that this is indeed the case when the entries of  $A$  satisfy a certain exponential decay condition. Also, in order to approximate a matrix function  $f(A)$ , we actually compute an approximation of  $f(\tilde{A})$ . How far this is from the actual  $f(A)$  depends on several factors, including of course on how far  $\tilde{A}$  is from  $A$ ; see [65] for a general theory of condition estimation for matrix functions. It is therefore desirable to bound the sparsification error in terms of the rate of decay of the entries of  $A$ . Furthermore, it follows from our decay bounds (see section 3.4) that  $f(\tilde{A})$  is itself a matrix whose entries satisfy certain off-diagonal decay bounds; since our methods produce banded (sparse) approximations to  $f(\tilde{A})$ , we need to be able to estimate the error committed when  $f(\tilde{A})$  is approximated by a matrix with a certain bandwidth or sparsity pattern.

We say that an  $n \times n$  matrix  $A = [a_{ij}]$  has the *exponential off-diagonal decay*

*property* if there is a constant  $c > 0$  such that

$$(3.1) \quad |a_{ij}| \leq c\lambda^{|i-j|}, \quad \text{where } 0 < \lambda < 1,$$

for all  $i, j = 1 : n$ . Corresponding to the matrix  $A$  we then define for a nonnegative integer  $m$  the matrix  $A^{(m)} = [a_{ij}^{(m)}]$  defined as follows:

$$(3.2) \quad a_{ij}^{(m)} = \begin{cases} a_{ij}, & \text{if } |i-j| \leq m; \\ 0, & \text{otherwise.} \end{cases}$$

The matrix  $A^{(m)}$  is  $m$ -banded, with  $m$  being the bandwidth of  $A^{(m)}$ ; according to this definition, diagonal matrices have zero bandwidth, tridiagonal ones have bandwidth  $m = 1$ , etc. Note that we do not require the matrix to be symmetric here, we only assume that the same pattern of nonzero off diagonals is present on either side of the main diagonal. The following simple result provides an estimate of the rate at which the sparsification error decreases as the bandwidth  $m$  of the approximation increases. In addition, it establishes  $n$ -independence when the constants  $c$  and  $\lambda$  in (3.1) are independent of  $n$ . For convenience, we work with the 1-norm.

**PROPOSITION 3.2.** *Let  $A$  be a matrix with entries  $a_{ij}$  satisfying (3.1) and let  $A^{(m)}$  be the corresponding  $m$ -banded approximation. Then for any  $\varepsilon > 0$  and for  $m$  sufficiently large, it is  $\|A - A^{(m)}\|_1 \leq \varepsilon$ .*

*Proof.* We have

$$\|A - A^{(m)}\|_1 = \max_{1 \leq j \leq n} \sum_{i=1}^n |a_{ij} - a_{ij}^{(m)}| \leq \sum_{k=m+1}^n c\lambda^k \leq \frac{c\lambda^{m+1}}{1-\lambda}.$$

Since  $0 < \lambda < 1$ , for any given  $\varepsilon > 0$  we can always make

$$\frac{c\lambda^{m+1}}{1-\lambda} \leq \varepsilon.$$

by taking  $m$  sufficiently large.  $\square$

The result of Proposition 3.2 also holds for matrices with other types of decay behavior, for instance algebraic decay of the form  $|a_{ij}| \leq c|i-j|^{-p}$  with  $p > 1$ . Moreover, the argument can be adapted to handle sparse matrices rather than banded ones, along the lines to be discussed in section 3.4.

Clearly, Proposition 3.2 is not very useful when applied to a single  $n \times n$  matrix  $A$  since it is obvious that taking  $m = n$  always results in a zero error. The result, however, shows that if we have a sequence  $\{A_n\}$  of matrices of increasing dimension  $n$ , with entries  $[A_n]_{ij}$  satisfying a *uniform* exponential off-diagonal decay property of the type

$$|[A_n]_{ij}| \leq c\lambda^{|i-j|}, \quad \text{where } 0 < \lambda < 1, \quad \text{with } c \text{ and } \lambda \text{ independent of } n,$$

then for any given  $\varepsilon > 0$  it is possible to find a bandwidth  $m$  independent of  $n$  such that the  $m$ -banded approximations  $A_n^{(m)}$  satisfy  $\|A_n^{(m)} - A_n\|_1 < \varepsilon$ , for all  $n$ .

We mention that although in this section we have limited our discussion to absolute errors relative to a certain matrix norm, in practice we adopt relative error measures when assessing the distance of a computed approximation to a desired quantity.

**3.3. Faber polynomials and series.** In this section we provide some background on approximation theory that will be needed in proving general decay bounds for the entries of functions of banded or sparse matrices. Our treatment follows closely that in [75], but see also [28, 94]. In the following,  $F$  denotes a continuum containing more than one point. By a *continuum* we mean a nonempty, compact and connected subset of  $\mathbb{C}$ . Let  $G_\infty$  denote the component of the complement of  $F$  containing the point at infinity. Note that  $G_\infty$  is a simply connected domain in the extended complex plane  $\bar{\mathbb{C}} = \mathbb{C} \cup \{\infty\}$ . By the Riemann Mapping Theorem there exists a function  $w = \Phi(z)$  which maps  $G_\infty$  conformally onto a domain of the form  $|w| > \rho > 0$  satisfying the normalization conditions

$$(3.3) \quad \Phi(\infty) = \infty, \quad \lim_{z \rightarrow \infty} \frac{\Phi(z)}{z} = 1;$$

$\rho$  is the *logarithmic capacity* of  $F$ . Given any integer  $N > 0$ , the function  $[\Phi(z)]^N$  has a Laurent series expansion of the form

$$(3.4) \quad [\Phi(z)]^N = z^N + \alpha_{N-1}^{(N)} z^{N-1} + \cdots + \alpha_0^{(N)} + \frac{\alpha_{-1}^{(N)}}{z} + \cdots$$

at infinity. The polynomials

$$\Phi_N(z) = z^N + \alpha_{N-1}^{(N)} z^{N-1} + \cdots + \alpha_0^{(N)}$$

consisting of the terms with nonnegative powers of  $z$  in the expansion (3.4) are called the *Faber polynomials* generated by the continuum  $F$ .

Let  $\Psi$  be the inverse of  $\Phi$ . By  $C_R$  we denote the image under  $\Psi$  of a circle  $|w| = R > \rho$ . The (Jordan) region with boundary  $C_R$  is denoted by  $I(C_R)$ . By Theorem 3.17, p. 109 of [75], every function  $f(z)$  analytic on  $I(C_{R_0})$  with  $R_0 > \rho$  can be expanded in a series of Faber polynomials:

$$(3.5) \quad f(z) = \sum_{k=0}^{\infty} \alpha_k \Phi_k(z),$$

where the series converges uniformly inside  $I(C_{R_0})$ . The coefficients are given by

$$\alpha_k = \frac{1}{2\pi i} \int_{|w|=R} \frac{f(\Psi(w))}{w^{k+1}} dw$$

where  $\rho < R < R_0$ . We denote the partial sums of the series in (3.5) by

$$(3.6) \quad \Pi_N(z) := \sum_{k=0}^N \alpha_k \Phi_k(z).$$

Each  $\Pi_N(z)$  is a polynomial of degree at most  $N$ , since each  $\Phi_k(z)$  is of degree  $k$ . We are now ready to state a result that will be instrumental in our proof of the decay bounds.

**THEOREM 3.3. (Bernstein's Theorem)** *Let  $f$  be a function defined on  $F$ . Then given any  $\varepsilon > 0$  and any integer  $N \geq 0$ , there exists a polynomial  $\Pi_N$  of degree at most  $N$  and a positive constant  $c(\varepsilon)$  such that*

$$(3.7) \quad |f(z) - \Pi_N(z)| < c(\varepsilon)(q + \varepsilon)^N \quad (0 < q < 1)$$

for all  $z \in F$  if and only if  $f$  is analytic on the domain  $I(C_{R_0})$ , where  $R_0 = \rho/q$ . In this case, the sequence  $\{\Pi_N\}$  converges uniformly to  $f$  inside  $I(C_{R_0})$  as  $N \rightarrow \infty$ .

