Reformulation of the Anderson method using singular value decomposition for stable convergence in self-consistent calculations

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Abstract

The Anderson method provides a significant acceleration of convergence in solving nonlinear simultaneous equations by trying to minimize the residual norm in a least-square sense at each iteration step. In the present study I use singular value decomposition to reformulate the Anderson method. The proposed version contains only a single parameter which should be determined in a trial-and-error way, whereas the original one contains two. This reduction leads to stable convergence in real-world self-consistent electronic structure calculations.

Keywords nonlinear simultaneous equations, least-square method, the Broyden method, the Pulay method, electronic-structure calculations

Research Activity Group Algorithms for Matrix / Eigenvalue Problems and their Applications

1. Introduction

In the past few years first-principles calculations based on density functional theory [1] have gained enormous interest among solid-state physicists, materials scientists, and quantum chemists. The Kohn-Sham equation [2], which plays a vital role within the density functional theory, is not only an eigenvalue problem, but also an implicitly defined, nonlinear fixed-point problem of interelecton potential [3–5] at least when local density approximation [2] is introduced. In other words, the Kohn-Sham equation is solved when the self-consistent interelectron potential is found. The Anderson method [6] is frequently employed for this purpose. It should be noted that the Pulay method [7] and limited-memory modifications [8–11] of the second Broyden method [12] are essentially equivalent to the Anderson method [13], while the first Broyden method can also be cast into the limited-memory form [14,15].

Suppose that for a system of nonlinear equations \( \mathbf{F}(\mathbf{x}) = 0 \), there are independent variable column vectors, 

\[
\{ \mathbf{x}_n, \mathbf{x}_{n-1}, \ldots, \mathbf{x}_{n-k} \}
\]

which are hopefully approaching a solution, and accompanying residual column vectors, 

\[
\{ \mathbf{y}_n, \mathbf{y}_{n-1}, \ldots, \mathbf{y}_{n-k} \}
\]

where subscripts denote iteration steps. In a simple iteration method, the independent vector at the \((n+1)\)th iteration step is given by 

\[
\mathbf{x}_{n+1} = \mathbf{x}_n + \alpha \mathbf{y}_n,
\]

where \( \alpha \) is a mixing factor ranging from a scalar to a preconditioning matrix [16,17] to a nonlinear procedure [18,19]. In the Anderson method, however, a virtual residual vector, 

\[
\mathbf{y}^*_n = \mathbf{y}_n + \sum_{1 \leq \nu \leq k} \gamma_\nu \frac{\mathbf{y}_{n+1} - \mathbf{y}_{n-\nu}}{\|\mathbf{y}_{n+1} - \mathbf{y}_{n-\nu}\|},
\]

is introduced. Here \( \gamma_\nu \) are parameters to be so determined that the virtual residual norm \( \| \mathbf{y}^*_n \| \) is minimized in a least-square sense. Then an accompanying virtual independent vector, 

\[
\mathbf{x}^*_n = \mathbf{x}_n + \sum_{1 \leq \nu \leq k} \gamma_\nu \frac{\mathbf{x}_{n+1} - \mathbf{x}_{n-\nu}}{\|\mathbf{x}_{n+1} - \mathbf{x}_{n-\nu}\|},
\]

is defined on assumption of linearity. \( \mathbf{x}^*_n \) is expected to be a minimizer for \( \| \mathbf{F} \| \) within the available subspace \( \{ \mathbf{x}_n, \mathbf{x}_{n-1}, \ldots, \mathbf{x}_{n-k} \} \). Last, the independent variable vector for the next step is predicted by applying the simple iteration method to \( \mathbf{x}^*_n \) and \( \mathbf{y}^*_n \) as 

\[
\mathbf{x}_{n+1} = \mathbf{x}^*_n + \alpha \mathbf{y}^*_n.
\]

In practice, a specialized linear solver should be used to determine the parameters \( \gamma_\nu \) reliably without encountering numerical instability. This means that a maximum condition number must be set for the linear solver beforehand. Moreover a limit for the number of the previous independent and residual vectors considered must be also set beforehand. Since the two parameters cannot be obtained a priori, they are determined in an ad hoc way. In the present study I eliminate the latter by reformulating the Anderson method based on singular value decomposition (SVD) [20]. This makes application of the Anderson method a little easier. Furthermore, stable convergence is achieved in the sense that the numbers of iteration steps required by self-consistency are

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Keywords: nonlinear simultaneous equations, least-square method, the Broyden method, the Pulay method, electronic-structure calculations

Research Activity Group: Algorithms for Matrix / Eigenvalue Problems and their Applications
less sensitive with respect to the remaining parameters as confirmed by test calculations.

