# Conjugate gradients acceleration of coordinate descent for linear systems 

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#### Abstract

This paper introduces a conjugate gradients (CG) acceleration of the coordinate descent algorithm (CD) for linear systems. It is shown that the Kaczmarz algorithm (KACZ) can simulate CD exactly, so CD can be accelerated by CG similarly to the CG acceleration of KACZ (Björck \& Elfving, BIT 1979). Experimental results were carried out on large sets of problems of reconstructing bandlimited functions from random sampling. The randomness causes extreme variance between different instances of these problems, thus causing extreme variance in the advantage of CGCD over CD . The reduction of the number of iterations by CGCD varies from about $50 \%$ to $90 \%$ and beyond. The implementation of CGCD is simple. CGCD can also be used for the parallel solution of linear systems derived from partial differential equations, and for the efficient solution of multiple right-hand-side problems and matrix inversion.


Keywords. Coordinate descent; CD; CGCD; CGMN; conjugate gradients acceleration; GaussSeidel; Kaczmarz algorithm; linear systems, matrix inversion, multiple right-hand-sides, parallelism.

AMS Categories. 65B99; 65F10; 65F20.

## 1 Introduction

The coordinate descent algorithm (CD) is a well-known and widely used optimization algorithm; see, for example, $[13,16]$ for details and applications. CD is also used for solving linear systems as follows: given a linear system $A x=b$, each step of CD consists of choosing a coordinate $x_{i}$ and minimizing $\|A x-b\|^{2}$ w.r.t. $x_{i}$ by setting $\frac{\partial}{\partial x_{i}}\|A x-b\|^{2}=0$ and extracting a new value for $x_{i}$. This approach is generally referred to as "CD for linear systems". There are several papers on various acceleration methods of CD, such as [1,9,10], and specifically for linear systems, see [8, 15]. CD for linear systems is identical to the Gauss-Seidel iteration on the system $A^{T} A x=A^{T} b$; see for example [11, p. 594] and [16, §1.4]. $A^{T} A x=A^{T} b$ is also called NR (for "normal" and "residual" associated with the minimization of $\|b-A x\|_{2}$; see Saad [12, §8.1]. In the following, "CD" will refer to CD for linear systems.

[^0]The rest of the paper is organized as follows. §2 presents the Kaczmarz algorithm (KACZ) [7] and shows the geometric identity between KACZ and CD. This is followed by the CG acceleration of KACZ, called CGMN, developed by Björck and Elfving [2], and the proof that this acceleration can be applied to CD. $\S 3$ presents the CG acceleration of CD and $\S 4$ presents the experiments. $\S 5$ presents an efficient construction of the pre-calculations for sparse systems, and $\S 6$ concludes with a summary and suggestions for the efficient solution of multiple right-hand-sides problems and matrix inversion.

## 2 Kaczmarz, CD and their CG acceleration

Consider a consistent system of linear equations

$$
\begin{equation*}
A x=b \tag{1}
\end{equation*}
$$

KACZ operates as follows: starting from an arbitrary point as the initial iterate, that point in space is orthogonally projected towards a plane defined by one of the equations, thus creating a new iterate. This process is continued sequentially for all the following equations, and then repeated until some stopping criterion is satisfied. There are several variations of this procedure, such as randomized selection of the equations and a symmetric approach in which the equations are alternately handled in a forward and backward sequence.

Let $x^{0}$ be a chosen initial iterate. For $k \geq 0, x^{k+1}$ is obtained from $x^{k}$ by projecting $x^{k}$ orthogonally towards the hyperplane determined by the $k$ th equation, i.e.,

$$
\begin{equation*}
x^{k+1}=x^{k}+\lambda \frac{b_{k}-\left\langle a^{k}, x^{k}\right\rangle}{\left\|a^{k}\right\|_{2}^{2}} a^{k}, \tag{2}
\end{equation*}
$$

