# Closer to the solution: Iterative linear solvers

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## 1 Introduction

The solution of dense linear systems received much attention after the second world war, and by the end of the sixties, most of the problems associated with it had been solved. For a long time, Wilkinson's "The Algebraic Eigenvalue Problem" [107], other than the title suggests, became also the standard textbook for the solution of linear systems. When it became clear that partial differential equations could be solved numerically, to a level of accuracy that was of interest for application areas (such as reservoir engineering, and reactor diffusion modeling), there was a strong need for the fast solution of the discretized systems, and iterative methods became popular for these problems.

Although, roughly speaking, the successive overrelaxation methods and the first Krylov subspace methods were developed in the same period of time, the class of Successive Overrelaxation methods became the methods of choice, since they required a small amount of computer storage. These methods were quite well understood by the work of Young [108], and the theory was covered in great detail in Varga's book [102]. This book was the source of reference for iterative methods for a long time.

It may seem evident that iterative methods gained in importance as scientific modeling led to larger linear problems, since direct methods are often too expensive in terms of computer memory and CPU-time requirements. However, the convergence behavior of iterative methods depends very much on (often *a priori* unknown) properties of the linear system, which means that one has to consider the origin of the problem. Since large linear systems may arise from various sources, for instance, data fitting problems, discretization of PDE's, Markov chains, or tomography, these properties may also vary widely. This means that one has to study the usefulness of iterative schemes for each class of problems separately and one can hardly rely on expertise gained from other classes. Most often preconditioning is required to obtain or improve convergence. Only for some classes of problems the effects of preconditioning are well understood; for

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most problems it remains largely a matter of trial and error. In our final section we will give some hints on preconditioning as well as pointers to literature. Also the (sparsity) structure of the matrix is important for the decision between algorithms: special algorithms can often be designed for special structures, e.g., Toeplitz matrices, banded matrices, red-black ordering, p-cyclic matrices, etc. Some important matrix properties can be easily deduced from the originating problem, for instance, symmetry and positive definiteness. Iterative methods that rely on these properties, such as the Lanczos method and the conjugate gradient method, have therefore the most widespread use.

The conjugate gradient method, and the method of Lanczos, were first viewed as exact projection methods, and since their behavior in finite precision arithmetic was not in agreement with that point of view, they were initially considered with suspicion. Through work of Reid [79], it became clear that these methods, if used as iterative techniques, could be used to advantage for many classes of linear systems. With preconditioning [21], these methods could be made even more efficient. Among the first popular preconditioned methods was ICCG [69, 63].

In the period after 1975, we have seen that symmetric positive definite sparse systems were usually solved by preconditioned CG methods, when *very* large, and by sparse direct solvers if moderately large. For indefinite symmetric sparse systems special variants were proposed, like MINRES, and SYMMLQ [74]. However, for very large unsymmetric systems, the SOR methods, and the method of Chebyshev [102, 54, 53], nicely tuned by Manteuffel [68], were still the methods of choice. Due to several poorly understood numerical problems, the two-sided Lanczos method, and a special variant Bi-CG [41], were not so popular at that time, but slowly more robust variants of Krylov subspace methods, with longer recursion formulas, entered the field. We mention GENCG [32], FOM [84], OR-THOMIN [103], ORTHODIR and ORTHORES [62].

In the mid-eighties we see the start of the popularity of Krylov subspace methods for large unsymmetric systems, and many new powerful and robust methods were proposed. The past 10 years have formed a very lively area of research for these methods, but we believe that by now this has come more or less to an end. We are now facing other challenging problems for the solution of very large sparse problems. For large linear systems, associated with 3-dimensional modeling, the iterative methods are often the only means we have for solution.

Of course, for structured and rather regular problems the multigrid methods have usually a better rate of convergence, but even from the CFD community we see an increasing attention to iterative schemes. We believe that it is not fruitful to see the two classes of methods as competitive, they may be combined, where depending on the point of view, multigrid is used as a preconditioner, or the iterative scheme as a smoother. We will not further discuss multigrid in our paper; it deserves separate attention. An excellent introduction to multigrid and the related multilevel methods, in the context of solving linear systems stemming from certain classes of problems, can be found in [61]

In the next section we will highlight the Krylov subspace methods as the source of one of the most important developments in the period 1985-1995. In subsequent sections, we will discuss a number of new variants.

For ease of presentation we restrict ourselves to real-valued problems. Matrices will be denoted by capitals (A, B, ...), vectors are represented in lower case (x, y, ...), and scalars are represented by Greek symbols  $(\alpha, \Theta, ...)$ . Unless stated differently, matrices will be of order  $n: A \in \mathbb{R}^{n \times n}$ , and vectors will have length n.

## 2 Krylov subspace methods: the basics

The past ten years have led to well-established and popular methods, that all lead to the construction of approximate solutions in the so-called Krylov subspace. Given a linear system Ax = b, with a large, usually sparse, unsymmetric nonsingular matrix A, then the standard Richardson iteration

$$x_k = (I - A)x_{k-1} + b$$

generates approximate solutions in shifted Krylov subspaces

$$x_0 + K^k(A; r_0) = x_0 + \{r_0, Ar_0, \dots, A^{k-1}r_0\},\$$

with  $r_0 = b - Ax_0$ , for some given initial vector  $x_0$ . The Krylov subspace projection methods fall in three different classes:

- 1. The *Ritz-Galerkin approach*: Construct the  $x_k$  for which the residual is orthogonal to the current subspace:  $b Ax_k \perp K^k(A; r_0)$ .
- 2. The minimum residual approach: Identify the  $x_k$  for which the Euclidean norm  $||b Ax_k||_2$  is minimal over  $K^k(A; r_0)$ .
- 3. The *Petrov-Galerkin approach*: Find an  $x_k$  so that the residual  $b Ax_k$  is orthogonal to some other suitable k-dimensional subspace.

The Ritz-Galerkin approach leads to such popular and well-known methods as Conjugate Gradients, the Lanczos method, FOM, and GENCG. The minimum residual approach leads to methods like GMRES, MINRES, and ORTHODIR. If we select the k-dimensional subspace in the third approach as  $K^k(A^T; s_0)$ , then we obtain the Bi-CG, and QMR methods. More recently, hybrids of the three approaches have been proposed, like CGS, Bi-CGSTAB, BiCGSTAB( $\ell$ ), TFQMR, FGMRES, and GMRESR.

Most of these methods have been proposed in the last ten years. GMRES, proposed in 1986 by Saad and Schultz [84], is the most robust of them, but, in terms of work per iteration step it is also the most expensive. Bi-CG, which

was suggested by Fletcher in 1977 [41], is a relatively inexpensive alternative, but it has problems with respect to convergence: the so-called breakdown situations. This aspect has received much attention in the past period. Parlett et al [75] introduced the notion of look-ahead, in order to overcome breakdowns, and this was further perfected by Freund and Nachtigal [45], and by Brezinski and Redivio-Zaglia [12]. The theory for this look-ahead technique was linked to the theory of Padé approximations by Gutknecht [59]. Other contributions to overcome specific breakdown situations were made by Bank and Chan [9], and Fischer [39]. We will discuss these approaches in our section on QMR.

The hybrids were developed primarily in the second half of the last 10 years; the first of these was CGS, published in 1989 by Sonneveld [89], and followed by Bi-CGSTAB, by van der Vorst in 1992 [98], and others. The hybrid variants of GMRES: Flexible GMRES and GMRESR, in which GMRES is combined with some other iteration scheme, have only been proposed very recently.