In the special case where  $F$  is a disk of radius  $\rho$  centered at  $z_0$ , Theorem 3.3 states that for any function  $f$  analytic on the disk of radius  $\rho/q$  centered at  $z_0$ , where  $0 < q < 1$ , there exists a polynomial  $\Pi_N$  of degree at most  $N$  and a positive constant  $c(\varepsilon)$  such that for any  $\varepsilon > 0$

$$(3.8) \quad |f(z) - \Pi_N(z)| < c(\varepsilon)(q + \varepsilon)^N,$$

for all  $z \in F$ . We are primarily concerned with the sufficiency part of Theorem 3.3. Note that the choice of  $q$  (with  $0 < q < 1$ ) depends on the region where the function  $f$  is analytic. If  $f$  is defined on a continuum  $F$  with logarithmic capacity  $\rho$  then we can pick  $q$  bounded away from 1 as long as the function is analytic on  $I(C_{\rho/q})$ . Therefore, the rate of convergence is directly related to the properties of the function  $f$ , such as the location of its poles (if there are any). Following [75], the constant  $c(\varepsilon)$  can be estimated as follows. Let  $R_0$ ,  $q$  and  $\varepsilon$  be given as in Theorem 3.3. Furthermore, let  $R'$  and  $R$  be chosen such that  $\rho < R' < R < R_0$  and

$$\frac{R'}{R} = q + \varepsilon,$$

then we define

$$M(R) = \max_{z \in C_R} |f(z)|.$$

An estimate for the value of  $c(\varepsilon)$  is asymptotically (i.e., for sufficiently large  $N$ ) given by

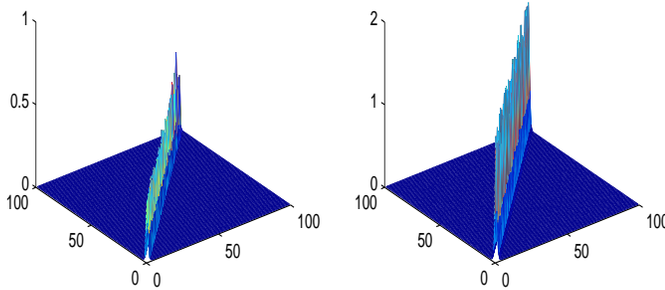
$$c(\varepsilon) \approx \frac{3}{2} M(R) \frac{q + \varepsilon}{1 - (q + \varepsilon)}.$$

It may be necessary to replace the above expression for  $c(\varepsilon)$  by a larger one to obtain validity of the bound (3.7) for all  $N$ . However, for certain regions  $F$  it is possible to obtain an explicit constant valid for all  $N \geq 0$ ; see [39] and section 5 below.

**3.4. General decay bounds.** Simple numerical experiments in MATLAB reveal that if  $A$  is a band symmetric matrix (e.g., tridiagonal) and  $f$  is a smooth function on an interval containing the spectrum of  $A$ , then most of the (relatively) large entries of  $f(A)$  are concentrated on or near the nonzero diagonals of  $A$ ; outside the band, the entries of  $f(A)$  tend to drop off rather quickly, albeit perhaps non-monotonically, and far from the main diagonal their magnitude becomes negligible. In other words,  $f(A)$  exhibits the off-diagonal decay property. This phenomenon is well-known to physicists, who sometimes call it the *nearsightedness principle*, see [67]. Further experiments suggest that a similar phenomenon holds for other types of diagonalizable matrices, including non-normal ones and often even for matrices whose eigenvectors are very far from being orthogonal. In Figure 3.1 we display “city plots” demonstrating the rapid decay in  $\log(A)$  and  $\exp(A)$  where  $A$  is a banded matrix with positive eigenvalues.

In the mathematical literature, the decay property has been first noticed for the inverse of banded, symmetric positive definite matrices; the seminal paper is [33], but see also [13, 52, 64, 77, 82]. The main result is an exponentially decaying upper bound of the form

$$|[A^{-1}]_{ij}| \leq c\lambda^{|i-j|}, \quad \text{where } c > 0 \quad \text{and} \quad 0 < \lambda < 1 \quad (1 \leq i, j \leq n).$$

FIG. 3.1. City plots for  $\log(A)$  and  $\exp(A)$ .

Here  $c$  and  $\lambda$  depend on the bandwidth and extreme eigenvalues of  $A$ , with decay being faster when the bandwidth is small and the matrix well-conditioned. The decay bound can be generalized to some extent to nonsymmetric (diagonalizable) matrices as well. Similar decay results have also been obtained for the resolvent  $(zI - A)^{-1}$ , the inverse square root  $A^{-1/2}$ , and the exponential  $\exp(A)$ ; see [63, 64, 83, 93]. In [11], Gene Golub and the first author have obtained a general decay result of the form

$$|[f(A)]_{ij}| \leq c\lambda^{|i-j|}, \quad \text{where } c > 0 \text{ and } 0 < \lambda < 1,$$

where  $A$  is banded symmetric and  $f$  is analytic on an ellipse containing the spectrum of  $A$ . This result is based on a classical result of Bernstein (different from Theorem 3.3) combined with the spectral theorem. The constants  $c$  and  $\lambda$  depend on the bandwidth of  $A$  and on the parameters of the ellipse and can be estimated explicitly. The decay result in [11] has recently found use in the study of the *entanglement* problem, which is fundamental to the field of quantum information processing; see [26, 27, 91]. The decay bound is susceptible of various immediate generalizations, for instance to matrices with a general sparsity pattern. The decay is now to be understood in terms of ‘distance’ in the graph  $G(A) = (V, E)$  associated with  $A$ , rather than distance from the main diagonal. In other words,  $|i - j|$  needs to be replaced by the graph-theoretic distance  $d(i, j)$ , defined as the length of the shortest directed path connecting node  $i$  with node  $j$ . (If there is no directed path from  $i$  to  $j$ , we set  $d(i, j) = \infty$ .) Also, the result in [11] can be immediately generalized to Hermitian matrices (rather than just real symmetric ones) and also to so-called block-banded matrices; see [27]. We further mention the decay bounds for functions of banded skew-symmetric matrices given in [31, 71] which, however, were obtained using different techniques from ours.

In the next sections we provide some further generalizations of these known results that extend and unify most existing bounds, using as our basic tool Theorem 3.3 above.

**4. Decay rates for the entries of diagonalizable matrices.** The decay bounds are obtained by translating Bernstein’s Theorem 3.3 in terms of matrices, via matrix diagonalization. Our treatment follows closely the ones in [11, 33]. Let  $A = [a_{ij}]$  be a diagonalizable matrix and let  $G(A) = (V, E)$  be the digraph associated to it with vertex set  $V$  and arc set  $E$ . We note that now the graph-theoretic ‘distance’  $d(i, j)$  is not necessarily a true distance, since in general  $d(i, j) \neq d(j, i)$ , unless  $A$  is structurally symmetric. Clearly, whenever  $d(i, j) > 1$  we have that  $a_{ij} = 0$ . More

generally, it is easy to see that  $a_{ij}^{(k)} = 0$  whenever  $d(i, j) > k$  for  $k = 0, 1, \dots$ , where  $a_{ij}^{(k)}$  denotes the  $(i, j)$  entry in the matrix  $A^k$ . To show this, we use induction over  $k$  and note that the situation is trivial for  $k = 1$ . Consider now the following identity:

$$a_{ij}^{(k+1)} = \sum_l a_{il}^{(k)} a_{lj}.$$

For pairs  $(i, l)$  such that  $d(i, l) > k$  we have that  $a_{il}^{(k)} = 0$ , by the inductive hypothesis. For pairs with  $d(i, l) \leq k$  we have that

$$k + 1 < d(i, j) \leq d(i, l) + d(l, j) \leq k + d(l, j)$$

and therefore  $d(l, j) > 1$ , hence  $a_{lj} = 0$ .

We are now in a position to state our general decay result. In the following,  $\sigma(A)$  denotes the spectrum of  $A$ .

