Myosin V is a vesicle transporter that unidirectionally walks along cytoskeletal actin filaments by converting the chemical energy of ATP into mechanical work. Recently, it was found that myosin V force generation is a composition of two processes: a lever-arm swing, which involves a conformational change in the myosin molecule, and a Brownian search-and-catch, which involves a diffusive “search” by the motor domain that is followed by an asymmetric “catch” in the forward actin target such that Brownian motion is rectified. Here we developed a system that combines optical tweezers with DNA nano-material to show that the Brownian search-and-catch mechanism is the energetically dominant process at near stall force, providing 13 k_BT of work compared to just 3 k_BT by the lever-arm swing. Our result significantly reconsiders the lever-arm swinging model, which assumes the swing dominantly produces work (>10 k_BT), and sheds light on the Brownian search-and-catch as a driving process.

Key words: Brownian motor, myosin, motor protein, nano machine

Myosin V is a dimeric motor protein that transports various cargos such as melanosomes, endoplasmic reticulum and messenger RNAs by moving processively along cytoskeletal actin filaments. Because of the cell’s dense and heterogeneous actin meshwork, the cargo itself should apply a dynamic load onto the myosin V. To overcome this load, a single myosin V converts the chemical free energy of 1 ATP into a directional motion of 36 nm and a maximum force of 2–3 pN. Its structure is composed of a motor domain (or head domain), which includes the actin and nucleotide-binding site, a lever arm, which has six calmodulin-binding sites, and a tail domain, which is responsible for dimerization. Finally, myosin V operates as a dimer.

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Fluctuation in lever-arm swing under load

We also observed -20 nm sub-steps that coupled with the 20 nm lever-arm swing (Fig. 2c), suggesting these sub-steps are the result of a lever-arm reversal (Fig. 1, state 3 → state 2). The duration of state 3 and 2 in Figure 1, \( \tau_1 \) and \( \tau_2 \), respectively, were load-dependent and fit to a single exponential curve. The inverse of the duration describes the transition rates to represent a 20 nm lever-arm swing and 57 nm Brownian search-and-catch.

**Figure 1** Hand-over-hand motion of myosin V. The rear head binds ATP and detaches from actin (state 1 → state 2 transition). A 20 nm sub-step of the rear (unbound) head results from a lever-arm swing (state 2 → state 3 transition). The detached head then undergoes a Brownian search-and-catch to bind to the next actin-binding site (state 3 → state 4 transition). ADP release from the rear head followed by attachment of new ATP molecule there returns us to state 1. At high load, the lever arm can also reverse its motion (state 3 → state 2 transition).

The DNA handle is composed of double-stranded DNA ~60 nm in length and tethers the myosin head directly to a 200-nm fluorescent polystyrene bead (Fig. 2a). The bead position is detected with nanometric and millisecond spatio-temporal resolution. Myosin V was observed walking along actin filaments using this setup.

In the presence of ATP, the observed 77 nm processive steps were decomposed into 20 and 57 nm sub-steps (Fig. 2b, inset). Because the duration of the sub-steps was 123 ms, which is comparable to the first-passage time of a Brownian head to reach the forward actin-binding site 77 nm away under load and because the expected bead displacement from the lever-arm swing is 20 nm, we concluded the sub-steps...
parameter, d, which relates to the force dependency of the reaction, indicates the lever-arm reversal is less sensitive to load than the lever-arm swing. Therefore, once the lever-arm has swung forward, it robustly maintains the post-lever-arm swing conformation when sensing load.

Lever-arm swing vs. Brownian search-and-catch

Given the individual rate constants, we can estimate the free-energy difference (ΔG) between the pre- and post-lever-arm swing states of the lead (bound) head using the formula

\[ \frac{k_{\text{reversal}}}{k_{\text{swing}}} = e^{\frac{\Delta G}{k_B T}} \]

where \( \Delta G \), F, d, \( k_b \) and T denote the free energy difference in the absence of load, force exerted by the optical tweezers, the characteristic distance, Boltzmann’s constant and absolute temperature, respectively. At no load, which describes the maximum energy bias of the lever-arm swing, \( \Delta G^0 \) was estimated to be \(-3.3\, k_b T\) (Fig. 3c).