2. Conventional Method

For simplicity I define a rectangular matrix as

\[
Y_n = \begin{pmatrix}
\tilde{y}_n - \tilde{y}_{n-1} & \tilde{y}_{n-1} - \tilde{y}_{n-2} \\
\|\tilde{y}_n - \tilde{y}_{n-1}\| & \|\tilde{y}_{n-1} - \tilde{y}_{n-2}\|
\end{pmatrix},
\]

and a column vector containing \(\gamma_v\) as

\[
\Gamma = \begin{pmatrix}
\gamma_1 \\
\gamma_2 \\
\vdots \\
\gamma_k
\end{pmatrix}.
\]

I omit the right-pointing arrow above \(\Gamma\) to emphasize that in general \(\Gamma\) is different form \(x_0\) and \(y_0\) in the number of rows. Using \(Y_n\) and \(\Gamma\), (2) is rewritten as

\[
\tilde{y}_n^* = \tilde{y}_n + Y_n \Gamma.
\]

The formal solution of \(\Gamma\) which minimizes \(\|\tilde{y}_n^*\|\) is given by

\[
\Gamma = - (Y_n^T Y_n)^{-1} Y_n^T \tilde{y}_n.
\]

Determining \(\Gamma\) using (8) literally should be discouraged, because of the potentially large condition number of \(Y_n^T Y_n\). Instead, \(\Gamma\) is computed in a following way. First, if the SVD is employed, \(Y_n\) is factorized into

\[
Y_n = \mathcal{Y}_n \Sigma_n V_n^T,
\]

where \(\mathcal{Y}_n\) and \(V_n\) are matrices containing the left and right singular vectors of \(Y_n\), respectively, while \(\Sigma_n\) is a diagonal matrix of the singular values. Then a corresponding truncated factorization,

\[
Y_n \approx \mathcal{Y}_n \Sigma_n \mathcal{V}_n^T,
\]

is considered. Here \(\Sigma_n\) is a diagonal matrix of the \(l\) largest singular values of \(\Sigma_n\), while \(\mathcal{Y}_n\) and \(\mathcal{V}_n^T\) contain the \(l\) column vectors of \(\mathcal{Y}_n\) and the \(l\) row vectors of \(V_n^T\) corresponding to the \(l\) largest singular values, respectively. \(l\), the effective rank of \(Y_n\), is the largest integer so determined that the condition number of \(\Sigma_n\) does not exceed the first predetermined limit \(s_{\text{max}}\). Of course, \(l\) can be equal to \(k\). Last, \(\Gamma\) is given by

\[
\Gamma = - \mathcal{V}_n \Sigma_n^{-1} \mathcal{Y}_n^T \tilde{y}_n.
\]

At the next iteration step, \(Y_{n+1}\) may be set to be

\[
Y_{n+1} = \begin{pmatrix}
\tilde{y}_{n+1} - \tilde{y}_n \\
\|\tilde{y}_{n+1} - \tilde{y}_n\|
\end{pmatrix} Y_n
\]

\[
= \begin{pmatrix}
\tilde{y}_n - \tilde{y}_{n-1} \\
\|\tilde{y}_n - \tilde{y}_{n-1}\|
\end{pmatrix} \begin{pmatrix}
\tilde{y}_{n-1} - \tilde{y}_{n-2} \\
\|\tilde{y}_{n-1} - \tilde{y}_{n-2}\|
\end{pmatrix}
\]

\[
\cdots
\begin{pmatrix}
\tilde{y}_1 - \tilde{y}_0 \\
\|\tilde{y}_1 - \tilde{y}_0\|
\end{pmatrix} \begin{pmatrix}
\tilde{y}_0 - \tilde{y}_{-1} \\
\|\tilde{y}_0 - \tilde{y}_{-1}\|
\end{pmatrix}.
\]

The usual practice is, however, that if \(k\) has reached the second predetermined limit \(k_{\text{max}}\), the rightmost (oldest) column of the right-hand side of (12) is removed as

\[
Y_{n+1} = \begin{pmatrix}
\tilde{y}_{n+1} - \tilde{y}_n \\
\|\tilde{y}_{n+1} - \tilde{y}_n\|
\end{pmatrix} \begin{pmatrix}
\tilde{y}_{n-1} - \tilde{y}_{n-2} \\
\|\tilde{y}_{n-1} - \tilde{y}_{n-2}\|
\end{pmatrix}
\]

\[
\cdots
\begin{pmatrix}
\tilde{y}_{k+1} - \tilde{y}_k \\
\|\tilde{y}_{k+1} - \tilde{y}_k\|
\end{pmatrix} \begin{pmatrix}
\tilde{y}_k - \tilde{y}_{k-1} \\
\|\tilde{y}_k - \tilde{y}_{k-1}\|
\end{pmatrix}.
\]