A nice overview of Krylov subspace methods, with focus on Lanczos-based methods, is given in [44]. Simple algorithms and unsophisticated software for some of these methods is provided in [10]. Iterative methods with much attention to various forms of preconditioning have been described in [5]. Recently a book on iterative methods was published also by Saad [83]; it is very algorithm oriented, with, of course, a focus on GMRES and preconditioning techniques, like threshold ILU, ILU with pivoting, and incomplete LQ factorizations. An annotated entrance to the vast literature on preconditioned iterative methods

## is given in [14].

#### 2.1 Working with the Krylov subspace

In order to identify optimal approximate solutions in the Krylov subspace we need a suitable basis for this subspace, one that can be extended in a meaningful way for subspaces of increasing dimension. The obvious basis  $r_0$ ,  $Ar_0$ , ...,  $A^{i-1}r_0$ , for  $K^i(A; r_0)$ , is not very attractive from a numerical point of view, since the vectors  $A^j r_0$  point more and more in the direction of the dominant eigenvector for increasing j (the power method!), and hence the basis vectors become dependent in finite precision arithmetic.

Instead of the standard basis one usually prefers an orthonormal basis, and Arnoldi [1] suggested computing this basis as follows. Start with  $v_1 \equiv r_0/||r_0||_2$ . Assume that we have already an orthonormal basis  $v_1, \ldots, v_j$  for  $K^j(A; r_0)$ , then this basis is expanded by computing  $t = Av_j$ , and by orthonormalizing this vector t with respect to  $v_1, \ldots, v_j$ . In principle the orthonormalization process can be carried out in different ways, but the most commonly used approach is to do this by a modified Gram-Schmidt procedure [53]. This leads to an algorithm for the creation of an orthonormal basis for  $K^m(A; r_0)$ , as in Fig 1.

It is easily verified that  $v_1, \ldots, v_m$  form an orthonormal basis for  $K^m(A; r_0)$ (that is, if the construction does not terminate at a vector t = 0). The or-

$$\begin{split} v_1 &= r_0 / \|r_0\|_2; \\ \text{for } j &= 1, ..., m-1 \\ t &= A v_j; \\ \text{for } i &= 1, ..., j \\ h_{i,j} &= v_i^T t; \\ t &= t-h_{i,j} v_i; \\ \text{end}; \\ h_{j+1,j} &= \|t\|_2; \\ v_{j+1} &= t/h_{j+1,j}; \\ \text{end} \end{split}$$

Figure 1. Arnoldi's method with modified Gram-Schmidt orthogonalization

tho gonalization leads to relations between the  $v_j$ , that can be formulated in a compact algebraic form. Let  $V_j$  denote the matrix with columns  $v_1$  up to  $v_j$ , then it follows that

$$AV_{m-1} = V_m H_{m,m-1}$$

The m by m-1 matrix  $H_{m,m-1}$  is upper Hessenberg, and its elements  $h_{i,j}$  are defined by the Arnoldi algorithm.

From a computational point of view, this construction is composed from three basic elements: a matrix vector product with A, innerproducts, and updates. We see that this orthogonalization becomes increasingly expensive for increasing dimension of the subspace, since the computation of each  $h_{i,j}$  requires an inner product and a vector update.

Note that if A is symmetric, then so is  $H_{m-1,m-1} = V_{m-1}^T A V_{m-1}$ , so that in this situation  $H_{m-1,m-1}$  is tridiagonal. This means that in the orthogonalization process, each new vector has to be orthogonalized with respect to the previous two vectors only, since all other innerproducts vanish. The resulting three term recurrence relation for the basis vectors of  $K_m(A; r_0)$  is known as the Lanczos method and some very elegant methods are derived from it. In the symmetric case the orthogonalization process involves constant arithmetical costs per iteration step: one matrix vector product, two innerproducts, and two vector updates.

## 3 The Ritz-Galerkin approach: FOM and CG

Let us assume, for simplicity, that  $x_0 = 0$ , and hence  $r_0 = b$ . The Ritz-Galerkin conditions imply that  $r_k \perp K^k(A; r_0)$ , and this is equivalent to

$$V_k^T(b - Ax_k) = 0.$$

Since  $b = r_0 = ||r_0||_2 v_1$ , it follows that  $V_k^T b = ||r_0||_2 e_1$  with  $e_1$  the first canonical unit vector in  $\mathbb{R}^k$ . With  $x_k = V_k y$  we obtain

$$V_k^T A V_k y = \|r_0\|_2 e_1.$$

This system can be interpreted as the system Ax = b projected onto  $K^k(A; r_0)$ .

Obviously we have to construct the  $k \times k$  matrix  $V_k^T A V_k$ , but this is, as we have seen, readily available from the orthogonalization process:

$$V_k^T A V_k = H_{k,k}$$

so that the  $x_k$  for which  $r_k \perp K^k(A; r_0)$  can be easily computed by first solving  $H_{k,k}y = ||r_0||_2 e_1$ , and forming  $x_k = V_k y$ . This algorithm is known as FOM or GENCG.

Note that for some  $j \leq n-1$  the construction of the orthogonal basis must terminate. In that case we have that  $AV_{j+1} = V_{j+1}H_{j+1,j+1}$ . Let y be the solution of the reduced system  $H_{j+1,j+1}y = ||r_0||e_1$ , and  $x_{j+1} = V_{j+1}y$ . Then it follows that  $x_{j+1} = x$ , i.e., we have arrived at the exact solution, since

$$Ax_{j+1} - b = AV_{j+1}y - b = V_{j+1}H_{j+1,j+1}y - b$$
  
=  $||r_0||V_{j+1}e_1 - b = 0$ 

(we have assumed that  $x_0 = 0$ ).

When A is symmetric, then  $H_{k,k}$  reduces to a tridiagonal matrix  $T_{k,k}$ , and the resulting method is known as the Lanczos method [65]. In clever implementations, it is possible to avoid storing all the vectors  $v_j$ . When A is in addition positive definite then we obtain, at least formally, the Conjugate Gradient method. In commonly used implementations of this method, one implicitly forms an LUfactorization for  $T_{k,k}$  and this leads to very elegant short recurrencies for the  $x_j$ and the corresponding  $r_j$ . The positive definiteness is necessary to guarantee the existence of the LU factorization, but it allows also for another useful interpretation. From the fact that  $r_i \perp K^i(A; r_0)$ , it follows that  $A(x_i - x) \perp K^i(A; r_0)$ , or  $x_i - x \perp_A K^i(A; r_0)$ . The latter observation expresses the fact that the error is A-orthogonal to the Krylov subspace and this is equivalent to the important observation that  $||x_i - x||_A$  is minimal<sup>1</sup> For an overview of the history of CG and main contributions on this subject, see [51].

<sup>&</sup>lt;sup>1</sup> The A-norm is defined by  $||y||_A^2 = (y, y)_A \equiv (y, Ay)$ , and we need the positive definiteness of A in order to get a proper innerproduct  $(\cdot, \cdot)_A$ .

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The local convergence behavior of CG, and especially the occurrence of superlinear convergence, was first explained in a qualitative sense in [21], and later in a quantitative sense in [94]. In both papers it was linked to the convergence of eigenvalues (Ritz values) of  $T_{i,i}$  towards eigenvalues of A, for increasing i. The global convergence can be bounded with expressions that involve condition numbers, for details see for instance [21, 53, 2]. In [2] the situation is analysed where the eigenvalues of A are in disjunct intervals.

