This is the accepted manuscript made available via CHORUS. The article has been published as:

**Force field preconditioned ab initio structure relaxation method**

Liping Liu, Zhanghui Chen, Yugui Yao, and Lin-Wang Wang

Phys. Rev. B **98**, 144109 — Published 16 October 2018

DOI: [10.1103/PhysRevB.98.144109](https://doi.org/10.1103/PhysRevB.98.144109)
Force field preconditioned ab initio structure relaxation method

Liping Liu¹,², Zhanghui Chen², Yugui Yao¹, Lin-Wang Wang*²

¹School of Physics, Beijing Institute of Technology, 5 South Zhongguancun Street, Haidian District, Beijing, 100081, People’s Republic of China

²Material Sciences Division, Lawrence Berkeley National Laboratory, One Cyclotron Road, Mail Stop 50F, Berkeley, California 94720, USA

*lwwang@lbl.gov

Abstract

We present a general method to accelerate ab initio atomic structural relaxations. In this method, the conjugate gradient ab initio relaxation is preconditioned by a force field relaxation. This force field is constructed on-the-flight with the information of the ab initio forces and the current atomic configuration. At each ab initio relaxation step, the so constructed force field is used to pre-relax the system, with the relaxation direction as the preconditioned direction for the ab initio conjugated gradient method. The force field model and its parameters are rather general making this method applicable for a wide range of systems commonly used in material simulations. More than 80 different systems have been tested representing different cases in material simulations. Across the board, we find accelerations mostly over a factor of 2 and for some large systems with a factor of 6~9. There is no case where the relaxation becomes worse. The code and lib for this method are provided, which can be used as a plugin in a standard ab initio atomic relaxation code.

Main Text:

Ab initio calculations based on quantum chemistry methods or density functional theory (DFT) have played a more and more important role in material simulations at. However, ab initio simulations for large systems can be rather expensive and can take an agonizingly long time to finish. The important role of ab initio simulation in high performance computing can be illustrated from surveys of major computer center time usage, where it is found that the ab initio simulations can take about 20% of all computer time [1]. Among all ab initio simulations, atomic relaxation is one of the most widely used methods and can also be one of the most expensive calculations. For large systems, hundreds of steps could be needed for the calculation to converge beyond a typical minimum atomic force criterion of 0.01 eV/A. As ab initio methods are increasingly used for new types of simulations, e.g., the genetic algorithm search of global structure minimum[2][3][4], high throughput search for optimal material properties, the local minimum atomic relaxation often becomes the bottleneck for all these simulations. It will thus be tremendously helpful if the ab initio atomic relaxation can be sped up in some degree. Even a factor of two or three speedup could have a major impact. In this work, we present a general method to speed up the ab initio atomic relaxation with guidance from a simple, unified and general classical force field. The speed up varies from 2 to 9, and it works for a wide range of systems from small molecules to large nanosystems.

Conjugate gradient (CG)[5] and Broyden–Fletcher–Goldfarb–Shanno(BFGS)[6] are two most widely used methods to relax the atomic structure following the DFT energy. The speeds of these two methods are similar. While BFGS can be slightly faster in some cases, if the initial
atomic position of the system is far away from the minimum, the CG method could be more stable. There are other methods, like the steepest descent and the damped molecular dynamics method. They could be more stable, but in general much slower. Recently, there is also a new method called FIRE [7], which combines the damped molecular dynamics with BFGS method. Such method can be useful for large difficult problems where several thousand steps are needed [7], however for problems as the ones tested in the current paper, we found that it sensitively depends on the adjustment of the parameters [8]. With unadjusted default parameters, it could be even slower than plain CG and BFGS. This makes it difficult to use especially for automatic and high throughput calculations. In the current work, we will base our approach on the CG method, which is general, stable and relatively fast. Note that this is a relaxation method reaching the local minimum of the energy surface, not a global minimum search method.