where $a^{k}$ is the $k$ th matrix row, $b_{k}$ is the $k$ th RHS element, and $0<\lambda<2$ is a relaxation parameter that determines whether the projection is exactly on the hyperplane determined by the equation ( $\lambda=1$ ), or before or after the hyperplane. It is more efficient to initially divide every equation by the $L_{2}$ norm of its corresponding row, thus avoiding the division by $\left\|a^{k}\right\|_{2}^{2}$ at every step. KACZ is also known as successive overrelaxation (SOR) on the system $A A^{T} y=b, x=A^{T} y$. Successive SOR (SSOR) is KACZ carried out in the forward direction and then in the backward direction. The robustness of KACZ follows from the following observation: after all the equations are normalized, the diagonal of $A A^{T}$ consists of ones, while the off-diagonal elements are all smaller than one. In a consistent system, the KACZ iterates converge towards a solution (there may be many solutions if the system is under-determined). We can now prove the following:

Lemma 1. Let CD operate on a consistent system of equations whose solution is $x_{1}^{*}, \ldots, x_{n}^{*}$. Then, $K A C Z$ can be applied (in parallel to $C D$ ) to the system of equations $x_{1}=x_{1}^{*}, \ldots, x_{n}=x_{n}^{*}$, with relaxation parameters that bring each iterate of $K A C Z$ to the same position in space that $C D$ brings it.

Proof: Let $\left(x_{1}^{k}, \ldots, x_{n}^{k}\right)$ be some current value of the coordinates obtained by CD, and consider one step of CD that changes some coordinate $x_{i}^{k}$ to a new value $x_{i}^{k+1} . x_{i}^{k+1}$ is now closer to $x_{i}^{*}$ than $x_{i}^{k}$, so the change of the $i$ th coordinate moves the current iterate $\left(x_{1}^{k}, \ldots, x_{n}^{k}\right)$ closer to the plane $x_{i}=x_{i}^{*}$. Only the $i$ th coordinate is changed so the direction of this movement is orthogonal to the plane $x_{i}=x_{i}^{*}$.

Consider now a specific KACZ operation on the system $x_{1}=x_{1}^{*}, \ldots, x_{n}=x_{n}^{*}$, running in parallel to CD. The initial iterate of KACZ is taken as the initial iterate of CD. For each step of CD on a coordinate $x_{i}^{k}$, KACZ performs one step on the equation $x_{i}=x_{i}^{*}$, and the relaxation parameter is chosen to bring the new iterate to the same position that CD brings it. Clearly, KACZ produces exactly the same sequence of iterates as CD.

In a seminal paper, Björck and Elfving [2] showed that KACZ can be accelerated by CG, and called this algorithm CGMN. Based on Lemma 1, we have the following:

Theorem 1. $C D$ can be accelerated by $C G$.
Proof: The original CGMN [2] assumed a constant relaxation parameter, but now we need to accelerate KACZ with variable relaxation parameters as in Lemma 1. In [6, §3] it was shown that the CG acceleration of KACZ can use different relaxation parameters for different equations, and this enabled a parallel version of CGMN (called CARP-CG) for solving huge linear systems derived from partial differential equations. See also [5, §2]. It is easy to see from [6, §3] that the cyclic relaxation parameters can be replaced by any relaxation parameter at every instance of Eq. (2). It now follows from Lemma 1 that the CG acceleration of (the specific) KACZ can be applied to CD.

The following are some related comments regarding CG. Given the system (1), consider the system NR mentioned previously:

$$
\begin{equation*}
A^{T} A x=A^{T} b \tag{3}
\end{equation*}
$$

The system (3) can also be solved directly by CG, and this is known as CGNR; see [12, §8.3.1]. There is also the CGNE algorithm, also known as Craig's Method, which is a CG acceleration of the system $A A^{T} u=b, x=A^{T} u$, which is referred to as NE (normal equations); see [12, §8.3.2]. In [4], CGNR and CGNE produced identical results on the problem of reconstructing bandlimited functions from random sampling, but they were significantly worse than CGMN, as can be seen from [4, Figs. $4 \& 6$ ].