**THEOREM 4.1.** *Let  $A$  be diagonalizable and assume that  $\sigma(A)$  is contained in a continuum  $F$ . Let  $\kappa(X)$  denote the spectral condition number of the matrix  $X$  of eigenvectors of  $A$ . Let  $f$  be a function defined on  $F$ . Furthermore, assume that  $f$  is analytic on  $I(C_{R_0}) \supset \sigma(A)$ , where  $R_0 = \frac{\rho}{q}$  with  $0 < q < 1$  and  $\rho$  is the logarithmic capacity of  $F$ . Then there are positive constants  $K$  and  $0 < \lambda < 1$  such that*

$$(4.1) \quad |[f(A)]_{ij}| < \kappa(X) K \lambda^{d(i,j)} \quad \text{for all } i, j = 1 : n.$$

*Proof.* From Theorem 3.3 we know that for any  $\varepsilon > 0$  there exists a sequence of polynomials  $\Pi_k$  of degree  $k$  which satisfies for all  $z \in F$

$$|f(z) - \Pi_k(z)| < c(\varepsilon)(q + \varepsilon)^k, \quad \text{where } 0 < q < 1.$$

We therefore have since  $A = XDX^{-1}$  that

$$\|f(A) - \Pi_k(A)\|_2 \leq \kappa(X) \max_{z \in \sigma(A)} |f(z) - \Pi_k(z)| < \kappa(X) c(\varepsilon)(q + \varepsilon)^k,$$

where  $0 < q < 1$ . For  $i \neq j$  we can write

$$d(i, j) = k + 1$$

and therefore, observing that  $[\Pi_k(A)]_{ij} = 0$  for  $d(i, j) > k$ , we have

$$|[f(A)]_{ij}| = |[f(A)]_{ij} - [\Pi_k(A)]_{ij}| \leq \|f(A) - \Pi_k(A)\|_2 < \kappa(X) c(\varepsilon)(q + \varepsilon)^{d(i,j)-1}.$$

Hence, choosing  $\varepsilon > 0$  such that  $\lambda = q + \varepsilon < 1$  and letting  $K_0 = c(\varepsilon)/(q + \varepsilon)$  we obtain

$$|[f(A)]_{ij}| < \kappa(X) K_0 \lambda^{d(i,j)}.$$

If  $i = j$  then  $|[f(A)]_{ii}| \leq \|f(A)\|_2$  and therefore letting  $K = \max\{K_0, \|f(A)\|_2\}$  we see that inequality (4.1) holds for all  $i, j$ .  $\square$

It is clear that when the matrix  $A$  is dense, the bound (4.1) is not very meaningful since the digraph  $G(A)$  will be dense as well. Hence, in practice the result is of some interest only in the (pseudo)sparse case. Also note that if  $A$  is normal, then  $X$  can be chosen to be unitary and therefore  $\kappa(X) = 1$ . This covers for example the Hermitian and skew-Hermitian cases; we have thus obtained a single proof for results that had

been previously proved using different tools. It should be clear from the proof of this result that in general one should not expect the bound to be very sharp. In particular, when  $A$  is highly non-normal (in the sense that  $\kappa(X) \gg 1$ ) the bound can be a severe overestimate of the actual size of the elements of  $f(A)$ , and in this case other tools, such as for example pseudospectra, may be better suited to characterize the actual decay behavior; see [97, 98].

We now specialize the result to the case of band matrices. As  $A$  is not assumed to have symmetric structure, it is necessary to account for the fact that for a nonsymmetric matrix, the rate of decay can be different above and below the main diagonal. For instance,  $A$  could have different numbers of nonzero diagonals below and above the main diagonal. An extreme case is when  $A$  is an upper (lower) Hessenberg matrix, in which case  $f(A)$  typically exhibits fast decay below (above) the main diagonal, and generally no decay above (below) it. In Figure 4.1 we display on the left the magnitude of the entries of  $f(A) = \cos(A)$  where  $A$  is a band matrix with one nonzero diagonal below and four nonzero diagonals above the main diagonal. The matrix is diagonalizable, but the condition number  $\kappa(X)$  of the eigenvector matrix is of the order of  $10^{10}$ . The plot on the left shows the magnitude of the entries on the 50th row of  $\cos(A)$ .

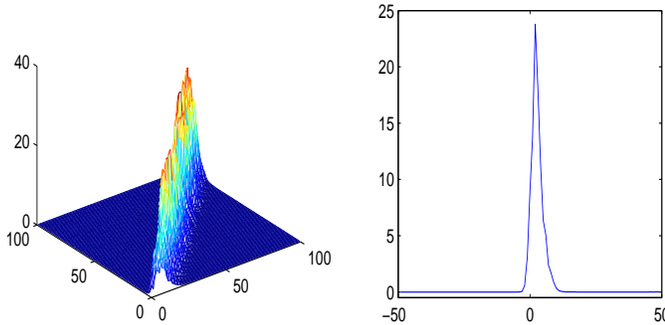


FIG. 4.1. *Cosine of a nonsymmetric band matrix.*

We establish the appropriate decay results next. We give the proof of the bound for entries in the lower part of  $f(A)$  only; the proof for the entries in the upper part of  $f(A)$  is identical, with the obvious changes.

We say that a matrix  $A$  has *lower bandwidth*  $p > 0$  if  $a_{ij} = 0$  whenever  $i - j > p$  and *upper bandwidth*  $s > 0$  if  $a_{ij} = 0$  whenever  $j - i > s$ . We note that if  $A$  has lower bandwidth  $p$  then  $A^k$  has lower bandwidth  $kp$  for  $k = 0, 1, 2, \dots$ , and similarly for the upper bandwidth  $s$ .

**THEOREM 4.2.** *Let  $A$  be diagonalizable with upper bandwidth  $p$  and lower bandwidth  $s$ . Assume that  $\sigma(A)$ , is contained in a continuum  $F$ . Let  $f$  be a function defined on  $F$ . Furthermore, assume that  $f$  is analytic on  $I(C_{R_0}) \supset \sigma(A)$  where  $R_0 = \frac{\rho}{q}$  with  $0 < q < 1$  and  $\rho$  is the logarithmic capacity of  $F$ . Then there are positive constants  $K, C$  and  $0 < \lambda, \eta < 1$  such that for  $i \geq j$*

$$(4.2) \quad |[f(A)]_{ij}| < \kappa(X)K\lambda^{i-j}$$

and for  $i < j$

$$(4.3) \quad |[f(A)]_{ij}| < \kappa(X)C\eta^{j-i}.$$

*Proof.* Observe first that for  $k = 0, 1, \dots$  the matrix  $\Pi_k(A)$  has lower bandwidth  $kp$  for all polynomials  $\Pi_k(z)$  of degree at most  $k$ . From Bernstein's Theorem we know that for any  $\varepsilon > 0$  there exists a sequence of polynomials  $\Pi_k(z)$  of degree  $k$  which satisfies for all  $z \in F$

$$|f(z) - \Pi_k(z)| < c(\varepsilon)(q + \varepsilon)^k, \quad \text{where } 0 < q < 1.$$

We therefore have since  $A = XDX^{-1}$  that

$$\|f(A) - \Pi_k(A)\|_2 \leq \kappa(X) \max_{z \in \sigma(A)} |f(z) - \Pi_k(z)| < \kappa(X)c(\varepsilon)(q + \varepsilon)^k$$

where  $0 < q < 1$ . For  $i > j$  write

$$i - j = kp + l \quad \text{for } l = 1, \dots, p;$$

then we have that

$$\frac{i - j}{p} - 1 \leq k$$

and hence observing that  $[\Pi_k(A)]_{ij} = 0$  for  $i - j > pk$ ,

$$|[f(A)]_{ij}| = |[f(A)]_{ij} - [\Pi_k(A)]_{ij}| \leq \|f(A) - \Pi_k(A)\|_2 < \kappa(X)c(\varepsilon)(q + \varepsilon)^{\frac{i-j}{p}-1}.$$

Hence, letting  $\lambda = (q + \varepsilon)^{\frac{1}{p}}$  and  $K_0 = c(\varepsilon)/(q + \varepsilon)$  we obtain

$$|[f(A)]_{ij}| < \kappa(X)K_0\lambda^{i-j}.$$

If  $i = j$  then  $|[f(A)]_{ii}| \leq \|f(A)\|_2$  and therefore letting  $K = \max\{K_0, \|f(A)\|_2\}$  we see that inequality (4.2) holds for all  $i \geq j$ .  $\square$

An immediate corollary from Theorem 4.2 is the following.

**COROLLARY 4.3.** *Let  $A$  be a banded diagonalizable matrix. Assume that  $\sigma(A)$  is contained in a continuum  $F$ . Let  $f$  be a function defined on  $F$ . Furthermore, assume that  $f$  is analytic on  $I(C_{R_0})$  ( $\supset \sigma(A)$ ), where  $R_0 = \frac{\rho}{q}$  with  $0 < q < 1$  and  $\rho$  is the logarithmic capacity of  $F$ . Then there are positive constants  $K$  and  $0 < \lambda < 1$  such that for all  $i, j=1:n$ ,*

$$(4.4) \quad |[f(A)]_{ij}| < \kappa(X)K\lambda^{|i-j|}.$$

We note that for  $A$  real and symmetric, this is essentially Theorem 2.2 in [11].

