Direct Generation of Electric Currents from Flowing Neutral Ionic Solutions

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We have discovered a new method of generating electric currents, directly from high pressure-induced flow of neutral ionic solutions. The mechanism is that the cations and anions have different flow velocities, if their atomic masses are dramatically different, due to different accelerations generated from the high applied pressure. The generated electric current is very sensitive to the strengths of the applied pressure, and it might be potentially used for detection of atomic masses and pressures.

Production and storage of electricity are among the most important activities in modern industrial processes. Conventional methods of generating electric current are electromagnetic induction, photovoltaic [1], thermoelectric [2], piezoelectric [3], electrokinetic generation [4], and so on. Recently, nanoscale electric devices have been developed extensively, such as nanoscale thermoelectric [5–7] and piezoelectric [8–10] devices. Nanoscale electrokinetic generation of electric current has also had a great progress, both in theories [11–13] and experiments [14–17]. Other methods of generating electric currents at the nanoscale have also been proposed, such as using electron tunnelling phenomena [18]. These methods of current generation usually require usage of solid materials with special electric response to external light, thermal and mechanical excitations, or require applying strong external electric fields. Therefore, the mechanisms of current generation often involve multisteps of processes, and the practical instruments for these methods might be also complicated.

In this work, we have discovered that electric currents can be directly generated from fast-flowing neutral ionic solutions under external pressures, using molecular dynamics (MD) simulations. The current is the result of dramatically different atomic masses, and hence very different accelerations, of the counterions in the flowing ionic solutions. This method represents a new type of one-step electric current generation, with possible applications in convenient electric energy storage, and highly sensitive detection of very large pressures.

In our MD simulations using the NAMD software package [19] and the CHARMM27 force field [20], a pair of counterions and 7948 water molecules were initially positioned inside a $6.2 \times 6.2 \times 6.2$ nm simulation box, with periodic boundary conditions applied, as shown in Figure 1. The water molecules were simulated using the TIP3P model [21]. The cation was N-heptyl-pyridine, and the anion was chloride. The partial charges of the atoms in the cation were calculated by ab initio methods, and the force field parameters were estimated similar to those in [12]. Pressure-induced flows of the ionic solution were generated by applying an external force of 0.02 kcal/(mol\( \cdot \)\AA) to all the nonhydrogen atoms in the region of $-0.5 \text{ nm} < x < 0.5 \text{ nm}$, in the positive $x$ direction. Given the number of water molecules and ions in the simulation box, this generates a pressure of $P = 500 \text{ atm}$ [11]. Langevin dynamics were performed, with 0.1 ps$^{-1}$ damping coefficient, which might be correct for polar groups and water, at $T = 300 \text{ K}$ [7]. The induced flow velocity of water molecules reached a constant value of around 85 m/s. When the atomic mass of the anion, $M_a$, was much larger than the atomic mass of any atom in the cation, the anion’s flow velocity became smaller than the cation, because of its smaller acceleration, supposing that the anion and cation were under the same external pressure, and the surface areas of
the nonhydrogen atoms in the cation and the anion were not very different. Therefore, as confirmed by our simulations, this method of current generation mainly depends on the atomic masses of the counterions, but does not qualitatively depend on the chemical structures or signs of charges of the counterions. This method could be applied similarly to the cases when the counterions are very large, such as colloids, but those are not the focus of this work.

We have estimated the influence of the anion’s atomic mass \( M \) on its flow velocity. In the simulations, we only changed the atomic mass of the anion from 35.5 Dalton, to 62.5, 125, 200, and 250 Dalton and left all the other parameters unchanged. The velocities of the cation were very close to the average velocities of the water molecules, since the atomic masses of the atoms in the cation were close to those of the atoms in water molecules. In Figure 2(a), we plot the flow velocities of the cation and the anion, depending on different values of \( M \). We can see that in general, the cation flowed faster than the anion, and the current was noticeable. The current first decreased and later increased with \( M \). The results were obtained from the simulations of pressurizing the solution for \( t \approx 20 \) ns, during which the ions pass the periodic boundary by \( \approx 300 \) times. The flow velocities of the water molecules and the counterions remained almost constant during the flow of the solutions. In Figure 3, we plot the distance travelled by the chloride ion \( L \), when \( M = 35.5 \) Dalton, depending on the length of the simulation. From the figure, we can see that the flow velocity of the chloride ion remained almost constant within about 4 ns of simulation. Please note that at the nanoscale, small molecules are subject to fluid pressure similar to single atoms. Although there are chemical bonds between atoms in a molecule, the fluid pressure will be distributed everywhere, including the internal region of molecules, so the atomic mass has the major influence on the flow velocity of ions, and chemical structures of ions are less important.

To study the generation of electric current in a nanoscale confinement, we also simulated the flow of the ionic solution with the same counterions and 6659 water molecules, inside a 6.2 nm long (76,0) carbon nanotube (diameter of 6.03 nm).

The system was also positioned inside a 6.2 × 6.2 × 6.2 nm simulation box, with periodic boundary conditions applied. We also only changed the values of \( M \) and plotted the velocities of the cation and the anion depending on the values of \( M \) in Figure 2(b). We can see that the cation also flowed faster than the anion, and the current first increased and later decreased with \( M \). We only simulated the condition when the N-heptyl-pyridine cation was always totally solvated and surrounded by water molecules, away from the carbon nanotube surface. Because the cation has a hexagonal ring and a hydrophobic tail, it might attach to the hydrophobic

**Figure 1:** Initial configuration of the simulated system. Two counterions were located randomly in the water box. The cation was nonprotonated N-heptyl-pyridine, and the anion was chloride ion (in green).

**Figure 2:** The flow velocities of the cation and anion depending on the different atomic masses of the anion, at an external pressure of \( P \approx 500 \) atm and at temperature \( T = 300 \) K. (a) Flow velocities of the cation and the anion in bulk solution. (b) Flow velocities of the cation and the anion in a (76,0) carbon nanotube.

**Figure 3:** The distance \( L \) travelled by the chloride ion (\( M = 35.5 \) Dalton), depending on the length of the simulation \( t \), at an external pressure of \( P \approx 500 \) atm and at temperature of \( T = 300 \) K.
The method of detection inside ionic solutions could be very sensitive to the atomic mass of the anion, as well as the external applied pressures. The induced electric current might also be very sensitive to the system's temperature, and we plan to investigate this in the future.

In summary, we have discovered a method to directly generate electric currents from fast-flowing neutral ionic solutions. This method is very convenient and has a simple mechanism. It could have potential applications in power generation and storage, as well as highly sensitive nanoscale detection of pressures and atomic masses [22–26]. It also provides a new possibility for separating mixed heavy element nuclear materials into different pure heavy elements.

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