## 4 The minimum residual approach

The creation of an orthogonal basis for the Krylov subspace, with basis vectors  $v_1, \ldots, v_{i+1}$ , leads to

$$AV_i = V_{i+1}H_{i+1,i}, (4.1)$$

where  $V_i$  is the matrix with columns  $v_1$  to  $v_i$ . We look for an  $x_i \in K^i(A; r_0)$ , that is  $x_i = V_i y$ , for which  $||b - Ax_i||_2$  is minimal. This norm can be rewritten as

$$||b - Ax_i||_2 = ||b - AV_iy||_2 = || ||r_0||_2 V_{i+1}e_1 - V_{i+1}H_{i+1,i}y||_2.$$

Now we exploit the fact that  $V_{i+1}$  is an orthonormal transformation with respect to the Krylov subspace  $K^{i+1}(A; r_0)$ :

$$||b - Ax_i||_2 = ||||r_0||_2 e_1 - H_{i+1,i}y||_2$$

and this final norm can simply be minimized by solving the minimum norm least squares problem for the i + 1 by i matrix  $H_{i+1,i}$  and right-hand side  $||r_0||_2 e_1$ .

In GMRES [84] this is done efficiently with Givens rotations, that annihilate the subdiagonal elements in the upper Hessenberg matrix  $H_{i+1,i}$ .

Note that when A is Hermitian (but not necessarily positive definite), the upper Hessenberg matrix  $H_{i+1,i}$  reduces to a tridiagional system. This simplified structure can be exploited in order to avoid storage of all the basis vectors for the Krylov subspace, in a way similar as has been pointed out for CG. The resulting method is known as MINRES [74].

In order to avoid excessive storage requirements and computational costs for the orthogonalization, GMRES is usually restarted after each m iteration steps. This algorithm is referred to as GMRES(m); the not-restarted version is often called 'full' GMRES. There is no simple rule to determine a suitable value for m; the speed of convergence may vary drastically for nearby values of m.

There is an interesting and simple relation between the Ritz-Galerkin approach (FOM and CG) and the minimum residual approach (GMRES and MIN-RES). In GMRES the projected system matrix  $H_{i+1,i}$  is transformed by Givens rotations to an upper triangular matrix (with last row equal to zero). So, in fact, the major difference between FOM and GMRES is that in FOM the last ((*i*+1)-th row is simply discarded, while in GMRES this row is rotated to a zero vector. Let us characterize the Givens rotation, acting on rows i and i + 1, in order to zero the element in position (i + 1, i), by the sine  $s_i$  and the cosine  $c_i$ . Let us further denote the residuals for FOM with an superscript F and those for GMRES with superscript G. Then we have the following relation between FOM and GMRES:

If  $c_k \neq 0$  then the FOM and the GMRES residuals are related by

$$||r_k^F||_2 = \frac{||r_k^G||_2}{\sqrt{1 - (||r_k^G||_2 / ||r_{k-1}^G||_2)^2}},$$
(4.2)

([23]: theorem 3.1). From this relation we see that when GMRES has a significant reduction at step k, in the norm of the residual (i.e.,  $s_k$  is small, and  $c_k \approx 1$ ), then FOM gives about the same result as GMRES. On the other hand when FOM has a breakdown ( $c_k = 0$ ), then GMRES does not lead to an improvement in the same iteration step. Because of these relations we can link the convergence behaviour of GMRES with the convergence of Ritz values (the eigenvalues of the "FOM" part of the upper Hessenberg matrix). This has been exploited in [100], for the analysis and explanation of local effects in the convergence behaviour of GMRES.

There are various different implementations of FOM and GMRES. Among those equivalent to GMRES are: Orthomin [103], Orthodir [62], GENCR [33], and Axelsson's method [3]. These methods are often more expensive than GM-RES per iteration step. Orthomin continues to be popular, since this variant can be easily truncated (Orthomin(s)), in contrast to GMRES. The truncated and restarted versions of these algorithms are not necessarily mathematically equivalent.

Methods that are mathematically equivalent to FOM are: Orthores [62] and GENCG [19, 106]. In these methods the approximate solutions are constructed such that they lead to orthogonal residuals (which form a basis for the Krylov subspace; analogously to the CG method). A good overview of all these methods and their relations is given in [83].

The GMRES method and FOM are closely related to vector extrapolation methods, when the latter are applied to linearly generated vector sequences. For a discussion on this, as well as for implementations for these matrix free methods, see [85].

#### 4.1 Inner-outer iteration schemes

Although GMRES is optimal, in the sense that it leads to a minimal residual solution over the Krylov subspace, it has the disadvantage that this goes with increasing computational costs per iteration step. Of course, with a suitable preconditioner one can hope to reduce the dimension of the required Krylov subspace to acceptable values, but in many cases such preconditioners have not been identified. The preconditioner  $K^{-1}$  is generally viewed as an approximation for the inverse of the matrix A of the system Ax = b to be solved. Instead of an optimal update direction  $A^{-1}r$ , we compute the direction  $p = K^{-1}r$ . That is, we solve p from Kp = r, and K is constructed in such a way that this is easy to do. An attractive idea is to try to get better approximations for  $A^{-1}r$ , and there are two related approaches to accomplish this. One is to try to improve the preconditioner with updates from the Krylov subspace. This has been suggested first by Eirola and Nevanlinna [31]. Their approach leads to iterative methods that are related to Broyden's method [13], which is a Newton type of method. The Broyden method can be obtained from this update-approach if we do not restrict ourselves to Krylov subspaces. See [104] for a discussion on the relation of these methods.

The updated preconditioners can not be applied immediately to GMRES, since the preconditioned operator now changes from step to step, and we are not forming a regular Krylov subspace, but rather a subset of a higher dimensional Krylov subspace. However, we can still minimize the residual over this subset.

The idea of variable preconditioning has been exploited in this sense, by different authors. Axelsson and Vassilevski [7] have proposed a Generalized Conjugate Gradient method with variable preconditioning, Saad [81, 83] has proposed a scheme very similar to GMRES, called Flexible GMRES (FGMRES), and Van der Vorst and Vuik have published a GMRESR scheme. FGMRES has received most attention, presumably because it is so easy to implement: only the update directions in GMRES have to be preconditioned, and each update may be preconditioned differently. This means that only one line in the GMRES algorithm has to be adapted. The price to be paid is that the method is no longer robust; it may break down. The GENCG and GMRESR schemes are slightly more expensive in terms of memory requirements and in computational overhead per iteration step. The main difference between the two schemes is that GENCG in [7] works with Gram-Schmidt orthogonalization, whereas GMRESR makes it possible to use Modified Gram-Schmidt. This may give GMRESR an advantage in actual computations. In exact arithmetic GMRESR and GENCG should produce the same results for the same preconditioners. Another advantage of GMRESR over Algorithm 1 in [7] is that only one matrix vector product is used per iteration step in GMRESR; GENCG needs two matrix vector products per step.

In GMRESR the residual vectors are preconditioned and if this gives a further reduction then GMRESR does not break down. This gives slightly more control over the method in comparison with FGMRES. In most cases though the results are about the same, but the efficient scheme for FGMRES has an advantage.