**5. More refined bounds.** The bounds obtained in the previous section can be improved by using sharper estimates on the Faber polynomials, provided that some (fairly reasonable) additional conditions are imposed on the continuum  $F$ . As already mentioned at the end of section 3.3, in [39] the author presents more quantitative estimates on the  $N$ th Faber polynomial for certain choices of  $F$ . Specifically, assume that  $F$  is a closed *Jordan region*. By a Jordan region we mean a region  $F$  that is bounded and whose boundary  $\Gamma$  consists of pairwise disjoint closed Jordan curves. If  $\Gamma$  is rectifiable, there exists at almost every point  $z \in \Gamma$  a tangent vector that makes

an angle  $\Theta(z)$  with the positive real axis. We say that  $\Gamma$  has *bounded total rotation*  $V$  if

$$V = \int_{\Gamma} |d\Theta(z)| < \infty.$$

It is apparent that  $V \geq 2\pi$  and equality holds if  $F$  is convex. In the case where  $F$  degenerates to a *Jordan arc*, the following results remain valid; we refer the reader to [39] for details.

**THEOREM 5.1.** *Let  $F$  be a Jordan region whose boundary  $\Gamma$  is of bounded total rotation  $V$ . Then the following holds:*

(a) *For  $N \geq 1$ ,*

$$\|\Phi_N\|_{\infty} \leq \rho^N V / \pi.$$

*This bound holds with equality when  $F$  coincides with the interval  $[-1, 1]$ .*

(b) *Let  $f$  be analytic on the closed region  $C_R$ ,  $R > \rho$ , then for  $N \geq 0$  we have*

$$\|f - \Pi_N\|_{\infty} \leq \frac{M(R)V}{\pi} \frac{(\rho/R)^{N+1}}{1 - \rho/R},$$

*where  $M(R) = \max_{z \in C_R} |f(z)|$  and  $\Pi_N(z)$  are the polynomials defined by equation (3.6).*

These bounds can be used to improve our earlier estimates in the case where  $F$  and  $f$  satisfy the conditions of Theorem 5.1. If in addition  $A$  satisfies the conditions of Theorem 4.1 we can obtain the following bound for the entries of  $f(A)$ :

$$(5.1) \quad |[f(A)]_{ij}| \leq \kappa(X) K \lambda^{d(i,j)},$$

where now  $K = \max \left\{ \|f(A)\|_2, \frac{M(R)V}{\pi(1-\rho/R)} \right\}$  and  $\lambda = \rho/R$ .

We now compare the above bound on the entries of  $f(A)$  with the magnitude of the actual entries  $[f(A)]_{ij}$ , for  $i < j$ , in a special case. The matrix  $A$  is tridiagonal, with spectrum in  $F = [-1, 1]$ . In this case  $\rho = 1/2$  and since  $F$  is convex we have that  $V = 2\pi$ . The function  $f$  must be analytic on the ellipse

$$\frac{x^2}{\left(\frac{R}{2\rho} + \frac{\rho}{2R}\right)^2} + \frac{y^2}{\left(\frac{R}{2\rho} - \frac{\rho}{2R}\right)^2} = 1.$$

This is an ellipse with foci at  $\pm 1$ , the sum of whose semiaxes is  $R/\rho$ . For our experiment we take  $f(z) = e^z$ , which is entire and therefore satisfies our assumptions. The constant  $M(R)$  is then given by

$$M(R) = e^{R + \frac{1}{4R}}.$$

It is important to note that (5.1) gives us a bound that changes with  $R$ . We can use small values of  $R$  to estimate the elements on the first few off-diagonals, and increase  $R$  as we move away from the diagonal. This adaptive strategy allows us to obtain sharper bounds than those described in the previous section. In Figure 5.1 we show the upper bound compared to the actual magnitude of the off-diagonal entries in the first row of  $f(A) = \exp(A)$ , in both the linear and logarithmic scale. Clearly, the

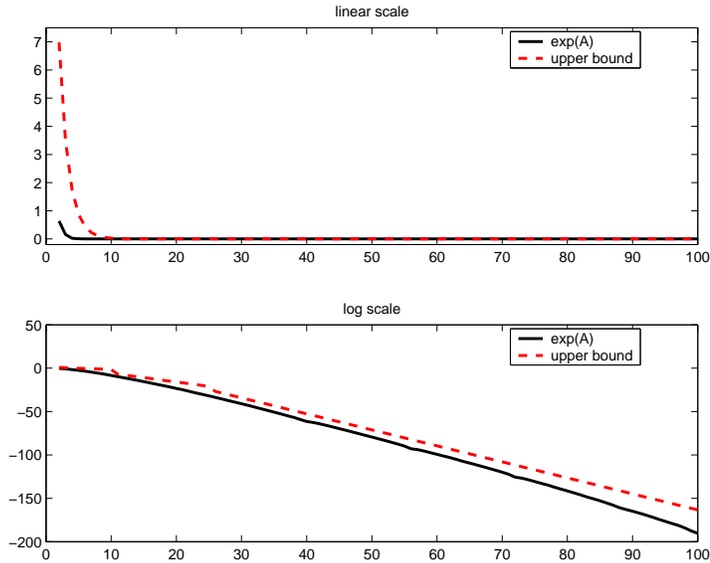


FIG. 5.1. Upper bound (5.1) vs. the entries of  $\exp(A)$ , first row.

bound does a good job capturing the decay behavior, especially as one moves away from the main diagonal.

The decay bounds obtained so far can be used to predict a minimum bandwidth (more generally, sparsity pattern) for the computed approximation to  $f(A)$  to satisfy a prescribed error tolerance; if the rate of decay, expressed in terms of the various quantities entering the upper bound, does not depend on  $n$ , then an  $O(n)$  approximation is feasible. A sufficient condition is that the  $n \times n$  matrix sequence  $\{A_n\}$  is such that for all  $n$ , the spectra  $\sigma(A_n)$  are all contained in a region that is bounded and bounded away from the singularities of  $f$  (if any); furthermore, the condition numbers of the corresponding eigenvector matrices must be bounded independent of  $n$ . Although these conditions seem rather restrictive, they are satisfied by large classes of matrices and functions that arise in applications, for example when  $f$  is entire and  $\{A_n\}$  is a sequence of normal (e.g., Hermitian) matrices with bounded spectra. If  $f$  is not entire (for example if  $f(z) = z^{-1}$ , or  $f(z) = z^{-1/2}$ , or  $f(z) = \log(z)$ ) then the spectra of the  $\{A_n\}$  must remain bounded away from the singularities of  $f$ . As an example,  $O(n)$  approximation of the inverse  $A_n^{-1}$  is possible if there is a compact set that contains all the spectra  $\sigma(A_n)$  and has positive distance from  $z = 0$ , provided that the eigenvector matrices have condition number that remains bounded as  $n \rightarrow \infty$ .

**6. Approximation algorithms.** In this section we give a description of the algorithms we use to compute approximations to functions of band matrices. As an integral part of the approach used, we also describe a procedure that can be used to determine a bandwidth for the approximation that captures all the relevant entries to a prescribed tolerance. In the following, when we write  $f(A)$  we are implicitly assuming that the function  $f(z)$  is defined on the spectrum of the matrix  $A$  and therefore the matrix function  $f(A)$  is well defined.

To approximate  $f(A)$  we consider algorithms that are based upon polynomial approximations. The problem is thus reduced to that of finding rapidly converging

polynomial approximations to the scalar function  $f(z)$  on a domain in the complex plane. This problem has been extensively studied in the approximation theory and numerical analysis literature; see, e.g., [40, 43, 92]. Depending on the location of the spectrum, different approaches can be used. If the eigenvalues of  $A$  lie on a line segment in  $\mathbb{C}$ , and estimates for the endpoints of the segment are available, it is possible to scale and shift the matrix  $A$  so that the resulting matrix has its spectrum in the interval  $[-1, 1]$ . For instance, let  $A$  have real spectrum with  $a = \lambda_{\min}(A)$  and  $b = \lambda_{\max}(A)$ . The affine linear function

$$\xi : \mathbb{C} \longrightarrow \mathbb{C} \quad \text{given by} \quad \xi(\lambda) = \frac{2\lambda - (a + b)}{b - a}$$

satisfies  $\xi([a, b]) = [-1, 1]$  and therefore the spectrum of  $B = \xi(A) = \frac{2}{b-a}A - \frac{a+b}{b-a}I$  is contained in  $[-1, 1]$ . Hence, we can use the well known three-term recurrence relation to generate the (optimal) Chebyshev polynomials and obtain an approximation to  $f(B)$  by truncating the Chebyshev series. The approximation for the original matrix function  $f(A)$  can then be easily obtained from that of  $f(\xi^{-1}(B))$  by inverting  $\xi$ . For simplicity of notation, we will assume that the matrix  $A$  has already been scaled and shifted so that its spectrum is contained in the interval  $[-1, 1]$ .