## 3 CG acceleration of CD

CD and its CG acceleration will now be presented in detail. Let $A$ be a matrix of size $m \times n$ ( $m$ rows by $n$ columns), and $b$ the right-hand side. Our purpose is to find a vector $x=\left(x_{1}, \ldots, x_{n}\right) \in \mathbb{R}^{n}$
such that $\|A x-b\|$ is minimal. CD proceeds by successively setting $\frac{\partial}{\partial x_{i}}\|A x-b\|^{2}=0$, for $i=$ $1, \ldots, n$. In the following, the dot product of two vectors is represented by "•". We introduce some notations:

- $A_{i}$ denotes the $i$ th column of $A$.
- $A_{i}^{j}=A_{i} \bullet A_{j}$ if $i \neq j$, else 0 .
- $A_{i}^{*}=\left(A_{i}^{1}, \ldots, A_{i}^{n}\right)$. (Note that the $i$ th element, $A_{i}^{i}$, is zero.)

In order to save some computation time, the columns of $A$ are normalized, i.e., each column is divided by its $L_{2}$-norm. It is easy to see that setting $\frac{\partial}{\partial x_{i}}\|A x-b\|^{2}=0$ produces the following iteration: $x_{i}=A_{i} \bullet b-x \bullet A_{i}^{*}$. The CD algorithm is the following:

## CD: The coordinate descent algorithm

1. compute all the terms $A_{i} \bullet b$ and $A_{i}^{*}$;
2. set $x \in \mathbb{R}^{n}$ to an arbitrary value ;
3. repeat until some termination goal is reached:
4. $\quad \mathbf{f o r} i=1$ to $n$ :
5. $x_{i}=A_{i} \bullet b-x \bullet A_{i}^{*}$;
6. end repeat

Note that in the term $x \cdot A_{i}^{*}$ in line $5, x_{i}$ is multiplied by zero, so $x_{i}$ does not appear in the RHS.
In the following CGCD, $\alpha, \beta, \gamma, \delta, \delta 1$ are auxiliary variables and $p, q, r$ are vectors of size $n$.
CGCD: CG acceleration of CD:

1. compute all the terms $A_{i} \bullet b$ and $A_{i}^{*}$ for $1 \leq i \leq n$;
2. set $x=r \in \mathbb{R}^{n}$ to an arbitrary value ;
3. forward and backward loops with $b$ :
4. for $i=1$ to $n$ :
5. $\quad r_{i}=A_{i} \bullet b-r \bullet A_{i}^{*}$;
6. $\quad$ for $i=n$ downto 1 :
7. $r_{i}=A_{i} \bullet b-r \bullet A_{i}^{*}$;
8. $p=r=r-x$;
9. $\delta=r \bullet r$;
10. CG iterations:
11. repeat until some termination goal is reached:
12. set $q=p$;
13. forward and backward loops without $b$ :
14. for $i=1$ to $n$ :
15. $\quad q_{i}=-q \cdot A_{i}^{*}$;
16. $\quad$ for $i=n$ downto 1 :
17. $q_{i}=-q \cdot A_{i}^{*}$;
18. $q=p-q ; \gamma=p \bullet q$;
19. $\alpha=\delta / \gamma ; x=x+\alpha p$;
20. $r=r-\alpha q ; \delta 1=r \bullet r$;
21. $\beta=\delta 1 / \delta$;
22. $p=r+\beta * p ; \delta=\delta 1$;
23. end repeat

Consider now the complexity question: does CG impose a great burden on the coordinate descent? Note that the computations of $A_{i}^{j}$ and $A_{i}^{*}$ are used for both CD and CDCG, so they are not considered in this comparison. From the CD algorithm, we can see that the number of products of one iteration in the loop of lines $4-5$ is of order $O\left(n^{2}\right)$, and so is the number of products (in one iteration) of the loops of CGCD (lines 15-16 and 18-19). In contrast, the number of products in one CG iteration needed for the CG in CGCD is only $5 n$, and this is negligible. In comparing the number of iterations required by CD and CGCD, for the same problem, each iteration of CGCD is counted as two iterations, due to the two loops in CGCD.

Note that both CD and CGCD require the pre-computation of two sets of constants:

$$
\begin{align*}
A B & =\left\{A_{i} \bullet b \mid 1 \leq i \leq n\right\}  \tag{4}\\
A A & =\left\{A_{i}^{*} \mid 1 \leq i \leq n\right\} \tag{5}
\end{align*}
$$

The space required by $A B$ is $O(n m)$, and the space of $A A$, in the case of a full rank matrix, is $O\left(n^{2} m\right)$. The details for constructing $A B$ and $A A$ for sparse systems are presented in Section 5.

