1 Introduction

Nano- and micron-scale motors that are able to propel themselves through solution by utilizing the energy derived from chemical reactions constitute an interesting class of objects with numerous potential applications. Such small self-propelled objects have been made in the laboratory and their properties have been investigated. In particular, a good deal of research has been devoted to some of the earliest examples of motors of this type, bimetallic and striped metallic rod motors.\(^1\)–\(^6\) Experimental studies of Janus colloidal motors, where one face of a polystyrene or silica sphere is coated with platinum,\(^7\)–\(^8\) and sphere dimer motors, containing two linked spheres, one silica and the other platinum,\(^9\) have also been carried out. Interest in such motors stems from their potential applications.\(^4\)–\(^6\),\(^8\)–\(^12\) For example, striped metallic rod motors that include Ni can be controlled by external magnetic fields and experiments have demonstrated that targeted cargo delivery can be achieved,\(^13\)–\(^15\) as has transport by Pt–Au nanomotors using photochemical stimuli.\(^16\) Self-propelled Janus particles have recently been investigated experimentally for cargo transport applications.\(^17\)–\(^18\)

Such motors are placed in a solution containing fuel of some type, most often hydrogen peroxide in the cases mentioned above, and chemical reactions on portions of the motor give rise to directed motion. In these examples the motors move by self-phoresis, where the gradient of some field across the motor, which is generated by asymmetrical chemical reactivity, induces fluid flow in the surrounding medium resulting in propulsion. The metallic rod motors exploit an electrochemical mechanism involving oxidation and reduction reactions at the rod components, leading to self-electrophoresis,\(^5\) while the Janus and sphere dimer motors move by self-diffusiophoresis. Motors that are propelled by the generation of bubbles have also been constructed and studied.\(^19\)–\(^21\) While many aspects of propulsion by self-phoresis are generic, the motor motion depends on the nature of the chemical reactions taking place on the motor, as well as those in the environment in which the motor moves.

A characteristic feature of such self-propelled motion is that the entire system is force-free. This has implications for motor propulsion and for the nature of the flows that develop in the system. The treatments of self-propulsion based on the motion of colloidal particles in gradients rely on various continuum approximations.\(^22\)–\(^26\) While these approximations are certainly valid for large particles, and even for quite small particles, as the nano and molecular levels are approached such approximations should be subject to scrutiny. There have been numerous studies of propulsion by self-phoretic mechanisms.\(^27\)–\(^32\) In common with our earlier studies of sphere dimer and polymer motors where a simple reaction \(A \rightarrow B\) occurs on the catalytic sphere,\(^33\)–\(^38\) we consider a particle-based mesoscopic description of the dynamics of self-propelled particles. The dynamics is such that the mass, momentum and energy conservation laws are satisfied and hydrodynamic flows are correctly described.

In this article we show how the mesoscopic dynamics can be extended to treat a general class of chemical reactions at the catalytic sites of the motor. The reactive dynamics is constructed to satisfy the conservation laws so the basic properties of the dynamical laws are preserved. As examples of these more general reactions, we present the results of simulations of...
self-propulsion of a composite Janus particle arising from a dissociation reaction \( A \rightarrow 2B \) and an exothermic reaction \( A \rightarrow B + \Delta u \) at motor catalytic sites. These examples illustrate some interesting effects; in particular, in the latter case of an exothermic reaction propulsion occurs by self-thermophoresis. The general strategy described in this paper should provide a framework that is useful in exploration of the effects of complex chemical kinetics on self-propelled motion at a mesoscopic level.

The outline of the paper is as follows: the system we consider comprises a motor and the environment in which it moves. In Section 2 we describe the (possibly reactive) environment in which the motor moves, the structure of the motor and how it interacts with the chemical species in the environment. A detailed description of the reactive processes at the motor catalytic sites is given in Section 3 and the results of simulations of the motor dynamics are presented in Section 4. The conclusions of the paper are given in Section 5.