For more general matrices, whose eigenvalues lie within a region  $F \subset \mathbb{C}$  more complicated than a line segment, it is possible to generalize the method above to some extent by means of Faber polynomials; see, e.g., [14, 80, 81]. For certain domains  $F$  the Faber polynomials satisfy a three-term recurrence relation similar to that of the Chebyshev polynomials; for instance, this is the case when the spectrum lies in the interior of an ellipse on which  $f$  is analytic. However, for more general regions  $F$  the  $k$ th Faber polynomial satisfies a  $k$ -term recurrence relation. From the point of view of storage requirements, this is a serious drawback. Some truncation and restarting strategies have been considered in the literature; here we consider instead an alternative approximation technique based on Newton-type interpolation, already used in [95].

**6.1. Chebyshev expansion.** Expansion in the Chebyshev basis has long been a popular approach for computing approximations to the density matrix in electronic structure calculations; see, e.g., [2, 48]. The method can be used to approximate a wide variety of matrix functions, as long as the matrix can be easily transformed to one whose eigenvalues lie in  $[-1, 1]$ . We start by recalling the matrix version of the classical three-term recurrence relation for the Chebyshev polynomials:

$$(6.1) \quad T_{k+1}(A) = 2AT_k(A) - T_{k-1}(A), \quad k = 1, 2, \dots$$

(with  $T_0(A) = I$ ,  $T_1(A) = A$ ). These matrices can be used to obtain an approximation

$$f(A) = \sum_{k=1}^{\infty} c_k T_k(A) - \frac{c_1}{2}I \approx \sum_{k=1}^N c_k T_k(A) - \frac{c_1}{2}I$$

to  $f(A)$  by truncating the Chebyshev series expansion after  $N$  terms. The coefficients  $c_k$  in the expansion only depend on  $f$  (not on  $A$ ) and can be easily computed numerically at a cost independent of  $n$  using the approximation

$$c_k \approx \frac{2}{M} \sum_{j=1}^M f(\cos(\theta_j)) \cos((k-1)\theta_j),$$

where  $\theta_j = \pi(j - \frac{1}{2})/M$ . Thus, most of the computational work is performed in (6.1). The basic operation in (6.1) is the matrix–matrix multiply. If the initial matrix  $A$  is  $m$ -banded, then after  $k$  iterations the matrix  $T_{k+1}(A)$  will be  $km$ -banded. In order to have a linear scaling algorithm, it is essential to fix a maximum bandwidth for the approximation  $P_N(A)$ , which must not depend on  $n$ . Then the cost is dominated by the matrix–matrix multiplies, and this is an  $O(n)$  operation provided that the maximum bandwidth remains bounded as  $n \rightarrow \infty$ . Similar conclusions apply for more general sparsity patterns, which can be determined by using the structure of successive powers  $A^k$  of  $A$ . In alternative, dropping elements by size using a drop tolerance is often used, although rigorous justification of this procedure is more difficult.

Let us now consider the error incurred by the series truncation:

$$(6.2) \quad \|e_N(A)\| = \|f(A) - P_N(A)\|,$$

where  $P_N(A) = \sum_{k=1}^N c_k T_k(A) - \frac{c_0}{2} I$  and  $\|\cdot\|$  is, for instance, the matrix 2-norm. We limit our discussion to the banded case, but the same arguments apply in the case of general sparsity patterns as well. Since  $|T_k(x)| \leq 1$  for all  $x \in [-1, 1]$  and  $k = 1, 2, \dots$ , we have that  $\|T_k(A)\| \leq 1$  for all  $k$ , since  $\sigma(A) \subset [-1, 1]$ . Using this well-known property to bound the error in (6.2), we obtain that

$$\|e_N(A)\| = \left\| \sum_{k=N+1}^{\infty} c_k T_k(A) \right\| \leq \sum_{k=N+1}^{\infty} |c_k|.$$

The last inequality shows that the error defined by (6.2) only depends on the sum of the absolute values of the coefficients  $c_k$  for  $k = N + 1, N + 2, \dots$ , but these in turn do not depend on  $n$ , the dimension of the matrix we are approximating. Hence if we have a sequence of  $n \times n$  matrices  $\{A_n\}$  with  $\sigma(A_n) \subset [-1, 1]$  for all  $n$ , we can use an estimate of the quantity  $\sum_{k=N+1}^{\infty} |c_k|$  and use that to prescribe a sufficiently large bandwidth (sparsity pattern) to ensure a prescribed accuracy of the approximation. As long as the bandwidth of the approximation does not exceed the maximum prescribed bandwidth, the error is guaranteed to be  $n$ -independent. In practice, however, we found that this strategy is too conservative. Because of the rapid decay outside of the bandwidth of the original matrix, it is usually sufficient to prescribe a much smaller maximum bandwidth than the one predicted by the truncation error. This means that numerical dropping is necessary (see section 6.3 below for our recommended dropping strategy), since the bandwidth of  $P_N(A)$  rapidly exceeds the maximum allowed bandwidth. Because of dropping, the simple error estimate given above is no longer rigorously valid. Our numerical experiments, however, suggest that  $n$ -independence (and therefore linearly scaling complexity and storage requirements) is maintained.

**6.2. General polynomial approximation.** We now turn to the problem of approximating  $f(A)$  when the spectrum of  $A$  is contained in an arbitrary continuum  $F \subset \mathbb{C}$ ; for a more detailed description of the technique we use, see [95]. The (Newton) interpolation polynomial we use is of the form

$$P_N(A) = c_0 I + c_1(A - z_0 I) + c_2(A - z_0 I)(A - z_1 I) + \dots + c_N(A - z_0 I) \dots (A - z_{N-1} I)$$

where  $c_k$  is the divided difference of order  $k$ , i.e.,

$$c_k = f[z_0, \dots, z_k], \quad k = 0, \dots, N.$$

For  $k = 0, \dots, N - 1$ , the interpolation points are chosen as  $z_k = \Psi(\omega_k)$ , where  $\omega_k$  are the  $N - 1$  roots of the equation  $\omega^{N-1} = \rho$  and  $\Psi(z)$  is the inverse of the map  $\Phi(z)$  that maps the complement of  $F$  to the outside of a disk with radius  $\rho$  and satisfies the normalization conditions (3.3). This method does not require the computation of Faber polynomials and their coefficients. However, it does require knowledge of the map  $\Psi(z)$ . For specific domains  $F$  this map can be determined analytically, see for example [14, 95]. In addition,  $\Psi(z)$  may require information on the convex hull of the eigenvalues of  $A$ . For more general domains one may have to resort to numerical approximations to compute  $\Psi(z)$ ; see [96]. Once again, the approximation algorithm requires mostly matrix–matrix multiplies with banded (or sparse) matrices and appropriate sparsification is generally required to keep the cost within  $O(n)$  as the problem size  $n$  grows. A rigorous error analysis that takes dropping as well as truncation into account is currently being attempted.

**6.3. Dropping.** We combine the above approaches with a dropping strategy. The idea applies to more general sparsity patterns, but we restrict our discussion to the case where  $A$  is a banded matrix with bandwidth  $m$ . In this case we only keep elements inside a prescribed bandwidth  $\hat{m}$  in every iteration. For given  $\rho$  and  $R$  we choose  $\hat{m}$  a priori so as to guarantee that

$$(\rho/R)^{\hat{m}} \approx \varepsilon/C$$

where  $C > 0$  is the constant for the bounds on  $[f(A)]_{ij}$  (with  $i \neq j$ ) discussed in section 5 and  $\varepsilon > 0$  is a prescribed tolerance. Note that  $C$  is independent of the norm of  $f(A)$  since we do not expect the diagonal elements to be insignificant.

As already noted, if  $A$  is banded with bandwidth  $m$ , then  $A^k$  has bandwidth  $km$ . This means that if we want the approximation to have a fixed bandwidth  $\hat{m}$ , where  $\hat{m}$  is (say) an integer multiple of  $m$  corresponding to a prescribed approximation error  $\varepsilon$ , then we ought to truncate the expansion at the  $N^*$ th term, with  $N^* = \hat{m}/m$ . It may happen, however, that this value of  $N$  is actually too small to reduce the error below the prescribed threshold. In this case it is necessary to add extra terms to the Chebyshev expansion; but this would lead to an increase of the bandwidth beyond the prescribed limit. A solution that has been used by physicists is simply to continue the recurrence but to ignore all entries in positions outside the prescribed bandwidth; see, e.g., [2, 48, 89]. The important fact is that by restricting all the terms in the Chebyshev expansion (6.1) to have a fixed bandwidth (independent of  $n$  and  $N$ ) we obtain an approximation scheme whose cost scales linearly in the size  $n$  of the problem.