Our approach is based on the idea that a simple Hamiltonian might be able to guide the ab initio atomic relaxation. It is well known that a good preconditioning can significantly speed up the CG. For example, for a nanosystem, a preconditioner can be constructed from the bulk dynamic matrix, which significantly accelerates the atomic relaxation [9]. However, such an approach can only be applied to nanostructures with bulk like internal degree of freedoms. In previous works, it has also been found that, a proper classical force field Hamiltonian can be used to provide either the preconditioning [10][11] or curved line minimization path [8] for the CG method with a speedup factor of 2 to 6. Unfortunately, most of these works require specific force fields and repeated on-the-flight fitting of the force field parameters [8][11]. In particular, while it works well for metallic systems with either embedded atom force field [12] or Lennard-Jones force field [13], it is difficult to be generalized to covalently bonded systems. Notice that, the commonly used CG or BFGS are agnostic regarding the physical meaning of the minimized variables (the atomic coordinates), such information (e.g., the distance between atoms) could be useful to construct approximated Hessian matrix of the system. In the current work, we present a method, which uses simple and general classical force field to provide a preconditioner for the ab initio atomic relaxation. More specifically, the ideal bond lengths and angle values for each individual bond and angle are calculated on-the-flight using the current atom position at each ab initio CG step. As a result, there is no need for their parameters, and the valence field model will have zero force. Only two generic bond strength and angle strength parameters are used. We have also combined the valence force field with embedded atom force field within a given system to deal with more complicated cases. The DFT force is used at an external force on top of the force field. This on-the-flight constructed force field potential is then used to relax the atomic position, with the resulting relaxation direction serves as the preconditioned direction of the DFT CG step. This avoids the need to calculate and invert the Hessian matrix of the force field model at every DFT CG step. The preconditioned direction is followed by the CG correction and DFT line minimization. More than 80 systems have been tested, including: molecules; bulk hetero structures; defects; surface; molecule absorbed on semiconductor and metal surface; metal clusters; as well as hybrid organic/inorganic systems. This covers most systems used in everyday ab initio atomic relaxations. The speedup ranges from a factor of 2 to a factor of 9. A simple valence force field (VFF) for the covalent bonding part of the system and a general embedded atom method (EAM) for the metallic part of the system are used. Besides the requirement for indicating which part (atoms) is covalent and which part (atoms) is metallic, there is no need to provide any other information. The code and lib are provided in the supplementary information, which can be used as a plugin in any standard DFT code. We believe the method can be widely adopted for different systems.

Our classical force field Hamiltonian consists of several terms as following:
\[ E_{\text{vff}} = E_{\text{vff}} + E_{\text{gupta}} + E_{\text{shift}} + E_{\text{delta}} \]

Here the \( E_{\text{vff}} \) is a unified valence force field (VFF) which has the following form:

\[
E_{\text{vff}} = \sum_{i=1}^{N_{\text{bond}}} k_b (b_i - b_{i0})^2 + \sum_{j=1}^{N_{\text{angle}}} k_d (\theta_j - \theta_{j0})^2
\]

where \( b_i \) is the bond length and \( \theta_j \) is the bond angle. What important is that, this \( E_{\text{vff}} \) is reconstructed at every step of the ab initio atomic relaxation \( R_0(t) \), \( b_0 \) and \( \theta_{00} \) are taken from the current bond distance and angle values from the current atomic configuration \( R_0(t) \) at this step. Note every bond and angle have their own \( b_0 \) and \( \theta_{00} \) values. These values are not type specific (e.g., different carbon-carbon bond pair can have different, \( b_0 \)). In another word, at the current step \( R_0(t) \), the \( E_{\text{vff}} \) is zero, and its force term is also zero. Nevertheless, it does provide a Hessian matrix which contains useful information for the connectivity of the atoms. We have used general parameter \( k_b \) and \( k_d \) for all bonds and angles, \( k_b = 30 \text{eV} / \text{Å}^2, k_d = 5 \text{eV} / \text{rad}^2 \), regardless of the atom types. Thus, to construct the above \( E_{\text{vff}} \), the only issue is the determination of the nearest neighbor list. We have used a bond distance cutoff to determine the bonds. This cut off is taken as the sum of the standard atomic radius, which are provided in Table.SI in the supplementary information (SI). Note, there are ionic bond systems where the valence bond topology is not well defined. Nevertheless, we found that the above \( E_{\text{vff}} \) can still be used in such cases. One can even have different number of bonds due to the use of different bond length cutoff, but the resulting speedups are similar. If two parts of a system are bonded with van der Walls force, there could be no covalent bonding between them. Their relative shift will be described by the \( E_{\text{shift}} \) term to be discussed later.