2 Mesoscopic dynamical system

The system we consider is a Janus particle motor comprising catalytic and noncatalytic faces in a fluid medium containing reactive and nonreactive chemical species. Chemical reactions may occur in the fluid medium as well as at the catalytic face of the Janus particle. The system is maintained in a nonequilibrium state, either by explicit fluxes of species into and out of the system at boundaries or by nonequilibrium reactions in the bulk fluid. The dynamical evolution of this system is carried out at a mesoscopic level that combines molecular dynamics for the reactive and nonreactive solvent particles in their interactions with the Janus particle, and multiparticle collision dynamics\(^{39,40}\) for the environment in which the Janus particle moves.

2.1 Dynamical model for the environment in which the motor moves

We first focus on the environment and outline how multiparticle collision dynamics (MPCD) can be used to provide a coarse grain description of its time evolution. Additional details concerning this method along with other applications can be found in reviews.\(^{41,42}\) We also discuss the generalization of MPCD to account for chemical reactions.\(^{41}\)

In general terms, molecules in a fluid move and interact or undergo collisions that change their velocities. In the mesoscopic level of description employed in MPCD, explicit details of the interactions among molecules are not considered. Instead, a coarse grained description of collisions is adopted that accounts for the effects of many real collisions and preserves the conservation laws that real collisions must satisfy. More specifically, \( N \) molecules with continuous positions and velocities are confined to a volume \( V \). The \( j \)th molecule has position \( \mathbf{r}_j \) and velocity \( \mathbf{v}_j \). There are no explicit intermolecular interactions among the particles and they move by free streaming between collisions which occur at discrete times \( \tau \). The effective multiparticle collisions take place in the following way: at the time intervals \( \tau \) the system is divided into \( N_c \) cells of volume \( V_c \) (see Fig. 1). The cells are labeled by the index \( \xi \). Rotation operators \( \omega_\xi \), which are chosen from some set of rotation operators, \( \Omega = \{\omega_1, \ldots, \omega_N\} \), are assigned to each cell. The post-collision velocity of particle \( i \) in cell \( \xi \), \( \mathbf{v}'_i \), is given by

\[
\mathbf{v}'_i = \mathbf{v}^\xi + \omega^\xi (\mathbf{v}_i - \mathbf{v}^\xi)
\]

where \( \mathbf{v}^\xi \) is the center of mass velocity of all particles in cell \( \xi \) and \( \omega^\xi \) is the rotation operator for the cell \( \xi \). This algorithm preserves mass, linear momentum and energy within each cell and preserves phase space volumes. The time interval \( \tau \) can be used to control the dynamical regime being simulated. Given the mean velocity of particles as specified by the temperature, the mean free path between collisions depends on the value of \( \tau \). For example, a system with a very small mean free path corresponding to a small fraction of a cell length models a dense fluid. The transport properties for MPCD depend on \( \tau \) and reflect this dependence on the mean free path.\(^{43,44}\) A MPCD fluid obeys an ideal gas equation of state. It also possesses a well defined viscosity that can be tuned using the collision rules.

Since we wish to allow for the possibility that chemical reactions may occur in the bulk of the solution,\(^{45}\) a modification of MPCD that accounts for reactions is employed in the simulations carried out in this study. Reactive multiparticle collision dynamics (RMPCD)\(^{35}\) combines multiparticle velocity-changing collisions among solvent species with local probabilistic rules for changing the identities of those solvent species that participate in reactive events. To carry out the dynamics, we make use of the same partition into cells \( \xi \) and time intervals \( \tau \) that are used for the MPCD algorithm.\(^{46}\) We suppose that various reactions, specified by the index \( j_i \), occur in the solution:

![Fig. 1](image_url) A diagram of the different events occurring in the combined MD–RMPCD dynamics. The small circles in the insets represent \( A \) (blue) and \( B \) (magenta) solvent particles. A Janus particle with a catalytic (C) and a noncatalytic (N) side is pictured within a RMPCD environment. The interaction range of the particle is the thin white shell around it. (a) A nonreactive event on the N side: an \( A \) particle collides with the Janus particle. (b) An \( A \rightarrow 2B \) reactive event on the C side; the \( A \) solvent particle is triggered for reaction when it enters the interaction region (dashed blue line) and the reaction takes place upon exit of the interaction region (dashed magenta line). (c) An \( A \rightarrow B \) reactive event. (d) A bulk MPCD cell before reactive collision. (e) Cell “d” after collision; the velocities of the particles have changed as well as the species of one of the particles (circled in gray). Within example cells (a–c), other solvent particles of the cell could participate in the reactive event.
\[ \sum_{a} v_{a}^{j} X_{a} = k_{j} \sum_{a} v_{a}^{j} X_{a} (j = 1, \ldots, r). \]  

