Transport properties and efficiency of elastically coupled Brownian motors

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As models for biological molecular motors, Brownian motors have been studied recently by many workers, and their physical properties such as velocity, efficiency, and so on, have been investigated. They have also attracted much interest in an application to nanoscale technology. It is significant to study more complex systems, that is, coupled Brownian motors, in detail, since Brownian motors with a single particle have been mainly studied until now. In this paper, we consider Brownian motors coupled mutually with elastic springs, and investigate the dynamics of the model and the efficiency of energy conversion. In particular, we find that the center of the mass of the elastically coupled particles moves faster than the corresponding single-particle model, and also that the efficiency of the coupled-particle model is larger than that of the single-particle model.

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I. INTRODUCTION

Brownian motors have attracted much attention as models of molecular motors [1] and their physical properties such as velocity, efficiency, and so on have been studied in detail [2]. Recently, some workers have also paid attention to them in an application to nanoscale technology [3]. For example, Porto et al. [3] studied microscopic engines on the atomic scale that transform the fed energy into directed motion through a dynamical competition between the intrinsic length of the moving object and supporting carrier.

The molecular motors are involved in cell locomotion, some cellular transport and muscle contraction, and so forth. Many models have been proposed to comprehend theoretically the mechanism of the molecular motors. Doering et al. [4] investigated “single-particle” Brownian motor models, a “rocking” ratchet model where a periodic or random external force is applied to the system. A famous one among the single-particle models is a so-called “flashing” ratchet model [5]. It is shown that in the model, only thermal noise and a proper asymmetric potential are enough to produce macroscopic motion of the particle toward a particular direction that depends on the asymmetry of the potential. Next, “coupled-particle” models, where particles interact mutually, have been investigated. Csaหók et al. [6] studied the dynamics of elastically coupled particles in a “rocking” ratchet model. On the other hand, Jülicher et al. [2] introduced and analyzed theoretically particles rigidly attached to a rigid backbone with equal spacing in a “flashing” ratchet. Recently, Elston and Peskin [7] investigated the characteristic of the elasticity between the motor and its cargo and showed that the elasticity allowed the motor to run faster than if they were linked rigidly. Klumpp et al. [8] studied the two harmonically coupled particles in the ratchet model and observed a driving mechanism different from the one in the case of a single particle, which does not need diffusion. As for the response to an external force, Reimann et al. [9] introduced another model of interacting Brownian particles and found some collective phenomena.

Studying the physical properties of coupled Brownian motors is interesting in itself. Moreover, it is quite significant to investigate the coupled systems, since in an application to nanoscale technology, coupled effects should be taken into account [3]. In an application to molecular motors, it is also interesting to study coupling effects, because, for example, molecular motors in muscle have a linear structure [1] that consists of many parts.

In this paper, we consider elastically coupled particles in a flashing ratchet model, in which each particle makes transition repeatedly between two states where interactions are expressed by their respective kinds of potential, and conduct numerical simulation. Unidirectional motion of the particles is confirmed in this model and the velocity for various values of the coupling constant, the temperature, and so on, is measured. Although an isolated single particle in flashing ratchet models cannot move in nonthermal conditions, that is, without thermal noise, elastically coupled particles in the flashing ratchet model may move due to the action of the interparticle springs restored to their natural length that is incommensurable with the period of the periodic potential, even if thermal diffusion is not allowed. It is also found that the velocity of the elastically coupled model under such conditions is larger than that of the corresponding single-particle model, and that the velocity has a maximum as a function of the coupling constant of the springs. Moreover, if we restrict the region where transition may be allowed, we find that the velocity of the model is enhanced by the restriction. We also apply various loads externally to our system to investigate the efficiency of energy conversion, and find that the efficiency has a peak as a function of the load. We also find that the peak values are larger when we restrict the region of transitions and are improved by the coupling effect in comparison with the single-particle model.