While the \( E_{\text{vff}} \) takes care of the covalent bonded, and even the ionic bonded regions, for metallic region (metal bulk/slab/cluster), one can take the advantage of the supreme quality of the embedded atom model (EAM). In particular, we have adopted the Gupta potential [14], which has a form as:

\[
E_{\text{gupta}} = \sum_{i=1}^{N} \left\{ \sum_{j \neq i}^{N} A_{ij} \exp \left( -p_{ij} \left( \frac{r_{ij}}{r_{ij}^0} - 1 \right) \right) - \sum_{j \neq i}^{N} \xi_{ij}^2 \exp \left( -2 q_{ij} \left( \frac{r_{ij}}{r_{ij}^0} - 1 \right) \right) \right\}^{1/2}
\]

Where, atom \( j \) is the neighbor of atom \( i \) within a sufficiently large cutoff, which is 2 times of the bond cutoff used in the \( E_{\text{vff}} \). \( r_{ij} \) is the first-neighbor distance in the lattice. \( A_{ij} \) and \( p_{ij} \) describe the Born-Mayer ion-ion repulsions, \( \xi_{ij} \) is the effective hopping distance, and \( q_{ij} \) describes its dependence on the relative interatomic distance. There are atomic specific parameters for most common metal elements as described in [15][16][17][18][19]. The parameters for many A-B metal atom pairs also exists, if they do not exist, an geometry average for the A, q, p, \( \xi \) parameters and an algebra average for \( r^0 \) of the two end atoms can be used. If the A-A parameter for a given element does not exist, but A1-A1, A2-A2 parameters for the two neighboring elements A1 and A2 (in the periodic table) exist, one can also take the average parameters of A1 and A2 to get an approximated parameter set for A. Some of the existing parameters are listed in Table.SII in the SI. Note that, one needs to specify, for each atom, whether it belongs to the metallic region, or nonmetallic region. The atom pairs A-B of the above GUPTA formula is included only when both A and B belong to the metallic region. While a covalent bond A-B in \( E_{\text{vff}} \) can exist when at least one atom belongs to the covalent bond region. If there is an aggregation of metallic elements in one place, we suggest assigning them as metal. However, if one metallic element is surrounded by other nonmetallic elements, as in covalent and ionic crystals, or embedded in a molecule or 2D systems, we suggest...
assigning it as non-metal. Theoretically, there could be cases where a metallic situation change to nonmetallic situation, or vice versa. Such cases are usually rare. Even when that happen, with the “wrong” assignment, the code will still work, although the acceleration might suffer. In the worse scenario when such rare cases happen, one can stop the relaxation, and reassign the metal/nonmetal types.

Besides the above two main force field terms, we have also introduced an absolute shift energy term:

\[ E_{\text{shift}} = \sum_{i=1}^{N_{\text{atom}}} k_s (R_i - R_{i0})^2 \]

Here \( R_i \) is the atomic coordinate, and \( R_{i0} \) is taken from the current atomic position \( R_0 \) (step) at this atomic relaxation step. We have used a general constant \( k_s = 1.5eV/Å \). This term prevents the shift of the system to large distance. It is like to have a diagonal term in the Hessian matrix. This is useful to fix the overall shift and rotation of isolated molecules or for cases where two parts are only connected by van der Waals interaction, which is not represented in both the \( E_{\text{vff}} \) and \( E_{\text{GUPTA}} \) potential.