**6.4. Stopping criteria.** In alternative to strategies that fix the number of terms in the polynomial expansion a priori (with or without dropping), one could decide to continue to add additional terms to the approximation to  $f(A)$  until some convergence criterion is satisfied. In other words, a dynamic criterion rather than a static, a priori one could be used. For instance, with the Chebyshev expansion method one can monitor the size of the coefficients  $c_k$  and stop iterating when these are sufficiently small. Also, for some matrix functions a natural notion of *residual* exists. For example in the case of  $f(A) = A^{-1}$ , stopping criteria could be formulated on the basis of the size (in some appropriate norm) of the matrix  $R = I - AM$  (or  $R = I - MA$ ), where  $M$  is the current computed approximation. For  $f(A) = A^{1/2}$ , where  $A$  is Hermitian and positive definite, one can use  $\|R\| = \|A - M^2\|$ , and  $\|R\| = \|I - MAM\|$  for  $f(A) = A^{-1/2}$ . Relative versions of these residuals can also be used, if an estimate of  $\|f(A)\|$  is available.

For many other types of matrix functions, however, no natural notion of residual seems to be readily available. In this case one may decide to stop the approximation process if significant progress is no longer being made. A possibility would be to stop iterating when the norm of the difference  $\|M_{new} - M_{old}\|$  between two successive approximations is below a prescribed threshold  $\delta > 0$ . Another possibility, which makes sense in some applications, could be to monitor the change in the trace; that is, the process is stopped when  $|\text{trace}(M_{new}) - \text{trace}(M_{old})| < \delta$ . Such criteria may fail in case of temporary stagnation; see [42] for some discussion of this issue, although in a somewhat different context. A rigorous justification of such criteria, however, is still lacking, and more work is needed to devise reliable dynamic stopping criteria.

**7. Some applications.** The approximation of matrix functions is of great importance in several areas including quantum chemistry (electronic structure calculations), quantum information processing, lattice quantum chromodynamics, numerical solution of time-dependent differential equations (both ODEs and PDEs), statistics, and so forth. Approximations of matrix functions such as  $A^{-1}$  and  $A^{-1/2}$  can also be useful in the context of developing preconditioners for  $Ax = b$ . See, e.g., [10, 11, 83]. In this section we briefly describe some of these applications.

**7.1. Density matrix computations.** In quantum mechanics the state of a system is described by a vector of unit norm in a (generally infinite-dimensional) complex Hilbert space  $\mathcal{H}$ , while physical quantities of interest (like momentum, energy, etc.) are usually represented by Hermitian linear operators acting on  $\mathcal{H}$ . The eigenvalues  $\{\lambda_i\}$  of an operator represent the possible values for the corresponding quantity, and the associated normalized eigenvectors  $\psi_i$  are the stationary states; see, e.g., [73, 74, 100]. If  $A$  is an operator representing the observable  $\alpha$ , then the expected value of  $\alpha$  in the state  $\psi$  ( $\psi \in \mathcal{H}$ ,  $\|\psi\| = 1$ ) is given by  $\langle \alpha \rangle = (A\psi, \psi)$  where  $(\cdot, \cdot)$  denotes the inner product in  $\mathcal{H}$ . The probability that  $\alpha$  will take the value  $\lambda_i$  is equal to  $|\langle \psi, \psi_i \rangle|^2$ . If the system happens to be in one of the stationary states  $\psi_i$ , then  $\alpha = \lambda_i$  with probability one. If  $\psi$  is expressed as a linear combination of eigenvectors  $\psi_i$  of  $A$ ,  $\psi = c_1\psi_1 + c_2\psi_2 + \dots + c_m\psi_m$ , then clearly  $\langle \alpha \rangle = c_1\lambda_1 + c_2\lambda_2 + \dots + c_m\lambda_m$ . An alternative expression that requires no explicit knowledge of the eigenvalues and eigenvectors of  $A$  is the following. If  $P$  is the orthogonal projector onto a subspace of  $\mathcal{H}$  spanned by an *ensemble* of states, the expected value of  $\alpha$  relative to that ensemble is given by  $\langle \alpha \rangle = \text{trace}(AP) = \text{trace}(PA)$ . For example, if the quantity of interest is the total energy  $\eta$  then  $\langle \eta \rangle = \text{trace}(HP)$  where  $H$  is the Hamiltonian operator. If the subspace is one-dimensional, spanned by the single state vector  $\psi$ , then  $P = \psi\psi^H$  and we recover the expression  $\langle \alpha \rangle = (A\psi, \psi)$  for the expected value of the observable corresponding to  $A$  in the state  $\psi$ . In practice, the infinite-dimensional operators are approximated by matrices by truncating a basis  $\{\phi_i\}_{i=1}^{\infty}$  for  $\mathcal{H}$  and representing the operators with respect to this finite basis  $\{\phi_i\}_{i=1}^n$ . The matrix  $P$ , originally introduced by von Neumann [100], is called the *density matrix* for the states of interest (the so-called *occupied states*, which span the range of  $P$ ). Once  $P$  is known, all statistical properties of the system under study are easily obtained. (In the *mixed case*,  $P$  is no longer a projector but a convex linear combination of projectors; it is still Hermitian and its eigenvalues lie between 0 and 1.) The important fact here is that  $P$  can be computed without knowledge of the eigenvalues and eigenvectors of  $A$ . Indeed,  $P$  can be expressed as a function of the Hamiltonian  $H$ . Although this function is discontinuous (essentially a step, or Heaviside function) it can be approximated very closely by a smooth function  $P \approx f(H)$ . Various expressions have been used for  $f$  in the literature; see, e.g., [2, 5, 48, 89]. An especially popular form is the so called

*Fermi–Dirac density matrix* (FDM):

$$P \approx f(H), \quad \text{where} \quad f(z) = \frac{1}{1 + e^{\beta(z-\mu)}}.$$

Here  $\mu$  is a shift parameter, called the *chemical potential*, which can be chosen so as to have  $\text{trace}[f(H)] = N_e$ , the number of electrons (occupied states) in the system. The parameter  $\beta$  is called the *inverse temperature* and controls how close the FDM is to the true density matrix; see, e.g., [2]. The exponential off-diagonal decay property of the density matrix, which is closely related to the localization of the eigenvectors of the Hamiltonian  $H$ , is well-established in the physics literature; see, e.g., [1, 48, 57, 67, 76]. Since the early 1990’s, this property has been exploited in the development of  $O(n)$  methods for electronic structure calculation [19, 44, 48, 67, 70, 78, 88, 89]. Here we note that since  $P$  is well-approximated by a smooth function of a sparse symmetric matrix  $H$  (the Hamiltonian is usually well-localized for a suitable choice of the basis set), the decay property also follows from Theorem 2.2 in [11]. The city plot in Figure 7.1 shows the rapid decay in the Fermi–Dirac density matrix corresponding to a “one-dimensional Anderson model,” see [1]. That is, the matrix function is  $f(H) = (\exp(\beta(H - \mu I)) + I)^{-1}$  where  $H$  is a  $200 \times 200$  symmetric tridiagonal matrix with random entries on the main diagonal (drawn from the uniform distribution on  $[0, 1]$ ) and with the remaining nonzero entries equal to  $-1$ . In this example we used  $\beta = 2$  and  $\mu = 0.5$ .

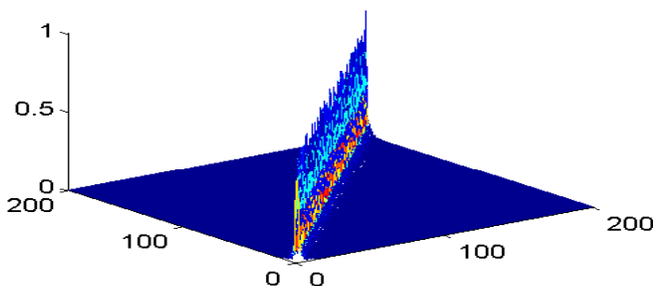


FIG. 7.1. City plot for a Fermi–Dirac density matrix.

Summarizing, in applications of quantum mechanics one is interested in computing quantities of the form  $\text{trace}[Af(H)]$  for given matrices  $A$  and  $H$ , for a smooth function  $f$ ; in particular, the expected value of the energy in a given configuration is given by an expression of the form  $\langle \eta \rangle = \text{trace}[Hf(H)]$ ; note that the function of interest now is  $g(z) = zf(z)$ . While some fast approximate methods for the evaluation of quantities of this kind have been developed by quantum chemists and physicists, virtually no rigorous justification of such algorithms was previously available. Some of the results in this paper can be regarded as providing such foundation.