The work done by a myosin dimer can be divided into two parts, work done by structural changes in the lever-arm and work done by the Brownian search-and-catch. We showed hand-over-hand stepping is triggered by a \( 3.3\, k_b T \) lever-arm

\[ \frac{1}{\tau_1} = k_{\text{total}} = k_{\text{reversal}} + k_{\text{catch}} + k_{\text{detach1}} \]

\[ \frac{1}{\tau_2} = k_{\text{total}} = k_{\text{swing}} + k_{\text{detach2}} \]

The ratio of the rates correspond to the frequency of the observed events (\( k_{\text{reversal}} : k_{\text{catch}} : k_{\text{detach}} = N_{\text{reversal}} : N_{\text{catch}} : N_{\text{detach}} \) and \( k_{\text{swing}} : k_{\text{detach2}} = N_{\text{swing}} : N_{\text{detach2}} \), where \( N_i \) is the number of observations for state i. Figure 3a shows the normalized frequencies of the subsequent transition state used to estimate the transition rates.

Figure 3b shows the load dependent transition rates for the lever-arm swing and lever-arm reversal. The lever-arm swing involves a structural change that is opposed by the load. Therefore, the reaction rate decreases with increasing load in a manner that obeys an Arrhenius-type transition,

\[ k(F) = k_0 e^{\left(\frac{-F \cdot d}{k_B T}\right)} \]

where \( k_0 \) is the transition rate in the absence of load, F is force, d is the characteristic distance (force sensitivity), and \( k_B T \) is the thermal energy.

For similar reasons, load should promote lever-arm reversal, which is consistent with our observation that the transition rate slightly increases with increasing load. The fitting
on average (data not shown). This equates to a Brownian search-and-catch contribution of 13 kBT (57 nm × 0.9 pN) of work, which is similar to the chemical free energy from Pi release (12 kBT) when a myosin head strongly catches forward actin. Consequently, myosin V predominantly works during the Brownian component by controlling Pi release, while the structural change of the lever-arm is a trigger for the forward catch. Supporting this theory, we reproduced 2–3 pN stall force and 18 kBT total work at the tail loading condition in a Langevin dynamics simulation using a 3.3 kBT energy bias in the lever-arm swing and experimentally obtained the transition rate constants.

swing in the lead (bound) head. The Brownian search-and-forward-catch contributes work to this system as load is exerted onto the myosin molecule. During physiological vesicle transport, myosin V apparently distributes the load from a cargo bound to the tail domain completely onto the lead (bound) head. However, when the rear (unbound) head undergoes Brownian motion, the lever-arm in the lead (bound) head should bend 10–15 nm backwards when a 2–3 pN load is exerted (bending stiffness = ~0.2 pN/nm)\(^2\). In these cases, the Brownian head will conduct work when it catches an actin target 77 nm forward. Therefore, the work done by the Brownian search-and-catch is physiologically relevant. We found that the majority of myosin V molecules completed the Brownian search-and-catch against 0.93 pN on average (data not shown). This equates to a Brownian search-and-catch contribution of 13 k_BT (57 nm × 0.9 pN) of work, which is similar to the chemical free energy from Pi release (12 k_BT)\(^2\) when a myosin head strongly catches forward actin. Consequently, myosin V predominantly works during the Brownian component by controlling Pi release, while the structural change of the lever-arm is a trigger for the forward catch. Supporting this theory, we reproduced 2–3 pN stall force and 18 k_BT total work at the tail loading condition in a Langevin dynamics simulation using a 3.3 k_BT energy bias in the lever-arm swing and experimentally obtained the transition rate constants\(^3\).
Strain sensor as a rectifier of Brownian motion

How does Brownian head ensure a strong catch? We have previously proposed the “strain sensor mechanism,” which explains the decision-making by the Brownian head. We describe it briefly below.

It is known that when the Brownian head is bound to both ADP and Pi, it undergoes weak interactions with actin, which can be described as an equilibrium state between rapid (sub millisecond) attachments and detachments. We constructed an ultrafast optical tweezers assay that can directly observe the weak binding state by detecting weak attachments between myosin and actin. We applied this system to myosin VI, which moves the opposite direction of myosin V, and found that Pi release, which corresponds with the transition from weak to strong binding, was very mechano-sensitive. When backward load (opposite direction to the myosin movement) was applied to a weak-binding head, the strong binding was greatly accelerated (~30-fold), whereas forward load did not affect the weak-to-strong transition rate. In the case of dimeric myosin V and VI, intramolecular strain should occur when forming a two-headed bound state (Fig. 1, states 1 and 4). When the Brownian head spans the actin helical pitch and forms a two-headed bound state, the rear head senses a forward strain and the weakly-attached front head senses a backward strain. Because the backward load accelerates strong binding, the forward catch is ensured.