Given the above three energy terms (\( E_{\text{vff}} \), \( E_{\text{GUPTA}} \), \( E_{\text{shift}} \)), one could calculate the Hessian matrix and use it for the preconditioner of the CG method. However, for large systems, the calculation of such Hessian matrix can be expensive. Besides, we like to update the Hessian matrix as the system evolves, especially if the initial configuration is far away from the local minimum. We also like to go beyond the use of linear information of the Hessian matrix. For example, for metallic cluster, when proper \( E_{\text{GUPTA}} \) is used, it is found that the \( E_{\text{GUPTA}} \) can guide the ab initio energy line minimization along a curved path, which is better than a plain use of Hessian matrix [8]. We have thus adopted this follow-the-force-field-path approach. In this approach, at each ab initio atomic relaxation step, the system will be fully relaxed based on the force field (FF) model constructed at the current step. The force field relaxed atomic positions will then be used for ab initio line minimization. This approach however, requires the force field atomic force at each ab initio step \( R_0 \) (step) to be the same as the DFT force. In our previous works [8][11], this has been achieved by explicit force field fitting at each step. However, the force field based on the above three general formalisms and parameters: \( E_{\text{vff}} + E_{\text{GUPTA}} + E_{\text{shift}} \) will not have the same atomic force as the DFT force. The \( E_{\text{vff}} \) and \( E_{\text{shift}} \) will have zero force, while the \( E_{\text{GUPTA}} \) might have a different force. To make the FF force the same as that of the DFT result, we have introduced a linear energy term:

\[ E_{\text{delta}} = - \sum_{i} \Delta F_i (R_i - R_{i0}) \]

Where, \( \Delta F_i \) is the force difference between the DFT force and the FF force (using the above three FF terms) evaluated at the current ab initio atomic relaxation step \( R_0 \) (step):

\[ \Delta F = F_{dft}^0 - F_{ff}^0 \]

The addition of this linear force correction term can make the FF minimum close to the DFT minimum. Fig. 1 shows a molecule case for the DFT minimum position, and the FF minimum position with and without the \( E_{\text{delta}} \) term.
Fig. 1 Energy curvature of DFT, force field, modified force field for the glycine system along the DFT force direction of the initial atomic configuration. This is for Glycine system, during one minimization step, x denotes the distance along the FF relaxation direction.

With the $E_{\text{FF}}$ set up at each step of the ab initio atomic relaxation (before line minimization), we can now use this $E_{\text{FF}}$ to guide our DFT line minimization search. The procedure is the following: at each ab initio atomic relaxation step $k$, its atomic coordinates $R_0(k)$ are used to construct $E_{\text{FF}}(k)$ following the above procedure. Then the atomic coordinates are relaxed following $E_{\text{FF}}(k)$ (this itself can use its own CG method, or BFGS method, it takes negligible amount of time compared to the ab initio part of the calculation). After it reached the minimum point $R_{0m}(k)$, we will take $F_p^k = R_{0m}(k) - R_0(k)$. Note, if everything is in the parabolic energy region, $R_{0m}(k) - R_0(k) = M^{-1} F_{\text{DFT}}$, where $M$ is the Hessian matrix of the $E_{\text{FF}}$. Thus our method will become a preconditioned CG method in that limit. After the $F_p^k$ is obtained, the CG feature is included by constructing the line searching direction as:

$$p_k = F_p^k + \frac{F_p^k \cdot F_k}{F_{k-1}^p \cdot F_{k-1}} p_{k-1}$$

An ab initio energy line minimization along the direction of $p_k$ will be performed. We have used Brent’s method for this line minimization [20]. In most cases, one trial step is sufficient for this line minimization step. The minimum position will be used as the starting point for step $k+1$: $R_0(k+1) = R_0(k) + \alpha p_k$, here $\alpha$ is the step size for this line minimization. The flow chart of the whole PCG procedure is shown below in Fig. 2.
We have tested 82 systems, all the calculations are done using the PWmat code [21][22], which is a GPU code for plane wave pseudopotential DFT calculations. SG15[23][24] norm conserving pseudopotentials are used. The typical plane wave energy cut-off is 50Ryd.