The stoichiometric coefficients for reaction \( j \) are \( v_{a}^{j} \) and \( v_{a}^{j} \) while \( k_{j} \) and \( k_{-j} \) are the rate constants for the forward and reverse reactions. Each reaction, \( \sum_{a} v_{a}^{j} X_{a} = k_{j} \sum_{a} v_{a}^{j} X_{a} \), is taken to occur with probability

\[ p_{j}^{f}(N) = a_{j}^{f} \left(1 - e^{-a_{j}^{f}}\right) / a_{j}^{f}, \]  

where \( N = \{N_{A}, N_{B}, \ldots\} \) is the set of numbers of the different species in cell \( \pi \), and \( a_{j}^{f} = k_{j}(V_{C}) \Pi \sum_{k} \left(N_{k}^{A} - v_{k}^{A}\right) \) accounts for the number of different ways the reaction can occur in the cell, with \( a_{j}^{f} = \sum_{k} a_{j}^{k} \). The rate constants \( k_{j}(V_{C}) \) are scaled to account for the cell volume \( V_{C} \). More specifically, \( a_{j}^{f} \) is the probability density that no reaction will occur up to the time \( t \) and the reaction \( R_{j} \) will occur between \( t \) and \( t + dt \). Since we are also coarse graining reactive collisions over time intervals \( \tau \), the integral of this quantity over the time interval \( \tau \) is \( p_{j}^{f}(N) \) given above in eqn (3). It can be interpreted as the probability that in the time interval \( \tau \) the reaction \( R_{j} \) will occur first, followed by any other events. The probability that no reaction occurs in the time interval \( \tau \) is \( p_{0}^{f}(N) = 1 - \sum_{j} p_{j}^{f}(\tau) = e^{-a_{j}^{f}\tau} \). Reaction rules may be constructed so that mass, momentum and energy are conserved in the reactive events.\(^{43} \)

Once reactive events that change the species through chemical reactions take place, the post-collision velocities of the particles are obtained using MPCD as described above (see Fig. 1). The combined effect of multiparticle collisions and reactive events is to mimic the effects of many real reactive and nonreactive collisions that take place in the time interval \( \tau \). One may show that for a system in local equilibrium and for reactions which occur slowly in the time scale \( \tau \) this dynamics yields the laws of mass action kinetics for the average species concentrations.\(^{43} \) Full RMPCD accounts for fluctuations in the reactive system.

### 2.2 Janus particle

The chemically powered motor we study is a composite Janus particle comprising a rigid assembly of catalytic and noncatalytic beads (see left panel of Fig. 2). One half of the Janus particle is made of catalytic (C) beads while the other half is made from non-active (N) beads. The beads in the Janus particle interact with solvent particles through specific intermolecular forces. The catalytic beads are able to catalyze various chemical reactions that are responsible for the propulsion of the motor.

The evolution of the entire system, the Janus particle and the surrounding solvent, is carried out by combining molecular dynamics (MD) with multiparticle collision dynamics. In this hybrid evolution scheme, Newton’s equations of motion are integrated for all interacting particles for time intervals \( \tau \). This includes the Janus particle and all solvent particles that interact with it. The remainder of the noninteracting solvent particles free stream during these time intervals. At the time intervals \( \tau \) (possibly) reactive multiparticle collisions take place as described above. The time scale of the MD portions of the hybrid MD-RMPCD depends on the intermolecular forces between the Janus and solvent particles. In this case the time scale \( \tau \) for the RMPCD portion of the dynamics is specified by the number of MD steps in \( \tau \). This number can be varied to tune the solvent transport properties as discussed earlier.

