Twirling motion of actin filaments in gliding assays with non-processive myosin motors

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We present a model study of gliding assays in which actin filaments are moved by non-processive myosin motors. We show that even if the power stroke of the motor protein has no lateral component, the filaments will rotate around their axis while moving over the surface. Notably, the handedness of this twirling motion is opposite from that of the actin filament structure. It stems from the fact that the gliding actin filament has “target zones” where its subunits point towards the surface and are therefore more accessible for myosin heads. Each myosin head has a higher binding probability before it reaches the center of the target zone than afterwards, which results in a left-handed twirling.

We present a stochastic simulation and an approximative analytical solution. The calculated pitch of the twirling motion depends on the filament velocity (ATP concentration). It reaches about 400μm for low speeds and increases with higher speeds.

Introduction

Gliding assays, also known as motility assays, represent the oldest in vitro technique to study motor proteins [1,2]. They consist of attaching motors (like myosins or kinesins) with their tails to a glass surface and adding the filaments (actin or microtubules). The motors will then pull the filaments and make them glide over the surface (Fig. 1A). Gliding assays are the most convenient way of testing motors for their functionality, measuring their speed in the absence of load and for testing their processivity. Several experimental and theoretical studies were dealing with the pathways of such filaments in the two-dimensional plane [3,4,5]. Interestingly, one group observed that gliding actin filaments move in a helical fashion [6,7]. In a subsequent experiment the pitch of rotation was determined as about 1μm, although the applied optical detection method did not allow discrimination between left- and right-handed rotation [8].

Helical motion of myosin motors has been very important in a somewhat different context. The processive motor myosin V has an average step size that is close, but not precisely equal to the actin periodicity. The helical motion of a motor around the actin filament therefore presents a very accurate way of measuring the difference between its step size and the filament pitch. Ali and coworkers have observed that myosin V walks on an actin filament along a left-handed helix with a pitch of 2.2μm [9] and thus has a step size slightly shorter than the actin half pitch (for a discussion of the myosin V step size see [10,11]).

Myosin VI, despite having a shorter lever arm than myosin V, showed either straight walking, or, in 20% of cases, a helical path with a pitch of 2.3μm [12]. Sun et al. [13] confirmed this result, but also showed that the relatively straight motion contains a large amount of random wiggling. New experiments on myosin X also show a left-handed helical motion with a pitch that is somewhat shorter than that of myosin V and VI [14].

In a recent experimental study Beausang and coworkers [15] used polarized total internal reflection microscopy to study the twirling motion of actin filaments in gliding assays with processive myosin V and non-processive muscle myosin (myosin II). While the twirling of filaments driven by myosin V agreed with the helical movement of single molecules mentioned above, myosin II interestingly showed a left-handed twirling motion as well. This result came as a surprise and the left-handed rotation is opposite from the observations by Nishizaka et al. [7]. But they are not in direct contradiction, as they were obtained with quite different ATP concentrations.

While the pitch of the twirling motion is a direct measure for the step size of processive motors, its interpretation is more complicated with non-processive ones. They could clearly generate twirling motion if there was a lateral component of the power stroke. In fact, there exists indirect evidence for such an asymmetry in myosin V [10]. However, we will show in this paper that there is another, more subtle, effect that can cause twirling motion of actin filaments in a gliding assay, even if the myosin heads exhibit no lateral motion. This effect stems from the fact that myosin heads can only bind to an actin filament in so called target zones, where the actin binding sites have approximately the right orientation (Fig. 1D) [17,18]. When a target zone is approaching a myosin head, the latter is more likely to bind at the beginning of the target zone than at its end, because it is more likely that it is already bound by that time. In this paper we will show simulation results and develop an approximative theory to estimate the pitch of helical motion resulting from this effect. Of course, we cannot exclude that there are other contributions towards the helicity. But, because the rotation is relatively weak as compared with the longitudinal motion, these effects can easily be treated separately and the total rotation is simply their superposition.