All the 82 systems have reached the DFT force criterion. However, there are a few cases where the PCG found minimum is slightly different from the CG found minimum, actually all with slightly lower energies (e.g., within 0.06 eV or 0.002eV). This is more relevant to the cases where the reported speedup is close to 1. In a way, this is not really the fault of PCG or CG. When there are multiple minima surrounding the initial starting atomic configuration, there is no guarantee different relaxation methods (CG, PCG, BFGS, steepest decent, FIRE etc.) will reach the same local minimum. One can indeed define a potential basin where the initial atomic position belongs. However, to guarantee to reach the bottom of that basin, steepest decent with very small time-steps (no line minimization) need to be used, which is hardly a practice method for atomic relaxation. The relaxation speedups for these 82 systems are shown in Fig. 3, along with the relaxation steps needed to converge the system when the preconditioner is not used. These systems have 5 to 216 atoms with various system types. Organic molecules include carbon chains, amino acids, and benzene rings. Metallic systems include copper cluster, silver cluster, copper surface, nickel-aluminide alloy. Semiconductors include gallium arsenide with
vacancies, diamond silicon with vacancies, GaAs-InAs heterostructure. 2D materials include silicene supercell and molecules absorbed on silicene; Ionic systems include sodium chloride supercell. All the systems, including their numbers of atoms are listed in Table. SIII. We have measured the speedup in two different ways. One is the number of steps needed to reach the 0.01eV/A force criterion, another is the number of steps needed for the ΔE (the total energy minus the final minimum energy) to drop to one half of its original values. The initial atomic positions of these systems are prepared either by universal force field (UFF) [26] relaxation (for the molecular system), or at their ideal crystal positions (for defect, heterostructure), or simply place a molecule on top of a surface. These are the typical initial positions people used in their material simulations. For most of the 82 systems, we have a speedup around 2.0, and some of them have a high accelerating factor up to 9. Without using the acceleration, most of these systems take in the order of 100 to 200 steps to reach the minimum, while some systems take about a 1000 steps. Thus a factor of 2-3 can mean a significant saving of time.

![Speedup](image)

Fig. 3 The speedup factors for different systems. (a) the speed up measured by the force criterion; (b) the speed up measured by reaching the half energy drop; (c) the CG steps to reach the force criterion without the use of the acceleration method. The types of systems are color coded in (a) and (b).

In the following, we describe these systems in more details. As listed in Table.SIII, simple carbon chains and 20 types of common amino acids are tested. The carbon chains have numbers of atoms ranging from 5 to 62. The amino acids contain common organic atomic types, like oxygen, nitrogen, sulfur, as well as different bonding topologies: single bonds, double bonds and rings. The number of atoms for the amino acids ranges from 10 to 26. The initial configurations are obtained by relaxation using Avogadro 1.2 [25] with Universal Force Field

Error! Reference source not found. Reference source not found.. Fig. 4 shows the relaxation of four amino acids cases. CG cannot converge them in 100 steps, BFGS method gives a slightly better result for some systems, and the new method converge the system in 40-60 steps with an accelerating factor of
2-4 relative to the CG method.

Fig. 4 First 100 relaxation steps for four amino acids for the three methods: CG (conjugated gradient), BFGS, and PCG (the current preconditioned conjugated gradient method). The vertical axis is ΔE = E(step) - E_final in eV, the horizontal axis is the CG steps.

We have tested metallic systems, including clusters, metallic heterojunctions, and metal surface with adsorbed molecule. As discussed before, EAM potential is used to describe the metallic part of the system in our accelerating scheme. Gupta EAM potential is relatively close to DFT potential energy surface, this leads to a good acceleration for the PCG method. We tested the metallic clusters Ag_{55}, Ag_{16}, 128 atom Ni_{3}Al_{Ni} metallic alloy, as well as Cu surface with absorbed molecules containing 122 atoms. These are real systems used for cluster studies, or metallic catalysis. The speedups for these systems are around 2-4.

It is also interesting to test the effects of our method for defects within semiconductors or strained heterojunction systems. To do this, we have taken the bulk GaAs and Si supercells containing 216 atoms, and remove some atoms to form vacancies. The initial structure is the ideal atomic positions of the perfect crystal. We also constructed a CdS-CdSe quantum well as well as a GaAs-InAs quantum well containing 64 atoms with the perfect crystal position using the average lattice constants as our initial atomic configurations. These are the typical mechanical problems for strain relaxation. The PCG method gives a 2-3 speedup for semiconductors with point defects and a speedup factor of 3-6 for the relaxation of the strained systems. These are typical problems encountered in defect calculations and nanostructure calculations. As a matter of fact, larger the system, higher speedup is expected in such bulk like systems. This is because for large systems, the soft acoustic phonon modes become lower in energy, which worsen the condition number of the Hessian matrix, and make the relaxation converges slower. However, our force field model describes well such mechanical acoustic modes, especially for their relationship to their corresponding wave numbers. As a result, the approximated Hessian matrix from the force field model can correct the soft modes through the preconditioning, and restore the speed of the atomic relaxation. To demonstrate this fact systematically, we have now provided the speed up of a semiconductor bar with increasing bar length in Fig.5. As one can see, the speed up increases with the bar length.
Fig. 5 GaAs-InAs hetero-structures. (a) Speedup factor of different sizes of systems. We construct GaAs supercells from 2x1x1 to 6x1x1, replace half or less half of the Ga atoms to In atoms, and randomly move all the atoms by a random shift less than 0.1 Angstrom. (b) Typical energy curve in main accelerate region (6x1x1)