Since the full hybrid MD-RMPCD preserves the basic conservation laws, hydrodynamic interactions and hydrodynamic flows induced by the self-propulsion are properly described so that this mesoscopic simulation scheme can capture all essential features of the Janus particle and solvent flow motions.

The last aspect that needs to be specified is the nature of the chemical reactions that take place at the catalytic face of the Janus particle. These reactions are a principal focus of this investigation and the description of how general reactions are implemented at the catalytic sites is the topic of the next section.

### 3 Chemical reactions at motor catalytic sites

In this section we show how different types of reaction can be implemented at a mesoscopic level at the catalytic face of the Janus particle. As noted above, in order to respect the basic conservation laws and ensure that fluid flows are properly described, the reactions must be constructed to satisfy mass, momentum and energy conservation laws. In earlier studies on sphere dimer motors we restricted our considerations to the simple thermoneutral catalytic reaction, \( A + C = B + C \). This reaction amounts to a “coloring” process that changes \( A \) to \( B \) on encounters with the catalytic \( C \) sphere. Mass, momentum and energy are simply conserved in this process. The propulsion arose from the fact that the \( A \) and \( B \) species interacted with different intermolecular potentials at the noncatalytic \( N \) sphere.

The actual catalytic reactions that take place on motor active regions may be quite complex, involving adsorption onto the surface with subsequent formation of products and their release, or binding to active sites in enzymes followed by
conformational or other changes and product release. All of these events could involve dissipation of energy into local solvent degrees of freedom, either directly or subsequent to various internal energy transfer processes within the motor. Our mesoscopic treatment of such reactive events coarse grains over most of these molecular-level details. In particular, when the reagents lie within the interfacial zone around the catalytic portion of the Janus particle where intermolecular forces between the reagents and Janus particle beads are non-zero, there is a possibility that a reaction will take place. We subsume all of the reaction details into effective reactive events where reactive species conversion and possible energy distribution in solvent degrees of freedom take place just as the species exit the reactive interfacial zone. The local solvent molecules that participate in any energy exchanges with reactive species are taken to lie in the MPC collision cell that contains the reactive species. Using this strategy it is possible to construct a general class of catalytic reactive events at the active face of the Janus particle.44

In this coarse-grain description of real reactions catalyzed on a Janus motor, the conservation of momentum and energy in a general reactive event can be written as

$$m_j v_{j'} + \sum_{i \in \xi} m_i v_{i'} = m_j v_j + \sum_{i \in \xi} m_i v_i,$$

(4)

for the momentum, where $v_j$ and $v_i$ are the velocities of the Janus particle and a solvent particle with corresponding masses $m_j$ and $m_i$, $\xi$ denotes the cell in which the reacting particles are found and the prime again denotes post-reaction quantities. Summing over all particles in the cell $\xi$ allows one to utilize the velocities of solvent particles, other than the reacting ones, to satisfy conservation rules. In order to model endothermic or exothermic reactions we associate an internal degree of freedom with the chemical species corresponding to an internal energy $u_a$. (For thermoneutral reactions this internal state label is not needed.) Allowing for such internal states, energy conservation reads

$$\frac{1}{2} m_j v_{j'}^2 + \frac{1}{2} \sum_{i \in \xi} m_i v_{i'}^2 + \sum_{i \in \xi} u_i = \frac{1}{2} m_j v_j^2 + \frac{1}{2} \sum_{i \in \xi} m_i v_i^2 + \sum_{i \in \xi} u_i.$$  

(5)