We will briefly discuss the GMRESR method. In [101] it has been shown how the GMRES-method, or more precisely, the GCR-method, can be combined with other iterative schemes. The iteration steps of GMRES (or GCR) are called outer iteration steps, while the iteration steps of the preconditioning iterative method are referred to as inner iterations. The combined method is called GMRES\*, where  $\star$  stands for any given iterative scheme; in the case of GMRES as the inner iteration method, the combined scheme is called GMRESR[101]. It was shown in [26, 25], that GMRESR can be implemented in a way that avoids about 30% of the overhead in the outerloop, which makes the method about as expensive per outer iteration step as FGMRES.

The GMRES $\star$  algorithm can be described as in Fig. 2.

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\begin{aligned} x_0 \text{ is an initial guess; } r_0 &= b - Ax_0; \\ \text{for } i &= 0, 1, 2, 3, \dots \\ \text{Let } z^{(m)} \text{ be the approximate solution} \\ \text{of } Az &= r_i, \text{ obtained after } m \text{ steps of} \\ \text{an iterative method.} \\ c &= Az^{(m)} \text{ (often available from the} \\ \text{ iteration method}) \\ \text{for } k &= 0, \dots, i-1 \\ \alpha &= c_k^T c \\ c &= c - \alpha c_k \\ z^{(m)} &= z^{(m)} - \alpha u_k \\ c_i &= c/||c||_2; u_i &= z^{(m)}/||c||_2 \\ x_{i+1} &= x_i + (c_i^T r_i)u_i \\ r_{i+1} &= r_i - (c_i^T r_i)c_i \\ \text{ if } x_{i+1} \text{ is accurate enough then quit} \\ \text{end} \end{aligned}
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Figure 2. The GMRES\* algorithm

A sufficient condition to avoid breakdown in this method  $(||c||_2 = 0)$  is that the norm of the residual at the end of an inner iteration is smaller than the norm of the right-hand side residual:  $||Az^{(m)} - r_i||_2 < ||r_i||_2$ . This can easily be controlled during the inner iteration process. If stagnation occurs, i.e. no progress at all is made in the inner iteration, then it is suggested in [101] to do one (or more) steps of the LSQR method, which guarantees a reduction (but this reduction is often very small).

The idea behind these flexible iteration schemes is that we explore parts of high-dimensional Krylov subspaces, hopefully determining essentially the same approximate solution that full GMRES would find over the entire subspace, but at much lower computational costs. For the inner iteration we may select any appropriate solver, for instance, one cycle of GMRES(m), since then we have also locally an optimal method, or some other iteration scheme, like for instance Bi-CGSTAB.

In [26] it is proposed to keep the Krylov subspace, that is built in the inner iteration, orthogonal with respect to the Krylov basis vectors generated in the outer iteration. Under various circumstances this helps to avoid inspection of already investigated parts of Krylov subspaces in future inner iterations. The procedure works as follows. In the outer iteration process the vectors  $c_0, ..., c_{i-1}$ build an orthogonal basis for the Krylov subspace. Let  $C_i$  be the *n* by *i* matrix with columns  $c_0, ..., c_{i-1}$ . Then the inner iteration process at outer iteration *i* is carried out with the operator  $A_i$  instead of A, and  $A_i$  is defined as

$$A_i = (I - C_i C_i^T) A. (4.3)$$

Of course,  $I - C_i C_i^T$  can be stored as a product of Householder transformations, which makes the updates more efficient. It is easily verified that  $A_i z \perp c_0, ..., c_{i-1}$ for all z, so that the inner iteration process takes place in a subspace orthogonal to these vectors. The additional costs, per iteration of the inner iteration process, are *i* inner products and *i* vector updates. In order to save on these costs, one should realize that it is not necessary to orthogonalize with respect to all previous *c*-vectors, and that "less effective" directions may be dropped, or combined with others. In [72, 8, 26] suggestions are made for such strategies. Of course, these strategies are only attractive in cases where we see too little residual reducing effect in the inner iteration process in comparison with the outer iterations of GMRES\*.

Note that if we carry out the preconditioning by doing a few iterations of some other iteration process, then we have *inner-outer iteration* schemes; these have been discussed earlier in [52]. It is difficult to analyse their convergence behavior and in a first attempt it is wise to look at the analysis for simplified schemes. We will give the flavor of this analysis and the type of results [52].

We consider a simplified inner-outer iteration scheme associated with the standard iteration for the splitting A = M - N, as given in Fig. 3; the inner iterations are carried out with M.

With  $e_k \equiv x - x_k$ , it follows that

$$e_{k+1} = (I - M^{-1}A)e_k - M^{-1}q_k$$

and

$$\begin{aligned} \|e_{k+1}\| &\leq \|I - M^{-1}A\| \|e_k\| + \|M^{-1}q_k\| \\ &\leq (\|I - M^{-1}A\| + \tau_k \|M^{-1}\| \|A\|) \|e_k\| \end{aligned}$$

Hence

$$\frac{\|e_k\|}{\|e_0\|} \le \left(\|I - M^{-1}A\| + \eta\right)^k,$$

**while**  $\frac{||q_k||_2}{||r_k||_2} < \tau_k$  do  $r_k = b - Ax_k$ Solve  $Mz = r_i$  approximately  $\Rightarrow \bar{z}_k$   $(q_k \equiv M\bar{z}_k - r_k)$   $x_{k+1} = x_k + \bar{z}_k$ if  $x_{k+1}$  is accurate enough **then** quit end

Figure 3. A simplified Inner-Outer iteration scheme

with  $\eta \equiv \max \tau_k ||M^{-1}|| ||A||$ , so that the convergence is largely determined by the norm of  $I - M^{-1}A$ , if  $\tau_k$  is small enough. It can be shown that for  $\tau_k = \Theta^k$ , for a fixed  $0 < \Theta < 1$ :

$$\limsup_{k \to \infty} \left( \frac{\|e_k\|}{\|e_0\|} \right)^{\frac{1}{k}} = \|I - M^{-1}A\|,$$

so that essentially the convergence rate for the splitting A = M - N, with exact inversion of M, is restored. A similar analysis can be carried out in relation with Chebyshev acceleration, see [52, 49]. In [49] the question is studied how  $\Theta$  can be chosen so that the *total* amount of work is minimized.

For an application of inner-outer iterations for Stokes problems, see the inexact Uzawa scheme, proposed in [36]; for Navier-Stokes related problems, see [55].

## 5 The Petrov-Galerkin approach: Bi-CG

For unsymmetric systems we can, in general, not reduce the matrix A to a symmetric system in a lower-dimensional subspace, by orthogonal projections. The reason is that we can not create an orthogonal basis for the Krylov subspace by a 3-term recurrence relation [38]. We can, however, try to obtain a suitable non-orthogonal basis with a 3-term recurrence, by requiring that this basis is orthogonal with respect to some other basis.

We start by constructing an arbitrary basis for the Krylov subspace:

$$h_{i+1,i}v_{i+1} = Av_i - \sum_{j=1}^{i} h_{j,i}v_j, \qquad (5.1)$$

which can be rewritten in matrix notation as  $AV_i = V_{i+1}H_{i+1,i}$ . The coefficients  $h_{i+1,i}$  define the norm of  $v_{i+1}$ , and a natural choice would be to select them such that  $||v_{i+1}||_2 = 1$ . In Bi-CG implementations, a popular choice is to select  $h_{i+1,i}$  such that  $||v_{i+1}||_2 = ||r_{i+1}||_2$ .