We also tested 3 organic/inorganic hybrid perovskite systems. Such systems (e.g., the methylammonium lead iodide, MAPbI₃) have been intensely studied in recent years for solar cell research. However, due to the existence of the organic molecule and their relative weak interaction with the inorganic framework, the atomic relaxations of such systems are quite challenging. In our cases, with about 100 atoms, the original CG method can only converge the systems after 800-1000 steps. Our PCG method can converges the systems within 400 steps.

Some of the actual relaxation curves of the above-discussed problems are shown in Fig. 6.
Fig. 6 Relaxation curves of different systems: (a) Ag16 cluster; (b) Ni3AlNi alloy; (c) Molecule C4H4N2S2 on Cu surface; (d) Point defect in bulk Si; (e) GaAs-InAs heterostructure; (f) MAPbS4 hybrid perovskite supercell. The horizontal axis is the CG steps.

Finally, we have tested a few surface-molecule adsorption cases. A 7x7 2D silicene adsorbing one H2O molecule (total 101 atoms); a 7x7 2D silicene adsorbing one SO2 molecule (total 101 atoms); 6x6 2D molybdenum disulfide adsorbing one NH3 molecule (total 112 atoms); and PbS (001) and (111) surfaces adsorbing one oleic acids molecule ((001) 85 atoms, (111) 105 atoms). The conventional CG method typically has very slow convergences for these systems due to the possible rotation and other soft configuration changes of the molecule on the substrate. The first three cases give speedup factors around 3-6. In the case of PbS adsorbing oleic acids cases, our PCG method converges in less than 300 steps, while the conventional CG method cannot converge after 1500 steps. The speedup at half energy is about a factor of 3. A few examples of the actual relaxation curves are shown in Fig. 7.
In summary, we have presented a force field guided DFT PCG method to speed up ab initio atomic relaxations. In this method, a simple force field model is constructed at every step of the ab initio atomic relaxation. Such a simple force field model allows the application of the current method to all common material simulation problems, provides accelerations to most of the cases we have tested. The force field is constructed in such a way that its atomic forces are the same as that of the DFT forces at each step. The system is then relaxed following the energy of the constructed force field (with negligible amount of computational time). The relaxed position is used as the preconditioning corrected gradient direction for the DFT PCG method. 82 different systems have been tested, covering a wide range of material simulation problems. Across the board, we found the speedup typically in the range of 2-6. Since ab initio atomic relaxation takes a major chunk of the computational time for material simulations, we expect the current method can be widely adopted in many studies. The code and lib for this method are provided in the GitHub [27], which can be used as a plugin in a standard ab initio atomic relaxation code. In our future work, an important direction is to develop an efficient automatic method to distinguish the local types of a system in a given region. Another is to determine whether we can consider Coulomb interaction and van de Walls interaction. More systems also need to be tested to fully confirm and understand the potential and limitation of this method. These will provide better force field parameters to make the PCG method more robust and efficient.

Acknowledgements
This work is supported by the Director, Office of Science (SC), Basic Energy Science (BES), Materials Science and Engineering Division (MSED), of the US Department of Energy (DOE) under Contract No.DE-AC02-05CH11231 through the Materials Theory program (KC2301). This work is also financially supported by the MOST Project of China (Grants No. 2014CB920903), the NSF of China (Grants Nos. 11734003, 11574029), and International Graduate Exchange Program of Beijing Institute of Technology. We use the resource of National Energy Research Scientific Computing center (NERSC) located in Lawrence Berkeley National Laboratory.

[27] See the static lib at https://github.com/llpjaj/DFT_PCG_atomic_relaxation.
See Supplemental Material at [URL] for the atomic radius and details of tests.