A more detailed notation will be required below and we shall make use of the following: $m_a$ is the mass of species $a$; $M_{a,b,c,}$ is the sum of the masses of the different species, for instance, $M_{c,ab} = m_a + m_b$ and $M_{a,m} = 2m_a$; $\mathbf{r}_a$ is the position of the particle $a$; $\mathbf{R}_b$ is the center of mass position of particles $a$ and $b$; $\mathbf{r}_{a,b} = \frac{m_a \mathbf{r}_a + m_b \mathbf{r}_b}{M_{a,b}}$ is the unit vector along the $\alpha - \beta$ direction; $v_{i,a}$ is the velocity of the particle $a\alpha$; $v_{\alpha,b} = \frac{m_a v_a + m_b v_b}{M_{a,b}}$ is the center of mass velocity of $a$ and $\beta$; $\mathbf{u}_{a,b} = v_a - v_b$ is the relative velocity of $a$ and $\beta$; and $\mu_{ab} = m_a v_{a,b} M_{a,b}$ is the reduced mass. For some of the quantities, for instance the sum of the masses or the center of mass velocity, arbitrary arrangements of indices can be made. For quantities where the order matters, such as $\mathbf{u}_{a,b}$ or $\mu_{a,b}$, grouping can be made with a parentheses, e.g., $\mu_{a,b} = m_a v_{a,b}/M_{a,b}$. The precise manner in which reaction rules are constructed depends on the chemical mechanism one wishes to simulate. We illustrate the procedure by considering two different reaction schemes: an exothermic unimolecular reaction $A + J \rightarrow B + J + \Delta u$, where $\Delta u$ is the energy release on reaction, and a dissociation reaction $A + J \rightarrow J + 2B$, where $J$ represents the Janus particle.

There is considerable freedom in the way one chooses to implement a reactive scheme that accounts for the conservation conditions in eqn (4) and (5). The number of participating solvent particles in the reacting cell is not prescribed by the method. Also, the exchange of velocities may involve random reorientations. For example, in the dissociation reaction discussed below, we chose to put all of the relative kinetic energy in the solvent particles. A redistribution between $J$, $B_1$, and $B_2$ could have been considered. Since the species-changing reactive event is followed by an MPC collision, energy is redistributed among all particles in the cell. As a consequence, on the mesoscopic time scale $\tau$, many details of the energy redistribution among reactive and other solvent species will not play a significant role and we do not expect major changes in our results for slightly different rules.

### 3.1 Exothermic unimolecular reaction

The exothermic unimolecular reaction involves the reactant and product species, $A$ and $B$, respectively, interacting with the Janus particle $J$. The reactive $A$ and $B$ species, while being treated as structureless objects, nevertheless carry internal energy labels $u_A$ and $u_B$, respectively. For example, this is a simplified description of a catalyzed isomerization reaction where reactants and products differ by a free energy difference of $\Delta u = u_B - u_A$. Mass balance requires that the masses of $A$ and $B$ be equal: $m_A = m_B$. Recall that the Janus particle is a rigid object so that no Janus internal degrees of freedom need to be taken into account. (This condition can be relaxed when flexible self-propelled particles are considered.)

For $A + C \rightarrow B + C + \Delta u$, we choose to account for the energy change $\Delta u$ on reaction by changing the post-collision relative kinetic energy $\mathbf{w}_{bs}$, of the product $B$ molecule and an additional solvent molecule $S_1$, and not the Janus particle. In this case the momentum conservation equations take the form,

$$m_A v_A + m_B v_B = m_A v_A' + m_B v_B' + m_{S_1} v_{S_1},$$

(6)

The second of these equations can be satisfied if the post-collision velocities are given by

$$v_B' = v_B + \frac{\mu_{bs}}{m_B}$$

$$v_{S_1}' = v_{S_1} - \frac{\mu_{bs}}{m_{S_1}}$$

(7)

where $s$ remains to be determined by energy conservation.

Supposing $u_A > u_B$ for now, we find that the norm of the relative post-collision velocity $\mathbf{w}_{bs}$ depends on the pre-collision relative velocity $\mathbf{w}_{AB}$ and also on the difference of internal
energies. As \( v_j \) remains unmodified by the reaction, the deposition of the excess energy \( \Delta u \) is made in the relative velocity \( w'_{BS1} \). Eqn (5) becomes, in the center-of-mass velocity frame of \( B-S_1 \),

\[
\frac{1}{2} M_{BS1} \dot{v}_{BS1}^2 + \frac{1}{2} \mu_{BS1} w_{BS1}^2 = \frac{1}{2} M_{AS1} \dot{v}^2 + \frac{1}{2} \mu_{AS1} w_{AS1}^2 + u_A - u_B.
\]