**7.2. Non-Hermitian matrices.** The bounds and algorithms for functions of non-Hermitian matrices will also be tested on other problems arising from physics. In recent years, the field of *non-Hermitian quantum mechanics* has emerged as one of great importance; see, e.g., [8, 9, 21, 55, 97]. In this physical theory, the operators and matrices that describe physical quantities (like the total energy) are non-Hermitian but have real eigenvalues; such matrices are called *R-matrices* in [73]. As in the

Hermitian case, this theory makes use of certain functions of the (non-Hermitian) Hamiltonian related to the eigenvectors, like the density matrix or the resolvent, which once again have been found to exhibit off-diagonal decay behavior. (Non-Hermitian matrices with real spectra also arise in fluid mechanics, see [12].) The observed decay behavior can be analyzed, in part, in the context of the general theory developed in this paper. As mentioned earlier, however, in the case of highly non-normal operators other tools (like pseudospectra, see [97, 98]) provide the necessary insight.

**7.3. Trace and determinant.** Additional possible applications of the theory and methods discussed in the present paper include fast algorithms for approximating the determinant of a large, sparse symmetric positive definite matrix; see [6, 62, 69, 85] for motivation and background. One possible approach that we are investigating is the reduction of the computation of the determinant to that of the trace of a matrix function; for instance, one could exploit the straightforward identity  $\det(A) = \exp(\text{trace}[\log(A)])$ . If the smallest eigenvalue of  $A$  is not too close to zero, approximating the trace of  $\log(A)$  is not difficult and good results can be expected.

**7.4. Other applications.** In addition, cheaply computed, low-accuracy approximations to  $f(A)$  may also be useful to construct initial guesses and/or preconditioners for certain iterative methods for computing  $f(A)b$ ; see [23, 84, 99] for recent ideas on the problem of preconditioning the evaluation of  $f(A)b$ . As another example, our decay bounds may be useful in devising stopping criteria for projection (Lanczos or Krylov subspace) methods for evaluating  $f(A)v$ . These methods construct low-dimensional subspaces and obtain successive approximations to  $f(A)v$  from these subspaces by explicitly computing  $f(H_m)$  where  $H_m$  is a tridiagonal or upper Hessenberg matrix of small dimension  $m$  (equal to that of the subspace). It turns out that the magnitude of the entries in the first column of  $f(H_m)$  plays an important role in these algorithms (e.g., in devising stopping criteria); hence, bounds on the size of these elements may be used to estimate the number of iterations needed to achieve a certain accuracy.

**8. Numerical experiments.** In this section we present the results of a few preliminary numerical experiments aimed at illustrating the performance of the approximation algorithms discussed in section 6, with particular attention to linear scaling behavior. We performed several tests involving different types of matrices and functions. We regard our approximation schemes as iterative methods, with each new iteration consisting of the inclusion of an additional term in the polynomial expansion. In our computations we exploit the decay property of the entries in  $f(A)$  to keep a fixed number  $\hat{m}$  of off-diagonals in every iteration. By  $N$  we denote the number of terms in the expansion that were necessary to obtain the indicated error. The errors were obtained by comparing the computed approximation to the exact computation via MATLAB's `funm` command. Unless otherwise specified, we measure the relative error in the Frobenius norm.

In Table 8.1 we show results for the Fermi–Dirac density matrix discussed in section 7.1. Here  $A$  is the 1-D Laplacian with Dirichlet end point conditions and the diagonal entries replaced by random entries from  $[0, 1]$ . We approximate the function  $f(z) = \frac{1}{1 + \exp(\beta(z - \mu))}$  for different values of  $\mu$  and  $\beta$  (which are physically realistic) using a Chebyshev series expansion. We impose a maximum bandwidth of  $\hat{m}$  and ignore entries outside such band. The results clearly show the  $n$ -independent behavior (and hence, linear scaling complexity) of the algorithm.

We now discuss how the values of  $N$  and  $\hat{m}$  were determined. Note that since

TABLE 8.1  
 Results for  $f(z) = \frac{1}{1+\exp(\beta(z-\mu))}$

$n$	$\mu = 2, \beta = 2.13$			$\mu = 0.5, \beta = 1.84$		
	error	$N$	$\hat{m}$	error	$N$	$\hat{m}$
100	9e-06	18	20	6e-06	18	22
200	4e-06	19	20	9e-06	18	22
300	4e-06	19	20	5e-06	20	22
400	6e-06	19	20	8e-06	20	22
500	8e-06	19	20	8e-06	20	22

the Fermi–Dirac density matrix is an approximation of an orthogonal projector of known rank, special stopping criteria can be used in this case. However, rather than using such criteria we adopted a more general strategy. The choice of the number  $N$  of terms to include in the Chebyshev series expansion is an easy matter, since the magnitude of the coefficients can be monitored. Note that if  $f$  is analytic, the coefficients  $c_k$  decay exponentially fast. Once the sum of the absolute values of three consecutive coefficients is sufficiently small, no additional terms are added; here we used the rather conservative tolerance of  $1e^{-15}$ . Although not very sophisticated, this criterion gave good results in practice. We obtained the a priori bandwidth  $\hat{m} = 20$  as follows. We first note that  $|f(z)| \leq 1$  for all  $z$ , therefore the constant  $M(R)$  in the bounds for  $|[f(A)]_{ij}|$  can be neglected. To perform the computation in the Chebyshev basis we scale and shift the matrix  $A$  so that its spectrum lies in  $[-1, 1]$ . To obtain the approximation to  $f(A)$  with the original  $A$  we shift and scale back the coefficients in the Chebyshev series. The logarithmic capacity of  $[-1, 1]$  is given by  $\rho = 1/2$ , and  $V \frac{1}{\pi(1-\rho/R)} = 4$  when  $R = 1$ . The entries  $(i, j)$  in  $f(A)$ , where  $i$  and  $j$  are such that  $|i - j| \geq 20$ , have magnitude at most  $3e-06$  and can therefore be neglected. For the second case ( $\mu = 0.5, \beta = 1.84$ ) we had to slightly increase  $N$  and  $\hat{m}$  to achieve a comparable error. As a rule, it is a good idea to use values of  $\hat{m}$  that are slightly greater than those predicted by the theory, also in view of the effects of dropping.

We performed many more experiments with the Chebyshev method and obtained excellent results whenever  $f$  is an entire function. If  $f$  has singularities, however, we found that the method can converge quite slowly, especially when the singularities are close to the spectrum of  $A$ . In these cases, the approach described in section 6.2 was found to be much better. Therefore, in the remaining tables we use the interpolation approach. The results in Table 8.2 were obtained by interpolating the “entropy” function  $f(z) = z \log(z)$  to obtain an approximation to  $f(A) = A \log(A)$ . As before, the values of  $N$  and  $\hat{m}$  that are guaranteed to result in a given error can be estimated a priori from knowledge of the parameters  $\rho$  and  $R$ . The matrix  $A$  is a tridiagonal matrix with random diagonal entries, shifted so as to be diagonally dominant (and therefore positive definite, so that the logarithm is well-defined). We also report the absolute error in the trace of the matrix function. Once again, the linear scaling behavior is clear.

For Table 8.3 we use a pseudospars matrix  $A$  with entries given by  $a_{ij} = e^{-\alpha|i-j|}$ , where  $\alpha > 0$ . This is a convenient way to generate matrices with a given rate of decay. Note that in the limiting case  $\alpha = 0$ , the matrix is “flat” (there is no decay). As  $\alpha$  is increased, the rate of decay also increases. It is known that  $A^{-1}$  is a tridiagonal  $M$ -matrix, hence  $A$  is positive definite. To compute  $f(A) = \log(A)$ , we first sparsify  $A$  by keeping enough diagonals to make the sparsification error  $\|A - \tilde{A}\|$  smaller than a

TABLE 8.2

	$A \log(A)$	$\text{trace}[A \log(A)]$		
$n$	error	error	$\hat{m}$	$N$
100	$5e-07$	$3e-04$	20	9
200	$6e-07$	$8e-04$	20	9
300	$1e-07$	$3e-04$	20	10
500	$2e-07$	$5e-04$	20	10

prescribed accuracy and then we approximate  $\log(\tilde{A})$ . The reported errors are relative to the matrix function  $\log(A)$  computed on the full  $A$ . For the first test we use  $\alpha = 2$  and we keep  $m = 15$  off-diagonals above and below the main diagonal. Again, the linear scaling behavior is apparent.