Clearly, we can not use  $V_i$  for the projection, but suppose we have a  $W_i$  for which  $W_i^T V_i = D_i$  (an *i* by *i* diagonal matrix with diagonal entries  $d_i$ ), and for which  $W_i^T v_{i+1} = 0$ . Then

$$W_i^T A V_i = D_i H_{i,i}, (5.2)$$

and now our goal is to find a  $W_i$  for which  $H_{i,i}$  is tridiagonal. This means that  $V_i^T A^T W_i$  should be tridiagonal too. This last expression has a similar structure as the right-hand side in (5.2), with only  $W_i$  and  $V_i$  reversed. This suggests to generate the  $w_i$  with  $A^T$ .

We start with an arbitrary  $w_1 \neq 0$ , such that  $w_1^T v_1 \neq 0$ . Then we generate  $v_2$  with (5.1), and orthogonalize it with respect to  $w_1$ , which means that  $h_{1,1} = w_1^T A v_1 / (w_1^T v_1)$ . Since  $w_1^T A v_1 = (A^T w_1)^T v_1$ , this implies that  $w_2$ , generated with

$$h_{2,1}w_2 = A^T w_1 - h_{1,1}w_1,$$

is also orthogonal to  $v_1$ .

This can be continued, and we see that we can create bi-orthogonal basis sets  $\{v_j\}$ , and  $\{w_j\}$ , by making the new  $v_i$  orthogonal to  $w_1$  up to  $w_{i-1}$ , and then by generating  $w_i$  with the same recurrence coefficients, but with  $A^T$  instead of A. Now we have that  $W_i^T A V_i = D_i H_{i,i}$ , and also that  $V_i^T A^T W_i = D_i H_{i,i}$ . This implies that  $D_i H_{i,i}$  is symmetric, and hence  $H_{i,i}$  is a tridiagonal matrix, which gives us the desired 3-term recurrence relation for the  $v_j$ 's, and the  $w_j$ 's. Note that  $v_1, \ldots, v_i$  form a basis for  $K^i(A; v_1)$ , and  $w_1, \ldots, w_i$  form a basis for  $K^i(A^T; w_1)$ .

We may proceed in a similar way as in the symmetric case:

$$AV_i = V_{i+1}T_{i+1,i}, (5.3)$$

but here we use the matrix  $W_i = [w_1, w_2, ..., w_i]$  for the projection of the system

$$W_i^T(b - Ax_i) = 0$$

or

$$W_i^T A V_i y - W_i^T b = 0.$$

Using (5.3), we find that  $y_i$  satisfies

$$T_{i,i}y = \|r_0\|_2 e_1,$$

and  $x_i = V_i y$ . The resulting method is known as the Bi-Lanczos method [65]

We have assumed that  $d_i \neq 0$ , that is  $w_i^T v_i \neq 0$ . The generation of the biorthogonal basis breaks down if for some *i* the value of  $w_i^T v_i = 0$ , this is referred to in literature as a serious breakdown. Likewise, when  $w_i^T v_i \approx 0$ , we have a nearbreakdown. The way to get around this difficulty is the so-called Look-ahead strategy, which comes down to taking a number of successive basis vectors for the Krylov subspace together and to make them blockwise bi-orthogonal. This has been worked out in detail in [75, 45, 46, 47].

Another way to avoid breakdown is to restart as soon as a diagonal element gets small. Of course, this strategy looks surprisingly simple, but one should realise that at a restart the Krylov subspace, that has been built up so far, is thrown away, and this destroys the possibility of faster (i.e., superlinear) convergence.

We can try to construct an LU-decomposition, without pivoting, of  $T_{i,i}$ . If this decomposition exists, then, similar to CG, it can be updated from iteration to iteration and this leads to a recursive update of the solution vector, which avoids saving all intermediate r and w vectors. This variant of Bi-Lanczos is usually called Bi-Conjugate Gradients, or for short Bi-CG [41]. In Bi-CG, the  $d_i$  are chosen such that  $v_i = r_{i-1}$ , similarly to CG.

Of course one can in general not be certain that an LU decomposition (without pivoting) of the tridiagonal matrix  $T_{i,i}$  exists, and this may lead also to breakdown (a breakdown of the *second kind*), of the Bi-CG algorithm. Note that this breakdown can be avoided in the Bi-Lanczos formulation of the iterative solution scheme, e.g., by making an LU-decomposition with 2 by 2 block diagonal elements [9]. It is also avoided in the QMR approach (see Section 5.1).

Note that for symmetric matrices Bi-Lanczos generates the same solution as Lanczos, provided that  $w_1 = r_0$ , and under the same condition Bi-CG delivers the same iterands as CG for positive definite matrices. However, the Bi-orthogonal variants do so at the cost of two matrix vector operations per iteration step. For a computational scheme for Bi-CG, without provisions for breakdown, see [10].

#### 5.1 QMR

The QMR method [47] relates to Bi-CG in a similar way as MINRES relates to CG. We start with the recurrence relations for the  $v_j$ :

$$AV_i = V_{i+1}T_{i+1,i}$$

We would like to identify the  $x_i$ , with  $x_i \in K^i(A; r_0)$ , or  $x_i = V_i y$ , for which

$$||b - Ax_i||_2 = ||b - AV_iy||_2 = ||b - V_{i+1}T_{i+1,i}y||_2$$

is minimal, but the problem is that  $V_{i+1}$  is not orthogonal. However, we pretend that the columns of  $V_{i+1}$  are orthogonal. Then

$$\|b - Ax_i\|_2 = \|V_{i+1}(\|r_0\|_2 e_1 - T_{i+1,i}y)\|_2 = \|(\|r_0\|_2 e_1 - T_{i+1,i}y)\|_2$$

and in [47] it is suggested to solve the projected miniminum norm least squares problem  $\|(\|r_0\|_2 e_1 - T_{i+1,i}y)\|_2$ . The minimum value of this norm is called the quasi residual and will be denoted by  $\||r_i^Q\|_2$ .

Since, in general, the columns of  $V_{i+1}$  are not orthogonal, the computed  $x_i = V_i y$  does not solve the minimum residual problem, and therefore this approach is referred to as a Quasi-minimum residual approach [47]. It can be shown that the norm of the residual  $r_i^{QMR}$  of QMR can be be bounded in terms of the quasi residual

$$|r_i^{QMR}||_2 \le \sqrt{i+1} ||r_i^Q||_2.$$

The sketched approach leads to the simplest form of the QMR method. A more general form arises if the least squares problem is replaced by a weighted least squares problem [47]. No strategies are yet known for optimal weights.

In [47] the QMR method is carried out on top of a look-ahead variant of the biorthogonal Lanczos method, which makes the method more robust. Experiments indicate that although QMR has a much smoother convergence behaviour than Bi-CG, it is not essentially faster than Bi-CG. This is confirmed explicitly by the following relation for the Bi-CG residual  $r_k^B$  and the quasi residual  $r_k^Q$  (in exact arithmetic):

$$||r_k^B||_2 = \frac{||r_k^Q||_2}{\sqrt{1 - (||r_k^Q||_2 / ||r_{k-1}^Q||_2)^2}},$$

see [23]: Theorem 4.1. This relation, which is similar to the relation for GMRES and FOM, shows that when QMR gives a significant reduction at step k, then Bi-CG and QMR have arrived at residuals of about the same norm (provided, of course, that the same set of starting vectors has been used).

