Ab initio current-induced molecular dynamics

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We extend the ab initio molecular dynamics (AIMD) method based on density functional theory to the nonequilibrium situation where an electronic current is present in the electronic system. The dynamics is treated using the semi-classical generalized Langevin equation. We demonstrate how the full anharmonic description of the inter-atomic forces is important in order to understand the current-induced heating and the energy distribution both in frequency and in real space.

Energy dissipation from an electronic current into the motion of atoms is an important and challenging scientific and technological problem.1,2 Especially, conductors consisting of a few atoms can be driven into a highly nonequilibrium vibrational state by a current.3,4 The local excitation of the atomic vibrations is hard to address in experiments5-10 and theoretically11-14. So far the formation of atomic “hot-spots” has been demonstrated using first principles calculations at the level of density functional theory (DFT) using the harmonic approximation thus neglecting inter-mode coupling.15 It was also shown how current-induced momentum transfer (”electron wind” forces) may displace the “hot-spots” up or down-stream w.r.t. the current. This behavior has been observed in recent experiments16-18. However, the fact that individual vibrational modes couple differently to the electronic current and phonon reservoirs makes this picture invalid in general. Moreover, the anharmonic inter-mode coupling distributes the energy among these and makes the problem challenging to treat beyond model calculations.

Here we present a computational method which allows for ab initio molecular dynamics (AIMD) simulation of atomic scale conductors in the presence of electronic current. It can address the effects of the current on the vibrational excitation (Joule heating), the current-induced ”wind” forces, the coupling to electrode phonon reservoirs, and the anharmonic couplings between the modes of the conductor. We demonstrate the method on two ”bench-mark” atomic scale conductors, an atomic gold chain and a benzene molecule, and analyze the role of the individual effects for the dissipation.

Langevin equation. – We consider an atomic scale conductor consisting of a central bottleneck junction and two electrodes, ”left” and ”right”, respectively. In the presence of an electrical current, electrons are scattered near the junction and deposit energy to the surrounding atoms. We are interested in the atomic dynamics driven by the electrical current across the junction. To this end, we define the ”system” as the coordinates of the nuclei in the junction. All remaining degrees of freedom (DOF) are treated as reservoirs/baths. We consider two kinds of reservoirs. The first is the electronic one, including all the electron DOF in the conductor and electrodes. Importantly, in the presence of electrical current, the electron reservoir is in a nonequilibrium state. This non-thermal reservoir serves as the energy source, depositing energy to the atoms. The second kind is the phonon reservoirs of the left and right electrode parts.

Employing the Feynman-Vernon influence functional approach, a semi-classical generalized Langevin equation (SCGLE) can be derived to describe the dynamics of the nonequilibrium system.14,21,29

\[ \ddot{Q}(t) = F(Q(t)) + \int_{-\infty}^{t} II'(t' - t) \dot{Q}(t') dt' + f(t). \] (1)

Here, \( \dot{Q}(t) \) is the mass-normalized displacement of system nuclei away from their equilibrium positions, \( F(Q(t)) \) is the forces acting on the system, \( \dot{Q}(t) \) is the mass-normalized velocity, and \( f(t) \) is the fluctuating/stochastic force with zero mean. It can be characterized by the correlation function,

\[ \langle f(t)f(t') \rangle = \hbar \Pi(t - t'). \] (2)
which includes thermal and quantum fluctuations. Both $\Pi$ and $\Pi'$ are given by a sum of contributions, i.e., $\Pi' = \Pi'_e + \Pi'_r$, with $e$, $L$ and $R$ representing the electron, the left and the right phonon reservoirs, respectively.

In nonequilibrium $\Pi'$ and $\Pi$ can be obtained using nonequilibrium Green’s functions.

3.1. Results

Previously Eq. (1) has been used for ab initio calculations in the harmonic approximation. Here, we go beyond this limitation and perform AIMD simulations by numerically solving Eq. (1), with all parameters obtained from the same DFT setup. In the following we consider two prototype atomic conductors and illustrate the importance of anharmonic vibrational coupling on the energy redistribution both in frequency and in real space.