TABLE 8.3

	$\log(A)$	$\text{trace}[\log(A)]$		
$n$	error	error	$\hat{m}$	$N$
100	$4e-07$	$7e-06$	10	9
200	$4e-07$	$1e-05$	10	9
300	$4e-07$	$2e-05$	10	9
500	$4e-07$	$3e-05$	10	9

Next we consider some non-Hermitian examples. In Table 8.4 we show results for the exponential, the sine and the cosine. The matrix  $A$  has entries  $a_{ij} = e^{-\alpha(i-j)}$  for  $i \geq j$  and  $a_{ij} = e^{-\beta(j-i)}$  for  $i < j$  with  $\alpha, \beta > 0$ . To obtain a non-symmetric matrix we choose  $\alpha = 1$  and  $\beta = 1.5$ . We sparsify the matrix by keeping 25 off-diagonals on either side of the main diagonal. The reported errors are relative to the “exact” computation using the full  $A$ .

TABLE 8.4

	$\exp(A)$			$\cos(A)$			$\sin(A)$		
$n$	error	$\hat{m}$	$N$	error	$\hat{m}$	$N$	error	$\hat{m}$	$N$
100	$6e-08$	30	12	$4e-07$	30	11	$9e-07$	30	11
300	$6e-08$	30	12	$4e-07$	30	11	$3e-08$	30	12
500	$4e-08$	30	11	$2e-08$	30	11	$2e-07$	30	10

In Table 8.5 we show results for the exponential function for a non-symmetric matrix with  $A$  generated as above, with  $\alpha = 1$  and  $\beta = 2$ . Now we keep only 15 off-diagonals on either side of the main diagonal. Again, we observe  $n$ -independence of the results.

TABLE 8.5

$\exp(A)$			
$n$	error	$\hat{m}$	$N$
100	$3e-07$	25	10
300	$4e-07$	25	9
500	$4e-07$	25	9
1000	$4e-07$	25	9

For the results in Table 8.6 we use two examples of banded non-symmetric matrices from the Matrix Market. Again, we show results for the exponential, the sine and the cosine. The matrices were scaled by dividing through by the largest eigenvalue.

TABLE 8.6

$f(A)$	$A$	$n$	error	$\hat{m}$	$N$
$\exp(A)$	<i>Bai/olm100</i>	100	$3e-06$	12	8
$\exp(A)$	<i>Bai/olm500</i>	500	$2e-06$	12	9
$\sin(A)$	<i>Bai/olm100</i>	100	$3e-06$	12	8
$\sin(A)$	<i>Bai/olm500</i>	500	$1e-06$	12	9
$\cos(A)$	<i>Bai/olm100</i>	100	$4e-06$	12	8
$\cos(A)$	<i>Bai/olm500</i>	500	$2e-06$	12	9

In Table 8.7 and Table 8.8 we give results for the approximation of the determinant of a symmetric positive definite matrix  $A$  using the identity  $\det(A) = \exp(\text{trace}[\log(A)])$ . For Table 8.7 we use the matrix  $A$  with entries  $a_{ij} = e^{-2|i-j|}$  as above and truncate it symmetrically to 15 off-diagonals. For Table 8.8 we use the same matrix used in Table 8.2. In both tables we report relative errors.

TABLE 8.7  
*Determinant, I*

$n$	error	$\hat{m}$	$N$
100	$7e-06$	10	9
200	$1e-05$	10	9
300	$2e-05$	10	9
500	$4e-05$	10	9

TABLE 8.8  
*Determinant, II*

$n$	error	$\hat{m}$	$N$
100	$4e-05$	20	11
200	$1e-04$	20	11
300	$2e-04$	20	11
400	$3e-04$	20	11

Finally, we consider banded approximations to  $f(A) = A^{-1}$  as preconditioners for the conjugate gradient method applied to linear systems  $Ax = b$  where  $A = T + D$  is a symmetric positive definite “Toeplitz-plus-diagonal” matrix. We note that no fast solvers are available for this problem unless  $D$  is nearly a multiple of the identity, but see the recent paper [83] for an approach that yields reasonable results. Here  $T$  has entries  $t_{ij} = e^{-0.1|i-j|}$  and  $D$  is a random diagonal matrix with entries in the interval  $[5, 6]$ . The right-hand side  $b$  is a random vector and  $x_0 = 0$  is used as the initial guess. We stop the preconditioned conjugate gradient (PCG) iteration when the residual norm is reduced to at least  $1e-07$ . The preconditioners are constructed as follows. We first truncate  $A$  to obtain a matrix with bandwidth  $m = 20$ , then we run a fixed number  $N = 10$  of iterations of the polynomial approximation method for  $f(z) = z^{-1}$  to construct a banded approximate inverse to the truncated matrix. We

fix the bandwidth of the approximate inverse to  $\hat{m} = 40$ , i.e., we allow the approximate inverse to have twice the bandwidth of the truncated matrix. Of course, the original matrix  $A$  is used for the matrix–vector multiplies in the PCG iteration. The results in Table 8.9 show that this simple preconditioner works quite well. Note that with an FFT-based implementation the cost of each matrix–vector multiply with  $A$  in the conjugate gradient method is  $O(n \log n)$ , whereas the cost of forming and applying the preconditioner in each iteration is just  $O(n)$ . In contrast, application of a circulant-plus-diagonal preconditioner costs  $O(n \log n)$  in general.

TABLE 8.9  
Number of PCG iterations

Precond.	$n$						
	100	200	600	800	1000	2000	3000
None	37	56	73	82	85	86	95
$M \approx A^{-1}$	8	12	13	13	14	14	16

**9. Conclusions and future work.** In this paper we have proved some general theoretical results on decay phenomena in functions of sparse matrices, and we have investigated some algorithms, based on polynomial approximation, for approximating matrix functions. Our decay bounds extend and unify several results that were known from the literature, and help understand various localization phenomena observed by physicists. Moreover, our results show that  $O(n)$  approximation is possible in many cases of practical interest. One of the goals of the paper is to call the attention of numerical analysts to this class of problems, which is ripe for treatment by experts in matrix computations. Of course, much work remains to be done. Here we elaborate on some possible directions of future research.

(1) The decay bounds and approximation algorithms for  $f(A)$  discussed in this paper do not make use of possible structural properties of  $A$ , except for bandedness or sparsity. However, the matrices that arise in applications are often endowed with additional structure. For example,  $A$  may have Toeplitz or block Toeplitz structure, or belong to the class of Hamiltonian or skew-Hamiltonian matrices, and so forth [32]. More generally,  $A$  may belong to a Lie or Jordan algebra corresponding to a matrix group (see [7]), and  $f$  may be a mapping from the algebra to the group. The best known examples are the exponential and the Cayley transform, which map skew-symmetric matrices to orthogonal ones. Similar relationships link (skew-)Hamiltonian and symplectic matrices, and pseudo-Hermitians with pseudo-unitary ones. Exploitation of this rich mathematical structure, when present, can be expected to lead to improved theoretical results and numerical algorithms. This could be especially useful for problems in quantum computing, where the matrices of interest usually belong to the classical matrix groups; see, e.g., [26, 91].

(2) There are important examples of situations where the decay behavior of  $f(A)$  is too slow to result in efficient approximations, for instance if the decay is algebraic (with small exponent  $p$ ) rather than exponential. In the area of electronic structure calculations, some authors have advocated the use of wavelet bases of various types, resulting in much faster off-diagonal decay of the entries of the density matrix than with the standard representation; see, e.g., [16, 49, 90]. As long as the entries of  $f(A)$  vary smoothly, a wavelet representation in which  $f(A)$  becomes extremely sparse is in principle possible. Similar wavelet compression ideas have been used by numerical analysts in many other situations, for instance, to improve the performance and

quality of sparse approximate inverse preconditioners: see [20, 25, 56]. It would be important to investigate the use of wavelet transforms and other tools from multiresolution analysis to improve the performance of approximation algorithms for matrix functions. For the case of functions of elliptic operators we also mention the approach in [46], where a different type of data compression is achieved by making use of the theory of hierarchical matrices.

(3) Finally, in this paper we have illustrated the theory by some simple MATLAB experiments involving mostly band matrices of fairly small size. Although this was enough for our purpose, which was to illustrate the decay behavior of  $f(A)$  and the  $O(n)$  scaling of the approximation algorithms, we plan to investigate more realistic problems in the near future. This includes larger matrices with general sparsity structure, and more complicated matrix functions. Efficient implementation of linear scaling algorithms for matrices with general sparsity patterns is going to be considerably more complicated than simpler banded approaches. Using bandwidth-reducing orderings (like reverse Cuthill–McKee, or more current ones) may be helpful in some cases, but by no means always; also, treating all entries within the band as nonzero would be wasteful. The main challenge is to develop efficient sparse matrix–matrix multiplies with adaptive sparsity patterns to incorporate dropping. This problem, which is of course important also in other contexts, is still without a completely satisfactory solution. We plan to investigate this further, building on the work in [24, 53].

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