II. THE MODEL

We consider elastically coupled particles (Fig. 1). In this paper, our model is described by dimensionless quantity. It is assumed that the particles are put in a heat bath represented by white noise. Particles are subjected to one of the two substrate potential stochastically. $W_j(x)$ $(j=1,2)$ defines the potential in state $j$ at point $x$. $W_1$ is a flat potential and we...
choose the following asymmetric interaction potential $W_2$:

$$W_2 = \left[ \frac{1}{2} \sin\left(\frac{2\pi x}{L}\right) + \frac{1}{8} \sin\left(\frac{4\pi x}{L}\right) \right] \times U,$$

where $U$ and $L$ represent the depth and period of the potential, respectively. In state one, no force from the substrate is exerted on the particles because the substrate potential $W_1$ is flat. Therefore, state one is called the detached state. Since, in state two, the particles feel periodic substrate potential, state two is called the attached state.

The equations of motion of the particles read

$$\gamma \ddot{x}_i = k(x_{i+1} - 2x_i + x_{i-1}) - h_i(t) \frac{\partial W_j(x_i)}{\partial x_i} + \sqrt{2D} \xi_i(t),$$

where $x_i$ denotes the position of the $i$th particle and we consider the overdumped case. $\xi_i(t)$ denotes white noise of zero

![FIG. 1. In our simulation, the period of the potential is not equal to the natural length of the springs. This “incommensuration” leads to the easy movement of the particles to a particular direction (to the right direction in this figure). Some particles are subjected to $W_1$ and the others to $W_2$.](image)

![FIG. 2. $x_1$, $x_N$, and the center of mass as a function of time. We find that finite temperature is not necessarily needed for the finite velocity of the elastically coupled particles in an asymmetric potential. This figure shows finite velocity for zero temperature ($k = 4.0, N = 20$).](image)

![FIG. 3. Velocity as functions of various parameters. The velocity shows various dependence on the parameters. From these figures, we can understand the characteristics of the model more profoundly.](image)
mean and correlation $\langle \xi_i(t)\xi_j(s)\rangle = \delta(t-s)\delta_{ij}$. $k$ is the spring constant and $N$ the number of particles. Since the boundary condition of the particles is free, the forces due to springs of Eq. (2.2), the first term of the right-hand side, is
\begin{equation}
k(x_2-x_1-a)
\end{equation}
for the first particle and
\begin{equation}
k(x_{N-1}-x_N+a)
\end{equation}
for the $N$th particle, where $a$ stands for the natural length of the springs. A friction constant $\gamma$ is set to be 1.0 and $D$ stands for the temperature. In our simulation, $U$, $L$, and $a$ are set to be 1.0, 1.0, and 1.35, respectively, if not mentioned.

$h_i(t)$ is a dichotomous random modulation that rules the time-dependent change expected 0 or 1. We determine $h_i(t)$ process as follows. An Ornstein-Uhlenbeck process $Z_i$ ($i=1,2,\ldots,N$), where
\begin{equation}
\langle Z_i(t)Z_j(s)\rangle = \delta_{ij}(D'/\tau)e^{-|t-s|/\tau},
\end{equation}
\begin{equation}
\langle Z_i(t)\rangle = 0, \quad D'=0.4
\end{equation}
is considered and if $Z_i(t)$ is less than 0, then $h_i(t)$ is set to be 0, if $Z_i(t)$ is more than 0, then $h_i(t)$ is set to be 1. Consequently, $h_i(t)$ changes with a correlation time $\tau$ stochastically. In our simulation, $\tau$ is set to be 1.0 if not mentioned. Thereby we can change only three parameters, that is, $k$, $D$, and $N$. If not mentioned, $k$, $D$, and $N$ are set to be 4.0, 0.2, and 20, respectively. At the beginning of our simulations, the $i$th particle is always located at $x=ia$.

III. NUMERICAL SIMULATION

A. Zero-temperature case

1. Dynamics of elastically coupled particles

First of all, we investigate a zero-temperature case ($D=0$). Figure 2 shows the motion of two boundary (first and $N$th) particles (thin lines), and the center of mass of the coupled particles (thick line). A particle trapped to the potential $W_2$ is likely to be near a minimum of it and going to move by the elasticity when trapped to $W_1$. When it is attached to the substrate again under the influence of $W_2$, it goes down to a minimum of the potential $W_2$ to the $+x$ direction more frequently than to the $-x$ direction because of the asymmetry of the potential $W_2$. Consequently, the particles move unidirectionally in an average sense.