Au chain – The first example is a 5-atom gold chain between two Au(100) electrodes [inset of Fig. 1(a)]. The temperature of the electrodes are 4K. In the harmonic approximation, Eq. (1) is linear and can be solved directly. We first check that the MD result in the harmonic approximation reproduces that from direct solution of Eq. (1) (Fig. S1). After validating the method, we proceed to study energy transfer from the nonequilibrium electronic bath to the system when the applied voltage bias is finite, $V \neq 0$. There are two energy transfer channels: The first is the stochastic Joule heating from the bias dependent fluctuating force correlation function $\Pi_e$. The second is the work done to the system through the energy non-conserving (NC) current-induced forces in $\Pi'_e$. The energy deposited in the system can be further transferred into the left/right equilibrium phonon reservoirs. If the system remains stable it reaches a steady state when the energy into the system is balanced with the energy out to the phonon reservoirs.

After reaching a steady state, we use the power spectrum of the velocity-velocity correlation function $C_{vv}$ (Eq. S1) to characterize the frequency/energy-resolved contribution to the system kinetic energy. Integrating $C_{vv}$ yields the kinetic energy stored in the system for frequencies below $\omega$,

$$E_K(\omega) = \int_0^\omega \frac{d\omega'}{2\pi} C_{vv}(\omega').$$

The calculated results for $C_{vv}$ along with $E_K$ are shown in Fig. 1(a) for the harmonic calculation using the dynamical matrix from DFT, and in Fig. 1(b) for the full AIMD calculation. Applying a bias voltage results in an increase in vibrational kinetic energy, which can be divided into separate heat and work contributions, shown in Fig. 1(c) and (d), respectively. The work part comes from the contribution of NC current-induced forces, while the heat part is from Joule heating.

Comparing results at $V = 0$V (equilibrium) and $V = 1$V (nonequilibrium) from the harmonic approximation, we find that electrons transfer energy mainly to the high frequency modes at $\sim 0.02$eV. This is indicated by a big step at $\sim 0.02$eV at 1V [red line in lower panel of Fig. 1(a)]. These modes are the alternating-bond-length (ABL) modes consistent with previous experimental and theoretical studies. In addition to stronger coupling to electrons, the ABL frequencies are located above the bulk phonon band of Au. This hinders direct harmonic energy transfer to the phonon reservoirs. Inclusion of the anharmonic coupling leads to an energy redistribution from the high to the low frequency modes.
which is absent in the harmonic case. The anharmonic coupling also reduces the excess kinetic energy in the system by opening an indirect channel of energy transfer from the ABL modes to the bulk electrode phonon reservoirs [Fig. 1(c-d)].

We now analyze the MD results in real space to gain further understanding of the bond properties. In Fig. 2(a) the kinetic energy per bond [Eq. S6] is shown for the harmonic and anharmonic cases. In each case, we compare the situations with and without work contribution from non-conservative (NC) current-induced forces. The inclusion of NC current-induced forces leads to a left-right symmetry breaking in the real space energy distribution. Specifically, more energy is accumulated in bonds near the left electrode from where the current enters the junction. This asymmetric concentration of energy ("hot spot") was explained by the asymmetric excitation of left- and right-traveling phonon waves by NC current-induced forces[15].

Due to this effect, some bonds in the chain heat up strongly. We have compared the statistics of bond fluctuations between atoms 2-3 and 4-5 in both the harmonic and full AIMD calculations [Fig. 2(b)]. The 2-3 bond shows larger broadening compared to the 4-5 bond in both cases. This is a direct result of asymmetric heating. In the AIMD calculation, the broadening is reduced compared to the harmonic result. This is due to the anharmonic energy transport channel to the phonon reservoirs, consistent with the analysis in Fig. 1. Moreover, we observe elongation of both bonds in the AIMD result. This is another manifestation of anharmonicity[28].

**Benzene molecular junction** – The second example we consider is a Benzene molecular junction connecting to Au(100) electrodes at 4K through direct Au-C bonds[29] [inset of Fig. 3(a)]. Due to different atomic mass and bonding type, there is a large frequency mismatch between Benzene vibrational modes (0−380 meV) and the gold electrode phonons (0−18 meV). Only low frequency vibrational modes (< 18 meV) couple directly to electrode phonons. The remaining high frequency modes (> 18 meV) appear as sharp peaks in the power spectrum [Fig. 3(a)]. Work done by current-induced forces in this junction is negligible due to low electrical conductance. The energy increase at 1 V is dominated by Joule heating. It involves mainly vibrational modes larger than 75 meV. Including the anharmonic vibrational coupling leads to large spectrum broadening [Fig. 3(b)], and an excess heating much smaller than the harmonic case [Fig. 3(c)].
Similar to the Au chain, we observe heating of the vibrational modes below 75 meV. This is a direct result of anharmonic vibrational coupling between different modes.