It is tempting to compare QMR with GMRES, but this is difficult. GMRES really minimizes the 2-norm of the residual, but at the cost of increasing the work of keeping all residuals orthogonal and increasing demands for memory space. QMR does not minimize this norm, but often it has a convergence comparable to GMRES, at a cost of twice the amount of matrix vector products per iteration step. However, the generation of the basis vectors in QMR is relatively cheap and the memory requirements are limited and modest. In view of the relation between GMRES and FOM, it will be no surprise that there is a similar relation between QMR and Bi-CG, for details of this see [47]. This relation expresses that at a significant local error reduction of QMR, Bi-CG and QMR have arrived almost at the same residual vector (similar to GMRES and FOM). However, QMR is preferred to Bi-CG in all cases because of its much smoother convergence behaviour, and also because QMR removes one break-down condition (even when implemented without look-ahead). Several variants of QMR, or rather Bi-CG, have been proposed, which increase the effectiveness of this class of methods in certain circumstances.

In a recent paper, Zhou and Walker [110] have shown that the Quasi-Minimum

Residual approach can be followed for other methods, such as CGS and Bi-CGSTAB, as well. The main idea is that in these methods the approximate solution is updated as

$$x_{i+1} = x_i + \alpha_i p_i,$$

and the corresponding residual is updated as

$$r_{i+1} = r_i - \alpha_i A p_i.$$

This means that  $AP_i = W_i R_{i+1}$ , with  $W_i$  a lower bidiagonal matrix. The  $x_i$  are combinations of the  $p_i$ , so that we can try to find the combination  $P_i y_i$  for which  $||b - AP_i y_i||_2$  is minimal. If we insert the expression for  $AP_i$ , and ignore the fact that the  $r_i$  are not orthogonal, then we can minimize the norm of the residual in a quasi-minimum least squares sense, similar to QMR.

### 5.2 CGS

It is well known that the bi-conjugate gradient residual vector can be written as  $r_j (= \rho_j v_j) = P_j(A)r_0$ , and, likewise, the so-called shadow residual  $\hat{r}_j (= \rho_j w_j)$  can be written as  $\hat{r}_j = P_j(A^T)\hat{r}_0$ . Because of the bi-orthogonality relation we have that

$$(r_j, \hat{r}_i) = (P_j(A)r_0, P_i(A^T)\hat{r}_0) = (P_i(A)P_j(A)r_0, \hat{r}_0) = 0,$$

for i < j. The iteration parameters for bi-conjugate gradients are computed from innerproducts like the above. Sonneveld [89] observed that we can also construct the vectors  $\tilde{r}_j = P_j^2(A)r_0$ , using only the latter form of the innerproduct for recovering the bi-conjugate gradients parameters (which implicitly define the polynomial  $P_j$ ). By doing so, the computation of the vectors  $\hat{r}_j$  can be avoided and so can the multiplication by the matrix  $A^T$ .

The resulting CGS [89] method works in general very well for many unsymmetric linear problems. It converges often much faster than BI-CG (about twice as fast in some cases) and has the advantage that fewer vectors are stored than in GMRES. These three methods have been compared in many studies (see, e.g., [78, 15, 76, 71]).

CGS, however, usually shows a very irregular convergence behaviour. This behaviour can even lead to cancellation and a "spoiled" solution [98]; see also Section 5.3. Freund [48] suggested a squared variant of QMR, which was called TFQMR. His experiments show that TFQMR is not necessarily faster than CGS, but it has certainly a much smoother convergence behavior.

#### 5.3 How serious is irregular convergence?

By very irregular convergence we refer to the situation where successive residual vectors in the iterative process differ in orders of magnitude in norm, and some of these residuals may even be much larger in norm than the starting residual.

We will give an indication why this is a point of concern, even if eventually the (updated) residual satisfies a given tolerance. For more details we refer to Sleijpen et al [86, 87].

We say that an algorithm is *accurate* for a certain problem if the *updated* residual  $r_j$  and the true residual  $b - Ax_j$  are of comparable size for the j's of interest.

In most iteration schemes the approximation for the solution and the corresponding residual are updated independently:

$$x_{j+1} = x_j + w_{j+1}$$
  
 $r_{j+1} = r_j - Aw_{j+1}$ 

In finite precision arithmetic, there will at least be a discrepancy between the two updated quantities due to the multiplication by A. We can account for that by writing

$$r_{j+1} = r_j - Aw_{j+1} - \Delta_A w_{j+1} \quad \text{for each} \quad j, \tag{5.4}$$

where  $\Delta_A$  is an  $n \times n$  matrix for which  $|\Delta_A| \leq n_A \overline{\xi} |A|$ :  $n_A$  is the maximum number of non-zero matrix entries per row of A,  $|B| \equiv (|b_{ij}|)$  if  $B = (b_{ij}), \overline{\xi}$  is the relative machine precision, the inequality  $\leq$  refers to element-wise  $\leq$ .

Note that we have ignored the finite precision errors due to the vector additions. In this case (i.e. situation (5.4) whenever we update the approximation), we have that

$$r_k - (b - Ax_k) = \sum_{j=1}^k \Delta_A w_j = \sum_{j=1}^k \Delta_A (e_{j-1} - e_j), \qquad (5.5)$$

where the perturbation matrix  $\Delta_A$  may depend on j and  $e_j$  is the approximation error in the *j*th approximation:  $e_j \equiv x - x_j$ . Hence,

$$\begin{aligned} |||r_k|| - ||b - Ax_k||| &\leq 2k \, n_A \,\overline{\xi} \, |||A||| \, \max_j ||e_j|| \\ &\leq 2k \, n_A \,\overline{\xi} \, |||A||| \, ||A^{-1}|| \, \max_j ||r_j||. \end{aligned} \tag{5.6}$$

Except for the factor k, the first upper-bound appears to be rather sharp. We see that approximations with large approximation errors may ultimately lead to an inaccurate result. Such large local approximation errors are typical for CGS; Van der Vorst[98] describes an example of the resulting numerical inaccuracy. If there are a number of approximations with comparable large approximation errors, then their multiplicity may replace the factor k; otherwise it will be only the largest approximation error that makes up virtually the bound for the deviation.

A more rigorous analysis, leading to essentially the same results, has been given by Greenbaum [56]. She also includes the errors introduced by the multiplication and subtraction in  $r_{i+1} = r_i - \alpha_i A w_i$ . However, the qualitative results

of this analysis are the same, since in general the errors in the matrix vector product dominate.

#### 5.4 Bi-CGSTAB

Bi-CGSTAB [98] is based on the following observation. Instead of squaring the Bi-CG polynomial, we can construct other iteration methods, by which  $x_i$  are generated so that  $r_i = \tilde{P}_i(A)P_i(A)r_0$  with other  $i^{th}$  degree polynomials  $\tilde{P}$ . An obvious possibility is to take for  $\tilde{P}_i$  a polynomial of the form

$$Q_i(x) = (1 - \omega_1 x)(1 - \omega_2 x)\dots(1 - \omega_i x), \tag{5.7}$$

and to select suitable constants  $\omega_j$ . This expression leads to an almost trivial recurrence relation for the  $Q_i$ . In Bi-CGSTAB  $\omega_j$  in the  $j^{th}$  iteration step is chosen as to minimize  $r_j$ , with respect to  $\omega_j$ , for residuals that can be written as  $r_j = Q_j(A)P_j(A)r_0$ .

Bi-CGSTAB needs two innerproducts more per iteration than CGS.

Bi-CGSTAB can be viewed as the product of Bi-CG and GMRES(1), or rather GCR(1). Of course, other product methods can be formulated as well, and this will be the subject of the next subsection.