We have proposed a model to explain how strong binding is accelerated in a strain-dependent manner. The opening and closing of the exit route for Pi, termed the “back door,” has been mechanically linked to that of the nucleotide (ATP or ADP)-binding pocket, or “front door.” The strain dependencies of the ADP release and ATP binding rates have previously been measured, revealing that external force applied to the head strains the front door. When the lever-arm is pulled backward, the head is bent forward, which closes the front door and opens the back door. Thus, backward strain accelerates Pi release and hence strong binding. At the same time, closing of the front door suppresses ADP release or ATP binding, which means the overall ATP turnover rate is slow under backward load. On the other hand, upon forward strain, the backdoor is closed or unaffected, resulting in a relatively rare transition to strong binding. This strain-dependent, asymmetric catch mechanism should be important for the Brownian head’s reliable forward movement.

We believe the structure-based mechanism of the strain sensor should be applicable to not only myosin VI but myosin V. It is known that the lever arm of myosin VI is bent 180 degrees, which distinguishes it from myosin V, and that this bend is responsible for the different direction of movement. Because the angle between the lever-arm and head should be important for the opening and closing of the front and back doors, we propose the relationship between the opening and closing between the two myosins are opposite based on the different positions of the lever arms.

To explicitly examine the relationship between the direction of myosin movement and the strain-sensor mechanism, we applied our ultrafast optical tweezers assay to myosin V. The results of the myosin V experiments are in agreement with our model, which assumes backward strain accelerates strong binding (unpublished data). Therefore, the strain-sensor mechanism should contribute to the directionality of the myosin motor.

The strain sensor mechanism is notable in that random Brownian motion is rectified in one direction by sensing the intensity and direction of the mechanical strain. Using this simple mechanism, myosin can autonomously sense positional information and adaptably respond to changes in its environment. These characteristics should be important in motor assembly systems like muscle.

Physiological advantages of the Brownian machine

Figure 4 shows the contribution of the lever-arm swing and Brownian search-and-forward catch to myosin V work at various loads when assuming physiological vesicle transport geometry (vesicle is attached at the tail domain). When 0.5 pN force is applied to the tail end, the lever-arm swing conducts 2.4 k_B T (20 nm × 0.5 pN) of work, whereas the total work done by myosin V is 4.4 k_B T (36 nm × 0.5 pN), meaning the Brownian search-and-catch component contributes 2.0 k_B T. Although the amount of work done by the lever-arm swing increases with load, it never exceeds 3.3 k_B T at saturating physiological ATP concentration. Any residual work at high loads then is done by the Brownian search-and-catch. Thus, our results argue the myosin V is a lever-arm-driven motor under low loads, but a Brownian search-and-catch driven motor under high loads. This result suggests that myosin V can change its force-generating mechanism depending on the external force.

Knowing that the vast majority of work done by myosin V at high load is a stochastic process (Brownian search-and-catch) and that the proportion of work done by the deterministic lever-arm swing and Brownian search-and-catch varies with load offers important insights into the mechanisms used by myosin V for its function. The hand-over-hand steps caused by deterministic lever-arm swings would be advantageous for smooth, rapid movement at near-zero load when no obstacles are present. However, cells contain a cytoskeleton meshwork and a number of dynamic molecules and vesicles that risk disturbing transport. In these cases, the Brownian search-and-catch mechanism should be advantageous for avoiding such obstructions. Thus, our findings indicate that myosin V is an optimized nanomachine highly adaptable to its intracellular environment, which should have significant implications on the design of artificial nanomotors.
Perspectives

To conclude, we have succeeded to quantify both the bias energy of the lever-arm swing and the work done by the Brownian search-and-catch. Surprisingly, the bias energy of the lever-arm swing (~3 k_BT) was much smaller than expected according to the lever-arm swinging model (>10 k_BT)^32,33. Instead, inherent Brownian motion is the dominant driving force. This motion is rectified by steric compatibility between the myosin head and actin filament^34 or by a strain sensor mechanism coupled with Pi release. The requisite energy to work is temporarily supplied from a thermal bath (Brownian motion) and forward diffusion is locked by consuming the chemical free energy of Pi release. In this sense, myosin V is a Brownian machine that can adjust its function in response to environmental change. This design should be advantageous for constructing artificial systems in which individual components autonomously coordinate to achieve efficient and adaptable function.

Accordingly, we are constructing an artificial muscle system using DNA nanotechnology called DNA origami^35. DNA origami is an attractive tool in that it can be used to create three-dimensional nanostructures^36. Regarding an artificial muscle system, we can attach myosin heads to DNA origami, which acts as an artificial thick filament (assembly of muscle myosin). Using DNA origami allows us to control not only the number of myosin heads, but also the spacing between heads. This model will be applied to studying the design principles of the muscle architecture, including how multiple Brownian machines achieve autonomous biological function in concert.

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