## 4 Experiments

### 4.1 Background

CD and CGCD were tested on the problem of recovering bandlimited signals from random sampling. A well-known paper on this topic is due to Strohmer and Vershynin [14], who got excellent results on this problem by using the Kaczmarz algorithm. Every sample contributes one equation to the problem. Before applying KACZ, the equations were sorted and every equation was assigned a probability weight proportional to the average distance of the sample from its nearest points. This concept is due to Feichtinger and Gröchenig [3]. In the experiments of [14], KACZ was applied randomly to the equations according to the weights. The experiments were done on a problem of 101 variables and 700 equations (samples). For convenience, the ratio between equations and variables will be denoted as EVR , so $\mathrm{EVR} \approx 7$ in this problem.

This problem was also studied by the author in [4], where it turned out that when EVR was smaller than 5 , the number of projections required by KACZ rises (seemingly) exponentially; see the KACZ plots in [4, Fig. 3]. However, CGMN provided excellent results for $2 \leq \mathrm{EVR} \leq 6$.

The problem of recovering bandlimited signals from random samples was chosen as a test-bed for comparing CD and CGCD due to the extreme variance between different instances of seemingly identical problems. The experimental problem that is being solved appears in [14, §4.1], and also in [4, §3.1]. It can be described as follows:

A bandlimited function $f(t)$ is known to be of the form

$$
\begin{equation*}
f(t)=\sum_{\ell=-r}^{r} x_{\ell} e^{2 \pi i \ell t} \tag{6}
\end{equation*}
$$

where $t$ is time, $x_{\ell} \in \mathbb{C}, r$ is a known positive integer (called the "bandwidth"), $n$ is an integer set as $2 r+1, m \geq n$, and $f\left(t_{j}\right)$ for $t_{j} \in \mathbb{R}$ and $1 \leq j \leq m$ are known samples of the values of $f(t)$ taken at irregular points in time. Our problem is to determine the values of $x_{1}, \ldots, x_{n}$ which will provide the values of $f(t)$ at any point in time. The linear system that provides the values of $x_{1}, \ldots, x_{n}$ is obtained by setting

$$
\begin{equation*}
f\left(t_{j}\right)=\sum_{\ell=-r}^{r} x_{\ell} e^{2 \pi i \ell t_{j}}, \quad \text { for } \quad 1 \leq j \leq m . \tag{7}
\end{equation*}
$$

The main problem of irregular sampling is that large gaps between samples make it more difficult to determine the values of $x_{i}$, hence the large computational differences between different instances of this problem; see $[4, \S 4.1]$ for the effects of large gaps. Note that every instance of setting up a linear system is derived from randomly selecting the values of $t_{1}, \ldots, t_{m}$; if these values are spread quite evenly, then CD will converge fairly quickly, otherwise, CD converges slowly, and this is where CGCD has a big advantage.

### 4.2 Results

The only way to compare CD and CGCD is to run a large number of experiments in which every instance is solved by both CD and by CGCD. Let $R$ denote the ratio of the number of iterations required by CGCD divided by the number of iterations required by CD on exactly the same problem. The experiments show that $R$ can vary in different cases from about 0.03 up to about 0.5 . In many cases CGCD provides a very significant improvement over CD . The C program for these experiments, called cgcd.c, is available for download ${ }^{1}$; the program is self-explanatory.

Figures 1 and 2 show comparisons between the number of iterations required by CD and CGCD for 50 and 100 runs, respectively. The relative residual goal was set at $10^{-13}$ and the maximum number of iterations was set at 100,000 . The results are sorted according to increasing numbers of iterations required by CD . The extreme variance stands out clearly, and in other experiments, some CD runs required significantly more iterations. The statistics show that the average number of iterations required by CGCD is around $13 \%$ of the average of CD. The most significant result

[^1]of these experiments is that while about $20 \%$ of the CD runs required thousands of iterations, CGCD did not need more than 350 iterations. Some of the runs fail to reach the prescribed relative residual goal, and the failure percentage is typically around $2 \%$ for CD and $4 \%$ for CGCD, but it may vary widely. The consequence is that CGCD is somewhat less stable than CD on some of these particular problems, so in practice, it may be useful to run both versions in parallel.