#### 5.4.1 Variants of Bi-CGSTAB

Because of the local minimization, Bi-CGSTAB displays a much smoother convergence behavior than CGS, and more surprisingly it often also converges (slightly) faster. One weak point in Bi-CGSTAB is that there is a break-down if  $\omega_j = 0$ . One may also expect negative effects when  $\omega_j$  is small. As soon as the GCR(1) step in Bi-CGSTAB (nearly) stagnates, then the BiCG part in the next iteration step cannot (or can only poorly) be constructed. Another dubious aspect of Bi-CGSTAB is that the factor  $Q_k$  has only real roots by construction. It is well-known that optimal reduction polynomials for matrices with complex eigenvalues may have complex roots as well. If, for instance, the matrix A is real skew-symmetric, then GCR(1) stagnates forever, whereas a method like GCR(2) (or GMRES(2)), in which we minimize over two combined successive search directions, may lead to convergence, and this is mainly due to the fact that then complex eigenvalue components in the error can be effectively reduced.

This point of view was taken in [60] for the construction of the Bi-CGSTAB2 method. In the odd-numbered iteration steps the Q-polynomial is expanded by a linear factor, as in Bi-CGSTAB, but in the even-numbered steps this linear factor is discarded, and the Q-polynomial from the previous even-numbered step is expanded by a quadratic  $1-\alpha_k A - \beta_k A^2$ . For this construction the information from the odd-numbered step is required. It was anticipated that the introduction of quadratic factors in Q might help to improve convergence for systems with complex eigenvalues, and, indeed, some improvement has been observed in practical situations (see also [77]).

In [88], another and even simpler approach was taken to arrive at the desired even-numbered steps, without the necessity of the construction of the intermediate Bi-CGSTAB-type step in the odd-numbered steps. In this approach the polynomial Q is constructed straight-away as a product of quadratic factors. In fact, it is shown in [88] that the polynomial Q can also be constructed as the product of  $\ell$ -degree factors, without the construction of the intermediate lower degree factors. The main idea is that  $\ell$  successive Bi-CG steps are carried out, where for the sake of an  $A^T$ -free construction the already available part of Q is expanded by simple powers of A. This means that after the Bi-CG part of the algorithm, vectors from the Krylov subspace s,  $As, A^2s, ..., A^\ell s$ , with  $s = P_k(A)Q_{k-\ell}(A)r_0$ are available, and it is then relatively easy to minimize the residual over that particular Krylov subspace. There are variants of this approach in which more stable bases for the Krylov subspaces are generated [87], but for low values of  $\ell$  a standard basis satisfies, together with a minimum norm solution obtained through solving the associated normal equations (which requires the solution of an  $\ell$  by  $\ell$  system. In most cases Bi-CGSTAB(2) will already give nice results for problems where Bi-CGSTAB may fail.

#### 5.5 Other product methods

In the paper by Sonneveld [89], it was made clear that we can construct product methods for which  $r_k = H_k(A)P_k(A)r_0$ , in which  $P_k$  denotes the Bi-CG iteration polynomial, and  $H_k$  is any other arbitrary monic polynomial of exact degree k, and he showed that this can be done with the same number of matrix vector multiplications as in k steps of Bi-CG. CGS and the Bi-CGSTAB methods are special instances of these hybrid Bi-CG schemes, but other product methods have been proposed as well.

In [42] it was suggested to take for  $H_k$  Bi-CG-like polynomials in an attempt to keep attractive convergence aspects of CGS, while avoiding the irregular convergence behavior. One idea is to take for  $H_k$  the Bi-CG polynomial obtained with a different starting vector  $w_1$ , which can be done for virtually no additional costs in Bi-CG; this leads to a generalized CGS process. Another idea is to take for  $H_k$  the product of  $P_{k-1}$  and a suitable fixed first degree factor. The idea is that the zeros of  $P_k$  and  $P_{k-1}$  do not coincide, so that it is likely that a zero of  $P_{k-1}$  will be close to a local maximum of  $P_k$ . This avoids the squaring effects in CGS, while it maintains the desired squaring effects in converged eigendirections. Both ideas are easily implemented in an existing CGS code.

In [109] it is suggested to generate  $H_k$  by a three term recurrence relation. This leads to an expression for  $r_k$  of the form

$$r_k = t_{k-1} - \eta_{k-1} y_{k-1} - \zeta_{k-1} A t_{k-1},$$

where  $t_{k-1}$  and  $y_{k-1}$  are auxiliary vectors in the hybrid Bi-CG iteration process, and  $\eta_{k-1}$  and  $\zeta_{k-1}$  represent the coefficients in a Bi-CG like three term recurrence for  $H_k$ . The idea suggested in [109] is to minimize the Euclidean norm of  $r_k$  as a function of  $\eta_{k-1}$  and  $\zeta_{k-1}$ , and this leads to product methods which, according to given examples, can compete with CGS and Bi-CGSTAB methods. It is not clear whether these generalized product methods offer an advantage over the higher order Bi-CGSTAB( $\ell$ ) methods.

## 6 Preconditioning

In order to improve the speed of convergence of iterative methods, one applies usually some form of preconditioning. Many different preconditioners have been suggested over the years, each of these preconditioners more or less successful for restricted classes of problems.

Among all these preconditioners the Incomplete LU factorizations [69, 21] are the most popular ones, and attempts have been made to improve them, for instance by including more fill [70], or by modifying the diagonal of the ILU factorization in order to force rowsum constraints [58, 6, 5, 73, 95, 34], or by changing the ordering of the matrix [96, 97]. A collection of experiments with respect to the effects of ordering is contained in [30]. More recently, it was discovered that a multigrid-inspired ordering can be very effective for discretized diffusion-convection equations, leading in some cases to almost grid-independent speeds of convergence [93, 11], see also [24]. In these publications the ordering strategy is combined with a drop-tolerance strategy for discarding small enough fill-in elements.

Red-black ordering is an obvious approach to improve parallel properties for well-structured problems, but it got a bad reputation from experiments, like those reported in [30]. If carefully done though, they can lead to significant gains in efficiency. Elman and Golub [35] suggested such an approach, in which Red-Black ordering was combined with a reduced system technique. The idea is simple, eliminate the red points, and construct an ILU for the reduced system of black points. Recently, DeLong and Ortega [28] suggested carrying out a few steps of red-black ordered SOR as a preconditioner for GMRES and Bi-CGSTAB. The key to success in these cases seems to be combined effect of fast convergence of SOR for red-black ordering, and the ability of the Krylov subspace to remove stagnations in convergence behaviour associated with a few isolated eigenvalues of the preconditioned matrix.

Saad [80] has proposed some interesting variants on the incomplete LU approach for the matrix A, one of which is in fact an incomplete LQ decomposition. In this approach it is not necessary to form the matrix Q explicitly, and it turns out that the lower triangular matrix L can be viewed as the factor of an incomplete Choleski factorization of the matrix  $A^T A$ . This can be exploited in the preconditioning step, avoiding the use of Q. The second approach was to introduce partial pivoting in ILU, which appears to have some advantages for convection-dominated problems. This approach was further improved by includ-

ing a threshold technique for fill-in [82].

Another major step forward, for important classes of problems, was the introduction of block variants of incomplete factorizations [92, 20, 4], and modified variants of them [4, 66]. It was observed, by Meurant, that these block variants were more successful for discretized 2-dimensional problems than for 3-dimensional problems, unless the '2-dimensional' blocks in the latter case were solved accurately. For discussions and analysis on ordering strategies, in relation to modified block incomplete factorizations, see [67].