Model definition

In order to concentrate on the effect of filament rotation, we define a model with simplified myosin kinetics, essentially containing a detached state, a bound pre-powerstroke state, a bound post-powerstroke state with ADP and a bound rigor state. At the same time, we take
into account the full helical structure of the actin filament. We propose an actin filament which can move in one direction and rotate around its axis. The position of the actin filament at a given time is therefore described with the coordinates \((X, \Theta)\). As follows from the helical structure of the actin filament, the \(x\)-coordinate of each bindings site is \(X + ia\) and its azimuth angle \(\Theta + i\delta_0\), where \(\delta_0 = -(13/28) \times 360\degree = -167.14\degree\) is the rotation and \(a = 2.75\) nm the axial rise per subunit. The definition of the model is illustrated in Fig. 1.

We assume that the myosin motors are distributed randomly directly under the gliding actin filament (a discussion how this simplified, one-dimensional model follows from the full, 2-D model is given in the Appendix). The motor numbered \(j\) is anchored at position \(x^j_M\). The elastic energy cost of binding a head to the site \(i\) consists of a longitudinal component with stiffness \(K\) and an angular component with stiffness \(K_\phi\) and can be written as

\[
U_i = \frac{1}{2}K(x + ia - x^j_M)^2 + \frac{1}{2}K_\phi(\Theta + i\delta_0 + 2\pi n)^2
\]  

(1)

with \(n\) chosen such that the angle \(\Theta + i\delta_0 + 2\pi n\) falls into the interval \([-\pi, \pi]\). The binding rate is then proportional to the Boltzmann factor

\[
k_A^i = k_A \exp \left[ \frac{U_i}{k_B T} \right].
\]  

(2)

This is essentially the expression used by Steffen et al. [17] to fit binding rates of a single myosin head to the actin filament. They determined the value of the angular stiffness expressed with the dimensionless coefficient

\[
\alpha = K_\phi / k_B T
\]  

(3)

as \(\alpha = 3.7\). However, this value needs to be regarded as a lower estimate, as it might partially result from torsional compliance of the actin filament, rather than myosin heads. We therefore use three different values of \(\alpha = 4, 6\) and 8 in the simulation. The angular contribution to the Boltzmann factor for a set of binding sites is shown in Fig. 2. For the longitudinal compliance, we use the value \(K = 0.5\) pN/nm, somewhat below the stiffness of myosin heads in muscle, which is about 2.5 pN/nm [19]. The lower stiffness reflects the additional compliance due to myosin tails and roughly corresponds to the value obtained with optical tweezers, 0.69 ± 0.47 pN/nm [20].

The force and the torque that a myosin head numbered \(j\), bound to site \(i\), exerts on the filament are

\[
F^j = K(x^j_M + \delta - X - ia)
\]

\[
M^j = -K_\phi(\Theta + i\delta_0 + 2\pi n).
\]

(4)

Here we introduced the displacement \(\delta\) that has the value 0 in the pre-powerstroke and \(d = 8\) nm in the post-powerstroke and rigor state.

The simplified model for the duty cycle of the myosin head is defined as follows. A head binds to an actin site...
with the rate \( k_i \) given by Eq. 2. The power stroke (a transition from \( \delta = 0 \) do \( \delta = d \)) takes place with the rate \( k_{PS} \). It is followed by the release of ADP with the rate \( k_{-ADP} \). Detachment follows after binding a new ATP molecule, therefore its rate depends on the ATP concentration, \( k_D = k_D^0[A\text{TP}] \). We neglect the strain dependence of those rates, as well as the existence of the reverse transitions. It should be noted that in this formulation the model is not thermodynamically consistent. However, as we are only interested in dynamics at low loads, this does not significantly affect the results.