The real space distribution of kinetic energy per bond at 1 V is shown in Fig. 4(a). It is symmetric due to the symmetry of the junction and the negligible contribution of NC current-induced forces. The bond length histograms for the different types of bonds in the junction are depicted in Fig. 4(b) for harmonic (lower panel) and full AIMD (upper panel) calculations. Comparing the two cases, we find that, the excess energy of C-H and C-C bonds is effectively transferred away to the Au-C bond in the presence of anharmonic coupling. This is evidenced from the reduced broadening of C-C and C-H bonds and the enhanced broadening of Au-C bond in the full AIMD results. As a result, the average Au-C bond length increases by \( \sim 0.2 \) Å. More interestingly, the Au-C bond distribution shows obvious skewness, thus the average length is larger than the most probable length. This is an important signature of bond anharmonicity, and can only be probed by the full AIMD calculation.

![Fig. 4](image)

**FIG. 4.** (a) Comparison of kinetic energy per bond in meV from the full AIMD and the harmonic calculation. (b) Length distribution of different bonds calculated from the two methods. In the full AIMD results, the Au-C bond exhibits the widest distribution with a maximum bond elongation of \( \Delta b = 0.2 \) Å and the largest shift of the mean bond length with bias. Interestingly, it also shows certain skewness with larger probability to longer length side.

In experiments an effective temperature, \( T_{\text{eff}} \), has been obtained from contact life-times \([30]\) or two-level conductance fluctuations \([31]\) via an Arrhenius type rate expression. The \( T_{\text{eff}} \) has been fitted to a \( V \)-dependence\([4]\), as \( T_{\text{eff}} = \gamma \sqrt{V} \) (zero ambient temperature). For both systems above we do not find the energy distribution among the modes is compatible with an overall effective temperature (cf. Fig. 5). On the other hand we find a maximum kinetic energy of interatomic Au-Au (Au-C) bonds corresponding to 80 K (320 K) which is comparable to experimental values\([30, 31]\) of 75 K (450 K).

In summary, we have presented a method for performing \textit{ab initio} molecular dynamics of an atomic scale conductor in the presence of an electrical current. We illustrated this approach by studying two prototype molecular conductors and analyzing the subtle interplay between Joule heating, current-induced forces and anharmonic vibrational coupling among different modes. We found that anharmonic coupling among different modes has an important effect on the energy redistribution both in frequency and in real space. Our study opens the possibility of studying current-induced atomic switches, chemical reactions and bond ruptures\([32, 33]\) beyond simple models.

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Supplementary Materials for “Ab initio current-induced molecular dynamics”

A. Velocity correlation and power spectrum

To study the energy distribution of the system at steady state, we define the velocity correlation function as

$$C_{v_i v_j}(t) = \frac{1}{T} \int_0^T dt' v_i(t + t') v_j(t'). \quad (S1)$$

Here, $T$ is the total time of one MD trajectory, and $v(t) = \dot{Q}(t)$ the mass-normalized velocity. Using the Wiener-Khinchin theorem, the power spectrum is then given by the Fourier transform of the velocity correlation function

$$\frac{1}{T} \langle |v(\omega)|^2 \rangle = C_{vv}(\omega) = \sum_i C_{v_i v_i}(\omega). \quad (S2)$$

with

$$C_{v_i v_i}(\omega) = \int dt \, C_{v_i v_i}(t)e^{i\omega t}. \quad (S3)$$

The cumulative integration of the power spectrum

$$E_K(\omega) = \int_0^\omega \frac{d\omega'}{2\pi} C_{vv}(\omega'), \quad (S4)$$

gives the kinetic energy contributed by modes with frequency $\leq \omega$. Thus, $E_K(+\infty)$ gives the total kinetic energy. To characterize the real space distribution, we define the velocity of the bond between atoms $n$ and $k$ as

$$\bar{v}_{nk}^\alpha = v_n^\alpha - v_k^\alpha, \quad (S5)$$

where $\alpha = x, y, z$. The kinetic energy of the bond is then

$$E_{b,nk}^{\alpha} = \int_0^{+\infty} \frac{d\omega}{2\pi} \sum_\alpha \bar{C}_{\bar{v}_{nk}^\alpha \bar{v}_{nk}^\alpha}(\omega). \quad (S6)$$