(8)

Since the center-of-mass velocity \( \dot{v}_{BS1} \) is not altered during the event, eqn (8) becomes after some algebra

\[
w_{BS1}^2 = w_{AS1}^2 + \frac{2 u_A - u_B}{\mu_{AS1}}.
\]

(9)

From eqn (7) we have \( w_{BS1} = w_{AS1} + s\). Using \( s = |s| \) and \( \delta = s/s \) we find

\[
s = -w_{AS1} \cos \theta_+ \sqrt{\left( w_{AS1} \cos \theta_+ \right)^2 + \frac{2 u_A - u_B}{\mu_{AS1}}}.
\]

(10)

where \( \theta_+ \) is the angle between \( s \) and \( w_{AS1} \). We decide to always pick the + solution. The orientation of \( s \) may be chosen at random.

As a consequence of the local increase in kinetic energy an inhomogeneous temperature gradient will be generated in the vicinity of the Janus particle. If the interaction energies of the \( A \) and \( B \) species with the Janus particle are same, this model exhibits the interesting feature that propulsion will be driven by thermophoresis alone.

3.2 Dissociation reaction

Here we suppose that a dissociation reaction, \( A \rightarrow B_1 + B_2 \), is catalyzed by interaction with the rigid Janus particle. The symbols \( B_1 \) and \( B_2 \) label the two \( B \) species molecules that result from the dissociation of \( A \). Mass conservation requires that the mass of a \( B \) molecule is half that of \( A \): \( m_B = m_A/2 \). Momentum conservation reads

\[
m_A \dot{v}_A + m_B \dot{v}_B = m_A \dot{v}_A + m_B (\dot{v}_B + \dot{v}_0).
\]

(11)

The energy conservation equation, eqn (5), can be written in the form

\[
\frac{1}{2} \mu_{AA} |w_{AA}|^2 = \frac{1}{2} \mu_{BB1} |w_{BB1}|^2 + \frac{1}{2} \mu_{BB2} |w_{BB2}|^2 + \frac{1}{2} \mu_{AB} \dot{v}^2 + \frac{1}{2} \mu_{AB} \dot{v}_0^2.
\]

(12)

We may choose to set \( w_{BB1} = 0 \), i.e., we may use all of the relative kinetic energy of the system to change the relative energy of \( B_1B_2 \). The direction of \( w_{BB1} \) may be chosen at random. The norm of \( w_{BB1} \) is then given explicitly by

\[
|w_{BB1}| = \sqrt{\frac{\mu_{BB2}}{\mu_{BB1}}} |w_{BB2}|^2.
\]

(13)

Other variants of these collision rules may be constructed.

4 Simulation of Janus particle motion

In this section we present the results of simulations of the self-propelled motion of the composite Janus particle when it catalyzes the exothermic \( A \rightarrow B \) unimolecular reaction and \( A \rightarrow 2B \) dissociation reaction. The simulations were carried out in a cubic box with periodic boundary conditions. This simulation volume contained \( N = N_A + N_B \) molecules. The \( A \) and \( B \) particles interact with all beads in the Janus particle through repulsive Lennard-Jones potentials,

\[
V(r) = 4\epsilon \left( \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 \right) + \Theta \left( r - r_c \right),
\]

(14)

where \( r_c = 2^{1/6} \sigma \) and \( \Theta \) is the Heaviside function. Note that we choose the interactions of both species with both catalytic and noncatalytic beads to be identical; consequently the particles are identical apart from species labels and no propulsion of the Janus particle can arise from the simple \( A \rightarrow B \) thermoneutral reaction considered in previous studies of sphere dimer motors. As described earlier, interactions between the \( A \) and \( B \) particles in the environment are accounted for by multiparticle collisions instead of direct intermolecular forces. In the MD portions of the simulation, Newton’s equations of motion are integrated using the velocity Verlet algorithm with a time step \( \Delta t \). The Shake and Rattle algorithms are used to maintain the rigid bond constraints in the composite Janus particle. In the MPC portions of the evolution, grid shifting is applied at each MPCD step to ensure Galilean invariance.