We assume that the filament position is quickly equilibrated after each step, therefore it always fulfills

\[
F = \sum F^j = 0 \quad M = \sum M^j = 0. \quad (5)
\]

**Simulation results**

Given the known structure of the actin helix and the power stroke size of myosin, our model essentially has two important parameters: the angular stiffness of myosin heads, \( K_\phi \), and the ratio between the detachment- and the attachment rate, \( k_D/k_A \). The latter is a function of the ATP concentration and is closely related to the duty ratio of motors. For other parameters we use the values given in Table I. The power stroke, connected with the phosphate (Pi) release, is assigned a very fast rate, \( k_{PS} = 10000\text{s}^{-1} \), and can be considered as taking place immediately after binding. The maximum attachment rate \( k_A \), i.e., the attachment rate to sites that do not require any elastic distortion, can be estimated as \( 50\text{s}^{-1} \). This reflects the estimated average attachment rate of \( 30\text{s}^{-1} \), or a maximum ATP turnover rate of \( 25\text{s}^{-1} \) in muscle [21]. The ADP release takes place with the rate \( k_{-ADP} = 1000\text{s}^{-1} \), characteristic for the fast myosin isoform [18]. For the detachment rate, which is determined by the ATP binding rate, we use \( k_D^0 = 5\mu\text{M}^{-1}\text{s}^{-1} \) [18].

**TABLE I: Simulation parameters**

| Parameter                  | Value                              | Description                        |
|----------------------------|------------------------------------|------------------------------------|
| \( k_A \)                 | \( 50\text{s}^{-1} \)             | Attachment rate                    |
| \( k_{PS} \)              | \( 10000\text{s}^{-1} \)           | Power stroke rate                  |
| \( k_{-ADP} \)            | \( 1000\text{s}^{-1} \)            | ADP release rate                   |
| \( k_D \)                 | \( (5\mu\text{M}^{-1}\text{s}^{-1}) \) | Detachment rate                    |
| \( d \)                  | \( 8\text{nm} \)                  | Power stroke size                  |
| \( K \)                  | \( 0.5\text{pN/\text{nm}} \)      | Myosin stiffness (longitudinal)    |
| \( K_\phi \)              | \( 4, 6, 8 \)                      | Dimensionless angular stiffness    |
| \( a \)                  | \( 2.75\text{nm} \)               | Distance between actin subunits    |
| \( \phi_0 \)              | \( -167.14^\circ \)               | Angle between actin subunits       |
| \( l \)                  | \( 5.5\text{\mu m} \)             | Actin filament length              |
| \( \rho \)                | \( 20\mu\text{m}^{-1} \)           | 1-d myosin density on surface      |
| \( k_BT \)                | \( 4.14 \times 10^{-21}\text{J} \) | Thermal energy                     |

The stochastic simulation essentially followed the following algorithm:

1. Distribute the positions of myosin motors \( x^j_M \) randomly along the distance covered by the actin filament, with an average linear density \( \rho \). Set \( X = 0 \) and \( \Theta = 0 \).

2. Determine the total rate of all possible transitions as

\[
 k_{\text{total}} = \sum_{j \in \text{Motors}} \begin{cases} 
 \sum_{i \in \text{Binding sites}} k_A^i(j) & \text{if motor } j \text{ detached} \\
 k_{PS} & \text{if motor } j \text{ in pre-PS state} \\
 k_{-ADP} & \text{if motor } j \text{ in post-PS state} \\
 k_D & \text{if motor } j \text{ in rigor state} 
\end{cases}
\]

\( k_A^i(j) \) is determined using Eq. 2 with the current values of \( X \) and \( \Theta \).

3. Determine the time until the next step as \( \Delta t = k_{\text{total}}^{-1} \ln(1/r) \), where \( r \) is a random number between 0 and 1.

4. Choose randomly one of the possible steps (attachment, detachment, power stroke, ADP release), so that the probability of choosing a certain step is given by its rate, divided by \( k_{\text{total}} \).

5. Change the state of the chosen motor and update the filament position \( X \) and angle \( \Theta \) according to Eq. 5.

6. Continue with step 2 until \( X \geq X_{\text{max}} \) (\( X_{\text{max}} = 1000\mu\text{m} \)).

7. Determine the average speed as \( v = X/t \) and pitch as \( \lambda = 2\pi X/\Theta \).

The results of this numerical simulation are shown in Fig. 3, which shows the inverse pitch of twirling as a function of the ATP concentration for three different values of
FIG. 3: (A) Inverse twirling pitch (number of rotations per micron traveled) as a function of the ATP concentration. The lines represent 3 different values of the angular stiffness ($\alpha = 4$, $\alpha = 6$ and $\alpha = 8$). Negative signs denote left-handed rotation. The upper scale shows the velocity (for $\alpha = 4$). (B) Gliding velocity as a function of the detachment rate. (C) Duty ratio $\eta$, giving the average fraction of myosin heads in the bound state.