B. Validation of the method

To validate our method, we show in Fig. S1 the comparison between MD and theoretical results at zero bias $V = 0$ within the harmonic approximation where the SCGLE can be solved exactly. We find good agreement between the MD (black) and theoretical (green) results.

C. Effective temperature

We estimate the effective temperature $T_{\text{eff}}$ from

$$\frac{C_{vv}(\omega, V)}{C_{vv}(\omega, 0)} = \frac{2n_B(\omega, T_{\text{eff}}) + 1}{2n_B(\omega, T_0) + 1}$$

with

$$n_B(\omega, T_{\text{eff}}) = \frac{1}{e^{\frac{\omega}{k_B T_{\text{eff}}}} - 1}$$

the Bose-Einstein distribution, $k_B$ Boltzmann’s constant, $\omega$ the phonon frequency, $T_0$ the temperature of the MD simulation and $V$ the applied bias voltage. The results for Au chain and benzene molecule are shown in Fig. S2 (a) and (b), respectively.
FIG. S1. The power spectrum and its cumulative integral for the 5-atom gold chain within the harmonic approximation at 0 V. The green line is the exact theoretical result from solving the SCGLE, while the noisy black lines are results of the MD simulation. They agree well with each other. The blue vertical lines mark the energy of different vibrational modes for the isolated chain. The red line is the cumulative integral of the power spectrum.

FIG. S2. Effective temperature over frequencies for the Au chain (left) and the Benzene molecule (right) for $T_0 = 4$ K, $V = 1.0$ V.

D. Computational details

The electronic structure of the relaxed structures is calculated using SIESTA[34] and TRANSIESTA[35]. The vibrational spectrum and the electron-vibration coupling of the system are calculated using Inelastica[27]. The electrode phonon self-energy is calculated using the method described in Ref. 38. From these, all parameters of the Langevin equation can be calculated. The details of the theory can be found in Ref. 22.

In the harmonic approximation, the dynamical matrix of the central system is used to calculate the potential force $F(Q(t))$ in Eq. (1), while for the full AIMD the anharmonic force is obtained from the *ab initio* code SIESTA. Specifically, to solve Eq. (1), we are using SIESTA in combination with the i-PI code[36], which is a python-based force engine for atomistic simulations using interatomic forces and potential energies calculated by an external driver code. This means that i-PI deals with the propagation of the nuclear motion, while the external driver SIESTA is used to compute the energy and forces. Communication between the client and i-PI runs through a socket interface. SIESTA receives the atomic coordinates from the python interface and returns forces $F(Q(t))$ that are used to integrate the Langevin equation, Eq. (1), which we have implemented in the integrator class of i-PI. For the MD in the harmonic
approximation, our Langevin-i-PI code communicates with an external driver that calculates the forces from the dynamical matrix.

In the SIESTA calculations, a single zeta polarized basis set is used for Au, and double zeta polarized basis is used for C, H. A mesh cut off of 100 Ry is used to accelerate the MD simulation. A comparison of the IETS of the Au chain using a cut off of 250 Ry to 100 Ry is depicted in Fig. S3, showing only slight influence of the cut off. A time step of 10 fs (1 fs) and total of 8192 (131072) MD steps are used for Au chain (benzene junction) for each independent MD run. The results shown in the main text are average over 10 (4) independent MD runs.

![IETS spectrum for Au chain using different mesh cut offs. They agree reasonably well with each other. To save the computational time, we have used a mesh cut off of 100 Ry in the MD simulation.](image)

FIG. S3. IETS spectrum for Au chain using different mesh cut offs. They agree reasonably well with each other. To save the computational time, we have used a mesh cut off of 100 Ry in the MD simulation.