Figure 1: Comparison of CD and CGCD on the same 50 problems. The results are sorted according to increasing number of iterations required by CD.


Figure 2: Comparison of CD and CGCD on the same 100 problems. The results are sorted according to increasing number of iterations required by $C D$.

## 5 Construction of $A B$ and $A A$ for sparse systems

The issue of setting up $A B$ and $A A$, Eqs. (4) and (5), needs some explanation. If the system matrix is full then $A B$ and $A A$ can be calculated easily as arrays, as in the above experiments. Otherwise, $A B$ and $A A$ can be calculated as explained below. An important issue here is that in each row of $A$, the nonzero elements can be accessed easily, i.e., without traversing the entire row. This should not be a problem because every row is made up by the parameters of a single equation and the sequential access can be enabled when the matrix is formed. Also, in huge matrices, it is customary to have some compact form of the non-zero coefficients of a matrix row.

Consider first the data structure required by $A B . A_{j} \bullet b$ is the dot product of column $A_{j}$ and $b$. We compute $A_{j} \bullet b$ for every $j$ as follows: for every $1 \leq j \leq n$, a variable $A B_{j}$ is set to zero. Next, for every matrix row $i$, we access all its non-zero elements, e.g., $a_{i, 5}, a_{i, 50}, a_{i, 90}$, multiply every one of them by $b_{i}$, and add these products to their corresponding $A B_{j}$, e.g., $A B_{5}+=a_{i, 5} \times b_{i}$, $A B_{50}+=a_{i, 50} \times b_{i}$, and $A B_{90}+=a_{i, 90} \times b_{i}$.

The data structure required by $A A$ consists of the sequences $A_{i}^{*}=\left(A_{i, 1}, \ldots, A_{i, n}\right)$, for $1 \leq i \leq n$. Every $A_{i, j}$ is $A_{i} \bullet A_{j}$ if $i \neq j$ and 0 otherwise. Consider the following approach: for every $A_{i}^{*}$, a linked list is constructed with a header $\left[i \mid A_{i}^{*}\right]$. The pointer of the header is initially set to null. This list will contain the non-zero elements of $A_{i} \bullet A_{j}$ together with the index $j$, and ordered by increasing values of $j$; for example:

$$
\begin{equation*}
\left[i \mid A_{i}^{*}\right] \rightarrow\left[5 \mid A_{i} \bullet A_{5}\right] \rightarrow\left[12 \mid A_{i} \bullet A_{12}\right] \rightarrow \cdots \rightarrow\left[340 \mid A_{i} \bullet A_{340}\right] \tag{8}
\end{equation*}
$$

This structure will enable the efficient calculation of dot products such as $q \cdot A_{i}^{*}$ in lines 16 and 19 of CGCD.

The construction of a list as in Eq. (8) proceeds as follows: We traverse the rows in sequence, for $1 \leq k \leq m$. The non-zero elements of row 1 are the parameters of one equation, e.g., $a_{1,5}, a_{1,20}$, and $a_{1,70}$. We now initiate 3 constants to zero, each constant labeled by a pair of the nonzero parameters: $A_{5,20}, A_{5,70}$, and $A_{20,70}$. We now compute all possible products of the parameters and add them to the 3 constants that were set up, according to the indices of the constants $A_{i, j}$ :

$$
\begin{align*}
A_{5,20}+ & =a_{1,5} \times a_{1,20} \\
A_{5,70}+ & =a_{1,5} \times a_{1,70}  \tag{9}\\
A_{20,70}+ & =a_{1,20} \times a_{1,70}
\end{align*}
$$