All quantities reported below are expressed in simulation units: mass \( m_A \), length \( a \), energy \( \epsilon \) and time \( \sqrt{\frac{m_A a^2}{\epsilon}} = t_{sim} \). In these units the system parameters were: simulation box size, \( 48 \times 48 \times 48 \); MPC time \( \tau = 0.5 \); average number of particles per MPC cell, \( 10 \); reduced temperature, \( k_B T = 1/3 \); and \( \Delta t = 0.01 \). The choice of the time step depends on the criterion \( \Delta t \ll t_{ij} \) where \( t_{ij} \) is the natural time for the Lennard-Jones parameters, \( t_{ij} = \sqrt{\frac{m_A a^2}{\epsilon}} \). The Janus particle is made from 36 individual spheres, each with Lennard-Jones radius \( \sigma = 3 \) and mass 50, so that \( m_A = 1800 \). All realizations of the dynamics started with 100 MPCD steps with an Andersen thermostat at chosen temperature \( k_B T = 1/3 \) in systems with all \( A \) particles. We note that for these parameters the thermal velocity for the Janus particle is

\[
v_{th} = \sqrt{\frac{k_B T}{m_A}} = 0.014.
\]

In order to sustain self-propelled motion in the steady state reactants must be supplied to the system and products removed. In the past this has been done by introducing fluxes of these species at the boundaries or far from the motor, or by bulk far-from-equilibrium reactions which are themselves assumed to be sustained by fluxes. (In simple descriptions these species feeds can be accounted for in pool chemical species that may be incorporated into effective rate constants.) In this study we maintain a supply of fresh reagents by taking the reverse of the reaction on the motor to occur in the bulk of the fluid, but with different reaction rates. The reverse reaction for the exothermic reaction is \( B \rightarrow A - \Delta \epsilon \) while for the dissociation reaction it is \( 2B \rightarrow A \). The reaction rates of both of these reactions were taken to be \( k_0 = 0.001 \). These reactions are implemented using the RMPCD algorithm. Momentum and energy are obeyed at the cell level as in eqn (4) and (5) but without the Janus particle.
Energy addition in a cell is obtained by a velocity scaling in the center-of-mass velocity frame.

4.1 Exothermic $A \rightarrow B$ reaction

As noted earlier, since the $A$ and $B$ particles are identical apart from species labels, the only possible source of directed motion (assuming instabilities do not occur) is the temperature gradient across the Janus particle generated by the exothermic reaction. In simulations starting from all $A$ molecules in the simulation box, after a transient of 3000 time units, the combined effect of reactions at the Janus particle and in the bulk leads to a steady average value of $\bar{N}_A$ and $\bar{N}_B$. For the system in this steady state regime, the number concentrations $c_A$ and $c_B$ are displayed in Fig. 3 and show a strong excess of $B$ at the top (catalytic side) of the particle. The polar axis of the Janus particle is used to define a coordinate system that moves along the particle. The definitions of the radius $r$ and angle $\theta$ are shown in Fig. 2 (right). The temperature field $T(r, \theta)$ around the compound is shown in Fig. 4. The temperature is computed cell-wise at regular intervals during the simulations and binned at the center of mass of the corresponding cell. One clearly observes a difference between the $C$ and $N$ sides of the Janus particle, a sign that a thermal gradient is present in the stationary regime of the system.

The velocity of the self-propelled Janus particle can be determined from the time and ensemble average of $V_z(t)$, the instantaneous velocity of the center-of-mass velocity projected on the instantaneous unit vector from the center of mass of the $N$ hemisphere to the center of mass of the $C$ hemisphere,

$$\langle V_z \rangle(t) = \left\langle \frac{1}{t} \int_0^t \mathbf{r}_{CN}(t') \mathbf{v}_j(t') dt' \right\rangle,$$

where the angular brackets denote an average over realization of the evolution. When the time argument is omitted, $\langle V_z \rangle = \langle V_z \rangle(t_{\text{max}})$ where $t_{\text{max}}$ is the duration of the simulation.