2. Parameter dependence of the velocity

Figure 3 shows the velocity of the center of mass of the $N$ particles as functions of various parameters. As a function of $k$, the velocity has a maximum at $k=4.0$ [Fig. 3(a)]. When we change $\tau$, the velocity has also a maximum [Fig. 3(b)], and the graph is bell shaped. The correlation time $\tau$ has to be appropriate if it is nearly equal to the time for particles

FIG. 4. $V$ vs $k$. The lower the temperature is, the faster the particles move in general. For high temperature, the peak almost disappears since the particles tend to move more randomly.

FIG. 5. $V$ vs $a$. When the natural length of the spring is almost equal to an integer times the period of the potential, the model moves so slowly because each particle is caught to a minimum of the potential and cannot escape from it easily.

FIG. 6. $V$ vs $h$. The model moves faster when the transition is restricted. Each line indicates the velocity of the center of mass at different temperature. We can see that the value of the peak and the position of the peak depend on $D$ ($k=4.0,N=20$).
FIG. 7. Load vs efficiency at $D=0.1$ and $k=4.0$. The efficiency has a maximum at $F_{\text{ext}}=0.06$ for the coupled system, and it is less than 0 for $F_{\text{ext}}\geq 0.11$, which means that the model moves in $-x$ direction. Efficiency for a single particle is lower than for the coupled model.

enough to go beyond maximums of $W_2$ in the detached state and for the springs to go back to the natural length in the attached state in order to pull the particles forward before they are detached again. The larger the depth of the potential $W_2$, $U$, becomes, the faster the particles go down toward the minimum of $W_2$. Since the correlation time $\tau$ is finite, the velocity of the particles increases as the depth increases until the velocity saturates for large $U$ such that particles probably arrive at minimums of the potential $W_2$ within the correlation time [Fig. 3(c)]. Finally, we investigate the velocity as a function of $N$ [Fig. 3(d)]. The velocity is almost independent of the number of particles except for small $N$. The dependence of velocity on the size in biological experiments of muscle system is similar to our results.

### B. Finite-temperature case

Figure 4 shows the velocity versus $k$ at $D=0.0, 0.1, 0.2, 0.3$. The velocity decreases as the temperature increases. At $D=0.0$, the velocity is almost independent on the spring constant $k$. The influence of the springs is weak in comparison with that of thermal diffusion for high-temperature $D=0.3$.

Until now, the natural length of the springs $a$ has been set to be 1.35. Now we investigate the relation between the velocity and the natural length in Fig. 5. The velocity disappears when $a/L$ is an integer for $D=0$. That is because particles are trapped at minimums of $W_2$ tightly and cannot escape without thermal noise. At $D=0.1$ and 0.2, however, the average velocity is finite even when $a/L$ is an integer, where particles in the detached state diffuse thermally enough to go over the maximum of $W_2$ against the force of springs.

### C. Restricted transition

Until now, transitions between detached and attached states has been allowed to occur everywhere in the potential. Now we restrict the region where transition toward the detached state can be allowed. This restriction is defined by parameter $h$. That is, when a particle is located at $x$ and if $W_2(x)<h$, transition may be allowed and if $W_2(x)\geq h$ it cannot be allowed. In our simulation, transitions from the detached state to the attached state may occur without any restriction of the positions of the particles. Figure 6 shows the velocity as a function of $h$. For a certain region of $h$, the velocity is larger than that of the unrestricted transition case, which is the right end of Fig. 6. One reason for this, we think, is that the possibility of the particles going over the maximum $W_2$ is larger when transition is restricted near minimums of the potential.

### IV. EFFICIENCY

#### A. Calculation of efficiency

Recently, Sekimoto [10] has defined the efficiency for thermal ratchet models with a load $F_{\text{ext}}$. Dere nyi et al. [11] has also defined the efficiency in another way.

#### TABLE I. $L_U$ and $\gamma v^2$ is shown for various loads. The energy used to the dissipation is much larger than that used to the total motion of the model.