FIG. 4: Simplified model used for the analytical solution. The filament travels with a stationary velocity $v$ and rotates with an angular velocity $\omega$. $x$ denotes the position of a myosin anchoring point relative to the center of the target zone.

The angular stiffness $K_\phi$. For reference, velocity (Fig. 3B) and duty ratio (Fig. 3C) are included as well. The behavior of the pitch is non-monotonous: it has a minimum of about 400 – 500 nm at intermediate speeds, but increases both at high, as well as very low speeds.

These results show that the helicity of the actin filament is sufficient to explain the twirling motion in a gliding assay. Somewhat counter-intuitively, this rotation is left-handed, and therefore opposite from the handedness of the actin filament. The effect becomes weaker for high speeds (where the distance traveled between two attachment events becomes longer), as well as for very low speeds, where motors have enough time to bind even to unfavorable sites outside the target zones.

Analytical approximation

In the following we will describe an approximative analytical solution with the aim of understanding and quantitatively reproducing the twirling dynamics. The essential simplification we will make is to neglect the discrete nature of binding sites on the actin filament and replace them with a continuous helical “groove”. The simplified model is shown in Fig. 4. We denote each head with its root position $x$ relative to the center of the target zone:

$$x = x_M - \left( X - \frac{L}{\pi} \Theta + nL \right)$$

(6)

with $n$ such that $-\frac{L}{\pi} < x < \frac{L}{\pi}$. A bound head is additionally characterized by the strain $\xi$, which is the position of its root relative to its binding site. For a head bound to site $i$, this is $\xi_i = x_M - X - \Theta i$.

In the original model the total binding rate for a motor positioned at $x$ is

$$\tilde{k}_A(x) = \sum_i k_A^i(x) = \sum_i k_A \exp \left[ -\frac{K_\xi_i^2 + K_\Theta \theta_i^2}{2k_BT} \right] \right].$$

(7)

In the sum, we only consider sites that are turned towards the motor-covered surface, therefore we can write the angle as

$$\theta_i = \left( x - \xi_i \right) \frac{\pi}{L}$$

(8)
We can now set up a Master equation for the probability that a motor positioned at \( x \) is in the attached state
\[
\partial_t A(x, t) - c\partial_x A(x, t) = \tilde{k}_A(x)(1 - A(x, t)) - k_D A(x, t)
\] (13)
and set \( \partial_x A = 0 \) to obtain the stationary solution. \( A \) also has to fulfill the periodic boundary condition
\[
A(-L/2) = A(L/2) .
\] (14)
The expectation value of the attachment position can be calculated as
\[
\langle x_A \rangle = \frac{\int_{-L/2}^{L/2} x \tilde{k}_A(x)(1 - A(x))dx}{\int_{-L/2}^{L/2} \tilde{k}_A(x)(1 - A(x))dx}
\] (15)
and the average strain at the time of attachment follows from (13):
\[
\langle \xi_A \rangle = \frac{K'_\vartheta}{K + K'_\vartheta} \langle x_A \rangle .
\] (16)
Because the strain on a motor changes with time as \( \dot{\xi} = -v \), the average strain of all bound motors is \( \langle \xi_A \rangle = v/k_D \).
The force per motor is then \( F = \frac{K}{\vartheta} \langle \xi + d \rangle = \frac{K}{\vartheta} \langle \xi_A \rangle = -v/k_D + d \). As the total force produced by all motors has to be zero, we obtain an expression for the velocity
\[
v = k_D (d + \langle \xi_A \rangle) .
\] (17)
The same type of calculation as for the velocity can be made for the angular velocity. Motors attach with an average angle \( \langle \vartheta_A \rangle \). As the filament rotates, their angle changes as \( \dot{\vartheta} = \omega \). The average angle of all motors is \( \langle \vartheta_A \rangle + \omega/k_D \) and needs to be zero because of torque balance, therefore
\[
\omega = -k_D \langle \vartheta_A \rangle = -k_D \frac{K}{K + K'_\vartheta} \left( \frac{\pi}{L} \right) \langle x_A \rangle .
\] (18)
These equations, together with the Master equation (13), the periodic boundary condition (14), the expression for \( \langle \xi_A \rangle \) (16) and for \( \tilde{k}_A \) (7) allow us to numerically determine the velocity \( v \) and the distribution of attached heads \( A(x) \) in a self-consistent manner. An example of the solution \( A(x) \), along with the attachment rate \( \tilde{k}_A(x) \) and attachment flux \( \tilde{k}_A(x)(1 - A(x)) \) is shown in Fig. 5A. Well visible is the asymmetry in the attachment flux. The expectation value of \( x \) at the time of attachment (\( \langle x_A \rangle \)) as a function of the ratio \( k_D/k_A \) is shown in Fig. 5B. It reaches its maximum when \( k_D/k_A \) is such that each motor travels an average path of \( \approx L/3 \) between two attachment events.