Fig. 5 displays $\langle V_z \rangle(t)$. After a short transient, the running average stabilizes around an approximately constant nonzero value, indicating the existence of directed motion. The value of the velocity is positive, consistent with the results obtained by Yang and Ripoll\(^7\) where the thermal gradient is generated externally on the fluid around one sphere of a sphere dimer.

The directed velocity $V_z$ is not negligible compared to the thermal velocity $v_{th}$ and the directed component of the motion plays an important role. Thus, we have demonstrated that the chemically induced temperature gradient by the chemical reaction leads to self-thermophoretic propulsion of the Janus motor.

The behavior for different values of $\Delta u$ is summarized in Table 1 where $\langle V_z \rangle$ and its standard deviation, which characterizes the coherence of $V_z$ between different runs of the same physical conditions, are given. As expected, increasing the energy $\Delta u$ associated with the reaction increases the propulsion velocity.

![Fig. 3](image3.png) The concentration field for $A$ and $B$ molecules, $c_A$ and $c_B$ for $\Delta u = 0.5$.

![Fig. 4](image4.png) The temperature field around the Janus particle for $\Delta u = 0.5$. The dark blue region is related to the very low solvent concentrations $c_{A/r, \theta}$ close to the Janus particle that leads to very poor statistics.

![Fig. 5](image5.png) The running average of the directed velocity for $\Delta u = 0.5$. The result is obtained from an average of 16 independent realizations of the dynamics.

| $\Delta u$ | $\langle V_z \rangle \times 10^3$ | $\sigma_{\langle V_z \rangle} \times 10^3$ |
|-----------|-----------------|-----------------|
| 0.1       | 1.5             | 0.8             |
| 0.2       | 2.8             | 0.9             |
| 0.3       | 3.8             | 1.2             |
| 0.5       | 5.7             | 1.4             |

Table 1 The average directed velocity and the associated standard deviation for the exothermic reaction for several values of exothermicity. Results are averages over 16 independent realizations.
5 Conclusions

The directed motions of small nano- and micron-scale motors that operate by chemically fueled phoretic mechanisms depend on the nature of the catalytic chemical reactions that occur on the motor. These reactions generate the gradients of fields that are responsible for the propulsion of the motor. The work described in this investigation demonstrated how a variety of chemical mechanisms can be incorporated into a mesoscopic description of the dynamics of the system. One of the important aspects of propulsion by phoretic mechanisms is the correct description of the fluid flow fields that are induced by the motor motion. In order to properly describe the hydrodynamics of the motor environment, the mesoscopic dynamics must preserve the basic conservation laws of mass, momentum and energy. The hybrid MD-MPCD dynamics employed in this study satisfies these criteria. We have shown how different catalytic reactions at the motor surface can be constructed that maintain these conservation laws.

The two examples we have chosen to illustrate the method present interesting features. For both the exothermic unimolecular and dissociation reactions we have chosen to consider the situation where the $A$ and $B$ reactive species interact with the same intermolecular forces with the Janus particle. This is a special but interesting case that highlights the role that the reaction mechanism plays in the motor propulsion. In the case of the exothermic reaction, the net effect of the reaction is to generate an inhomogeneous temperature gradient in the vicinity of the Janus particle, giving rise to propulsion by a thermophoretic mechanism. This scheme is easily generalized to allow for different interaction potentials for the chemical species on the different hemispheres of the Janus particle. In such a situation both thermophoretic and diffusiophoretic mechanisms will operate, and they can act in either the same or opposite directions. For the dissociation reaction, it is the increased number of product molecules produced in the reaction, even when the potential energies of interaction of reactant and products with the Janus particle are the same, that leads to propulsion by self-diffusiophoresis. Again if these potential energies were different the propagation velocity would change.

The work presented here provides a method that allows one to explore self-propulsion for a variety of motor geometries and catalytic reaction mechanisms at a mesoscopic level. Through such studies one can investigate motor dynamics on scales where the validity of continuum theories should be tested and study situations where analytical methods are difficult to carry out due to the complex nature of the reaction dynamics.

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