Domain decomposition methods were motivated by parallel computing, but it appeared that the approach could be used with success also for the construction of preconditioners. Domain decomposition has been used for problems that arise from discretization of a PDE over a given domain. The idea is to split the given domain into subdomains, and to solve the discretized PDE's over each subdomain separately. The main problem is to find proper boundary conditions along the interior boundaries of the subdomains. Domain decomposition is used in an iterative fashion and usually the interior boundary conditions are based upon information on the approximate solution of neighboring subdomains that is available from a previous iteration step.

It was shown by Chan and Goovaerts [17] that domain decomposition can actually lead to improved convergence rates, provided the number of domains is not too large. A splitting of the matrix with overlapping subblocks along the diagonal, which can be viewed as a splitting of the domain, if the matrix is associated with a discretized PDE and has been ordered properly, was suggested by Radicati and Robert [78]. They suggested to construct incomplete factorizations for the sub-blocks. These sub-blocks are then applied to corresponding parts of the vectors involved, and some averaging was applied on the overlapping parts. A more sophisticated domain-oriented splitting was suggested in [105], for SSOR and MILU decompositions, with a special treatment for unknowns associated with interfaces between the subdomains.

The isolation of sub-blocks was done by Tang [91] in such a way that the sub-blocks corresponded to subdomains with proper internal boundary conditions. In this case it is necessary to modify the sub-blocks of the original matrix such that the subblocks could be interpreted as the discretizations for subdomains with Dirichlet and Neumann boundary conditions in order to force some smoothness of the approximated solution across boundaries. In [90] this was further improved by requiring also continuity of cross-derivatives of the approximate solution across boundaries. The local fine-tuning of the resulting interpolation formulae for the discretizations was carried out by local Fourier-analysis. It was shown that this approach could lead to impressive reductions in numbers of iterations for convection dominated problems.

Another approach that received attention is the concept of constructing an

explicit approximation for the inverse of a given matrix A. The idea is to find a sparse matrix M such that ||AM - I|| is small for some convenient norm. Kolotilina and Yeremin [64] presented an algorithm in which the inverse was delivered in factored form, which has the advantage that singularity of M can be easily detected. In [22] an algorithm is presented which uses the 1-norm for the minimization. We also mention Chow and Saad [18], who use GMRES for the minimization of  $||AM - I||_F$ . This approach has the disadvantage that one does not have easy control over the amount of fill-in in M, and drop-tolerance strategies have to be applied. The approach has the advantage that it can be used to correct explicitly some given implicit approximation, like for instance an ILU decomposition.

An elegant approach was suggested by Grote and Huckle [57]. They also attempt to minimize the F-norm, which is equivalent to the Euclidean norm for the errors in the columns  $m_i$  of M:

$$||AM - I||_F^2 = \sum_{i=1}^n ||Am_i - e_i||_2^2$$

Based on this observation they derive an algorithm that produces the sparsity pattern for the most error-reducing elements of M. This is done in steps, starting with a diagonal approximation, each steps adds more non-zero entries to M, and the procedure is stopped when the norms are small enough or when memory requirements are violated.

## 7 Quo Vadis?

In the previous sections we have seen a large number of new iterative methods, which have come in addition to older methods like Richardson's method, Gauss-Jacobi, Gauss-Seidel, SOR, SSOR, BSOR, S2LOR, Chebyshev iteration, and many more. From the positive point of view, this huge variety in methods has provided us with a powerful toolbox from which we can select the appropriate tool for a given problem. From the point of view of the poor user however, this toolbox is confusingly large. In many practical situations it is not clear at all what method to select. One is often faced with the question from outside the expert community which method is the best, but there is in general no best method. This is nicely illustrated in [71], where examples are given of different types of linear systems with quite different convergence behavior for a few archetypes of the methods discussed before. It is shown that for each method there is a linear system for which the given method is the clear winner, as well as a linear system for which the same method is the clear looser. For instance, there is an example for which the, in general slowly converging, LSQR method is better than the, in general fast converging, GMRES method by orders of magnitudes. This also puts the efforts in perspective where the methods are compared on the basis of some examples. Hopefully this gives us some insight in the properties of these methods, but what we really need is some insight into the characteristics of the linear systems that discriminate between methods.

At present the situation is even more complicated due to the existence of distributed memory parallel computers. These architectures can be used efficiently only if one succeeds in keeping communication between processors below certain levels, and it turns out that the innerproducts, which are needed for the Krylov subspace methods, become a bottleneck for the efficiency for larger numbers of processors [27]. This has led to a revival of interest in innerproduct-free methods, like Chebyshev's method and SOR. It has been suggested to use these methods in combination with the Krylov subspace methods (as a polynomial preconditioner, see [53, 83] for discussions on this), see also our section on preconditioning. An interesting approach to construct an innerproduct-free variant of CG from information obtained after a few steps of CG is hinted in [40], where orthogonal polynomials are constructed based on partial information obtained from CG-iteration polynomials. It has been shown by Golub and Kent [50] that the optimal parameters for the Chebyshev method can also be deduced from the Chebyshev iteration vectors. This has been further worked out by Calvetti et al [16], who show how adaptive Chebyshev methods can be constructed with iteration parameters that are obtained as in [50].

It is well-known that the distribution of eigenvalues has to do with the speed of convergence of Krylov subspace methods, and a way to improve this distribution is known as *Preconditioning*. With a good preconditioner there is not much difference between the various Krylov subspace methods, but this shifts the problem to the identification and construction of good preconditioners. Although much work has been done in this area, there is still a strong need for identifying good preconditioners, in particular for highly non-normal matrices and for indefinite problems. These problems play an important role in *Computational Fluid Dynamics*, and also *Helmholtz* problems lead to indefinite problems. For particular cases, for instance the *Stokes* problem, effective indefinite preconditioners have been proposed [55, 37].

The main difficulty in indefinite preconditioning can be explained as follows. The Krylov subspace methods converge fast when the eigenvalues are clustered, say around 1. This means that the preconditioner, often viewed as an approximation to the inverse of the given matrix, must transfer the eigenvalues to 1. It may easily happen that, due to approximation errors that are intentionally made in order to keep the process efficient, eigenvalues are transferred to values close to 0. In that case the convergence of Krylov subspace methods will be slow, and it may happen that the unpreconditioned iteration process converges faster than the preconditioned iteration.

Complex-valued problems can be treated as the problems discussed in this paper, in particular Hermitian problems can be treated as real symmetric ones, by replacing  $A^T$  by  $A^H$ , and by the appropriate change of innerproduct. Some

problems, for instance in *Electromagnetics*, lead to symmetric complex matrices. Krylov subspace methods that take advantage of this property have been suggested in [43, 99], but preconditioning for this type of problems has remained an open problem.

The approximation error in iterative schemes is not directly available, and has to be estimated or bounded in terms of information obtained in the iteration process. This can be done for special cases, for instance for the Conjugate Gradient method (see [10] and references therein). For most cases, however, the problem of getting realistic error bounds is yet unsolved. In some applications we need information on errors in specified components of the solution, this is also mainly an open problem.

Parallel computing is a very important issue in solving large linear systems, but we have treated this issue only in passing. One might easily dedicate an entire paper for highlighting developments in this area. We restrict ourselves to referring to the overview paper by Demmel et al [29], in which parallel approaches are discussed as well as open problems in this area.

Although the list of interesting open problems can be made arbitrarily longer, space limitations forced us to mention only a few of them.

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