We now add the three elements $A_{i, j}$ of Eq. (9) to the corresponding lists of $A_{5}^{*}, A_{20}^{*}, A_{70}^{*}-$ see Eq. (8). Every $A_{i, j}$ in the lists must carry the column number which differs from the header, and the
lists are sorted by the indices:

$$
\begin{align*}
{\left[5 \mid A_{5}^{*}\right] } & \rightarrow\left[20 \mid A_{5,20}\right] \rightarrow\left[70 \mid A_{5,70}\right] \\
{\left[20 \mid A_{20}^{*}\right] } & \rightarrow\left[5 \mid A_{5,20}\right] \rightarrow\left[70 \mid A_{20,70}\right]  \tag{10}\\
{\left[70 \mid A_{70}^{*}\right] } & \rightarrow\left[5 \mid A_{5,70}\right] \rightarrow\left[20 \mid A_{20,70}\right]
\end{align*}
$$

Note that $A_{i, j}=A_{j, i}$. Continuing with the following rows, we get two types of new products. For example, if in row 2 we get a product $c=a_{2,5} \times a_{2,70}$, then $c$ should be added to $A_{5,70}$ in the list of $A_{5}^{*}$ and also to $A_{5,70}$ in the list of $A_{70}^{*}$. Otherwise, new elements of type $A_{i, j}$ are created, the remaining constants are added to them as above, and they are inserted into the corresponding lists $\left[i \mid A_{i}\right]$ as in Eq. (9). Note that if there are $k$ non-zero elements in a row, then there will be $\binom{k}{2}$ products.

At the end, the linked lists can be converted to arrays to save some computation time. In some cases, the sizes of the lists can be quite large, so it would be more efficient to use binary trees instead of linked lists because the insertion time is $\log _{2}$ of the time in lists. At the end, the inorders of the trees need to be converted to linked lists or arrays.

## 6 Conclusions and further applications

It has been shown that the CG acceleration of coordinate descent for linear systems, denoted CGCD, can be significantly faster than plain CD, so it can benefit applications that already use CD for linear systems, and perhaps other applications for which CD is too slow.

CGCD can also be useful for the problematic issue of multiple right-hand-sides because the time consuming setup of $A A$ depends only on the matrix, and setting up new structures for $A B$ is simple. As a consequence, CGCD can also deal with matrix inversion efficiently by successively choosing right-hand-sides as $(1,0, \ldots, 0),(0,1,0, \ldots, 0), \ldots,(0, \ldots, 0,1)$. In a parallel setting, after $A A$ has been set up, the computations with the various RHSs can be distributed among several processors.

Parallelism for sparse matrices: in a full-rank matrix, such as in the experiments of this paper, CD (and CGCD) cannot be paralleled because CD is inherently sequential. In a sparse matrix, CD can be parallelize as follows: we define two variables $x_{i}$ and $x_{j}$ as dependent if they appear together in some equation, and otherwise they are independent. We will first show that if $x_{i}$ and $x_{j}$ are independent, then they can be handled in parallel. Consider the dot product $A_{i} \cdot A_{j}=0$ and also line 5 of the CD algorithm $x_{i}=A_{i} \bullet b-x \bullet A_{i}^{*}$ : the vector $A_{i}^{*}$ is defined as $\left(A_{i, 1}, \ldots, A_{i, n}\right)$ and $A_{i, j}=A_{i} \bullet A_{j}=0$. Hence, $x_{j}$ has no effect on the new value of $x_{i}$ (obtained from line 5), and vice versa. It follows that the two updates of $x_{i}$ and $x_{j}$ can be done in parallel.

To deal with dependent variables, we need to divide the variables into disjoint sets $S_{1}, \ldots, S_{k}$, which will be handled sequentially, and, if a variable $x_{i}$ is in set $S_{j}$, then all its dependent variables belong
to different sets. This way, all the required updates of a set can be done in parallel. The number of sets should be as small as possible. As an example, consider the problem of solving a partial differential equation in two dimensions using a finite difference, second-order scheme, in which every equation consists of 5 variables forming a 5-point stencil on the grid - see the central cross in Fig. 3. The division of the variables into two sets is shown by the diagonal red (solid) and blue (dashed) lines. Note, for example, that the central grid point is in the red set and all its 4 neighbors are in the blue set. A $3 \times 3$ stencil for higher order accuracy will require three sets.


Figure 3: Coloring a 2D grid with diagonal lines of two colors for 5-point stencils for PDE problems.

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Data availability: The link https://cs.haifa.ac.il/~gordon/cgcd.c provides the C-program for the experiments. The text at the beginning of the program explains in detail how to run it.


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