3. Proposed Method

Along with (5), I define a rectangular matrix containing the independent variable vectors as

\[
X_n = \begin{pmatrix}
x_n - x_{n-1} \\
\|x_n - x_{n-1}\|
\end{pmatrix} \begin{pmatrix}
x_{n-1} - x_{n-2} \\
\|x_{n-1} - x_{n-2}\|
\end{pmatrix}
\]

\[
\cdots
\begin{pmatrix}
x_{n-k+1} - x_{n-k} \\
\|x_{n-k+1} - x_{n-k}\|
\end{pmatrix} \begin{pmatrix}
x_{n-k} - x_{n-k-1} \\
\|x_{n-k} - x_{n-k-1}\|
\end{pmatrix}.
\]

Since \(\mathcal{Y}_n = Y_n \mathcal{V}_n \Sigma_n^{-1}\) holds, a similar quantity,

\[
\mathcal{X}_n = X_n \mathcal{V}_n \Sigma_n^{-1},
\]

is introduced. \(\mathcal{X}_n\) and \(\mathcal{Y}_n\) are computed by working with \(\mathcal{X}_n\) and \(\mathcal{Y}_n\) as

\[
\tilde{x}_n = \tilde{x}_n + \mathcal{X}_n \Gamma'
\]

\[
\tilde{x}_n^* = \tilde{y}_n + \mathcal{X}_n \Gamma',
\]

respectively, where \(\Gamma'\) is obtained by

\[
\Gamma' = - \mathcal{X}_n^T \tilde{y}_n.
\]

At the next iteration step, \(X_{n+1}\) and \(Y_{n+1}\) are updated by

\[
X_{n+1} = \begin{pmatrix}
x_{n+1} - x_n \\
\|x_{n+1} - x_n\|
\end{pmatrix} \begin{pmatrix}
x_n - x_{n-1} \\
\|x_n - x_{n-1}\|
\end{pmatrix}
\]

\[
\cdots
\begin{pmatrix}
x_{n+1-k+1} - x_{n+1-k} \\
\|x_{n+1-k+1} - x_{n+1-k}\|
\end{pmatrix} \begin{pmatrix}
x_{n+1-k} - x_{n+1-k-1} \\
\|x_{n+1-k} - x_{n+1-k-1}\|
\end{pmatrix}.
\]

4. Test Calculations

The conventional and proposed methods are compared by applying them to first-principles calculations for wurtzite ZnO based on plane-wave, pseudopotential
Table 1. Iterations required to reach self-consistency for wurzite ZnO with various maximum singular values $s_{\text{max}}$ and history data limits $k_{\text{max}}$. Maximum numbers of history data reached in the proposed method are shown in parentheses. Note that lattice parameters and atomic positions in the unit cell are optimized.

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$\alpha = 1.6$

approach [21, 22]. Lattice parameters and atomic positions in the unit cell are also optimized. Remaining technical details are explained elsewhere [23]. The mixing factor $\alpha$ is chosen to be a scalar parameter.

The parameters and results are shown in Table 1. For $\alpha = 0.8$, both the methods have achieved fast convergence of iteration steps below 40. Clearly, however, the proposed method is the less sensitive to the selection of $\alpha$ and $s_{\text{max}}$. Almost always the self-consistency is reached within about 40 steps. In contrast, when the parameters are chosen poorly, for example at $\alpha = 1.6$ and $1/s_{\text{max}} = 3 \times 10^{-1}$, the conventional method requires more than 100 iteration steps depending on $k_{\text{max}}$. More importantly, finding the optimal $k_{\text{max}}$ seems to be difficult, because though the iteration steps increase with $k_{\text{max}} \leq 20$, the fastest convergence is achieved at $k_{\text{max}} = 40$. As a whole whereas the larger $s_{\text{max}}$ is desirable for the conventional method, a guiding principle for $k_{\text{max}}$ is unclear. Table 1 shows also the maximum $l$ reached within the proposed method. These values might be taken as the best $k_{\text{max}}$ for the conventional method. At $k_{\text{max}}$ near these values, however, the conventional method does not necessarily show the comparable performance of the proposed one. This is likely because discarding the oldest column is not the best strategy to keep $X_n$ and $Y_n$ from excessive growth as pointed out in the previous section.

5. Conclusion

Reformulation of the Anderson method for a system of nonlinear equations has been described. The Anderson method in practice requires two empirical parameters commanding to what extent stably the least-square problem appearing at each iteration step is solved and how many vectors containing the convergence history information are retained. In the proposed method the SVD is used to extract the effective information from the history vectors, rather than as a black-box tool for solving the least-square problem. The extracted vectors are chosen to play a role of storage space for the history information. Thereby the latter empirical parameter is no more needed. This makes the proposed method be the less sensitive to the selection of the remaining parameter and the mixing factor and the more efficient because of a smarter way of discarding a redundant part of the history information, as supported by the stable convergence in the test calculations.
References