<table>
<thead>
<tr>
<th>Load</th>
<th>$L\times v$</th>
<th>$\gamma \times v^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.000000</td>
<td>0.000000</td>
<td>1.386913</td>
</tr>
<tr>
<td>0.010000</td>
<td>0.002441</td>
<td>1.383389</td>
</tr>
<tr>
<td>0.020000</td>
<td>0.004427</td>
<td>1.392557</td>
</tr>
<tr>
<td>0.030000</td>
<td>0.006206</td>
<td>1.388566</td>
</tr>
<tr>
<td>0.040000</td>
<td>0.007744</td>
<td>1.393150</td>
</tr>
<tr>
<td>0.050000</td>
<td>0.008313</td>
<td>1.397241</td>
</tr>
<tr>
<td>0.060000</td>
<td>0.008928</td>
<td>1.397302</td>
</tr>
<tr>
<td>0.070000</td>
<td>0.008471</td>
<td>1.398886</td>
</tr>
<tr>
<td>0.080000</td>
<td>0.006630</td>
<td>1.404158</td>
</tr>
<tr>
<td>0.090000</td>
<td>0.004963</td>
<td>1.412609</td>
</tr>
<tr>
<td>0.100000</td>
<td>0.001967</td>
<td>1.420848</td>
</tr>
<tr>
<td>0.110000</td>
<td>$-0.000569$</td>
<td>1.417309</td>
</tr>
</tbody>
</table>

FIG. 8. Load vs efficiency at $D=0.1$, $k=4.0$, and $h=0, -0.35$ where $h$ is the parameter to restrict the transition region. The maximum is located at larger $F_{\text{ext}}$ than in the unrestricted case of Fig. 7, and the model moves $+x$ direction even for larger $F_{\text{ext}}$ than the load where motions to $-x$ direction occurs in the unrestricted transitions. Efficiency for the single-particle model is lower than the coupled-particle model in this case as well as in the case of Fig. 7.
If we define $v$ as an average velocity of the model, the conventional expression of the efficiency $\eta$ is

$$\eta = \frac{F_{\text{ext}}v}{P_{\text{in}}},$$

(4.1)

where $P_{\text{in}}$ is the input power [10]. If we define $P_{\text{out}}$ as the power output, it seems more appropriate to define it as

$$P_{\text{out}} = F_{\text{ext}}v + \gamma v^2,$$

(4.2)

rather than only $F_{\text{ext}}v$, the numerator of Eq. (4.2) [11]. The term $\gamma v^2$ denotes, of course, the dissipation via friction. In our simulation, the load $F_{\text{ext}}$ is dispersedly exerted to each particle, that is, $-F_{\text{ext}}/N$ is applied to each particle.

We show in Fig. 7 the efficiency defined by Eq. (4.2). It has a maximum as a function of $F_{\text{ext}}$, and at a certain threshold it has a value less than 0, which means that the model moves in the $-x$ direction. The efficiency for a single particle is shown also in Fig. 7. The efficiency for the coupled model is higher than for the single-particle model.

In Table I, we compare $F_{\text{ext}}v$ with $\gamma v^2$. We find that $\gamma v^2$ is always much larger than $F_{\text{ext}}v$ for any load, which makes sure that most of the energy of the molecular motor is used for the Brownian motion and therefore for the dissipation. This is a reason why the efficiency of the molecular motor is not so high.

Since muscle is known to have much higher efficiency than in our simulation, we do not succeed in reproducing the real situation if we apply our model to muscle contraction. This is mainly because of the simplicity of our model.

**V. CONCLUSION**

We demonstrate that elastically coupled particles in a flashing ratchet model move co-operatively. An important result is the generation of directed motion in a no-thermal condition. Though an isolated particle in flashing models cannot move in a no-thermal condition, elastically coupled particles in flashing models may move by the effects of springs even if thermal diffusion is not allowed. It is also found that the velocity of the particles has a maximum at a specific coupling constant of springs. The maximum of the velocity as a function of $h$, the restriction parameter, appears when we restrict the region of transitions as well. Changing other various parameters, we confirm the complicated behaviors of the elastically coupled particles.

On the efficiency of energy conversion, the effects of the coupling between particles are also found to be very important. The elastically coupled particles may pull a heavier load than a single particle, and the efficiency for the coupled model is larger than the single-particle model. Most of the energy, however, is used for Brownian motion, that is, for the dissipation.

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