This finally gives us the expression for the twirling pitch
\[
\lambda = \frac{2\pi v}{\omega} = -\frac{2Ld \left( 1 + \frac{K'_\vartheta}{K} \right)}{\langle x_A \rangle} - \frac{2L K'_\vartheta}{K} .
\] (19)
FIG. 5: (A) Top panel: probability $A(x)$ that a motor with a position $x$ relative to the center of a target zone is in the bound state. The parameters are $k_D/k_A = 0.01, 0.1, 1, 10$ (each a different line type) and $\alpha = 4$. Middle panel: dimensionless attachment rate $\bar{k}_A(x)/k_A$. Bottom panel: attachment flux, obtained as the product of the probability that a motor is in the detached state and the attachment rate. Note that the asymmetry is most pronounced for $k_D/k_A = 0.1$. (B) The expectation value of $x$ at the time of binding, $\langle x_A \rangle$, as a function of $k_D/k_A$.

The results are shown in Fig. 6 and compared with simulation data from Fig. 4. The simulation results are well reproduced, although there is a certain discrepancy which is more pronounced for low values of the angular stiffness $\alpha$. The main reason for this discrepancy is the extrapolation beyond the boundaries of one period, which was used in the derivation of Eq. 10. Other (minor) sources of deviation are the neglected discrete nature of binding sites and of the fact that each binding site can only be occupied by one head at a time.

Discussion

In this study we have demonstrated that the helical actin structure, along with the fact that myosin heads preferentially bind to those sites oriented towards them, is sufficient to explain left-handed rotation in a gliding assay. The maximum twirling motion is achieved at relatively low speeds (below 100 nm/s). Twirling is reduced with higher speeds, achieved at higher ATP concentrations. Interestingly, it is also reduced under extremely low ATP concentrations, when the velocity drops under 10 nm/s. However, in this regime the results depend strongly on the choice of the angular stiffness $\alpha$, which is less well known. The minimum pitch resulting from this effect lies in the order of 400–500 nm, which is in good agreement with recent experimental results [13]. The model also makes a testable prediction that the pitch should increase with a higher ATP concentration. Because pitch only depends on the ratio between the attachment and detachment rate, addition of ADP should have the same effect on the pitch as a reduced ATP concentration that yields the same filament speed. If this turns out not to be the case, it will be a strong indication of a lateral conformational change in the myosin head connected with the release of ADP.

Although the qualitative aspects of our theory are generic and practically independent of any assumptions other than the helical actin structure, there are alternative effects that could well contribute to the twirling motion. One such possibility is that the power-stroke of the myosin head contains a lateral (azimuthal, off-axis) component. A similar effect could also result from an asymmetric attachment rate, which could cause the attached motors to exert a certain torque on the filament immediately after binding. Such a torque does not contradict the laws of thermodynamics because the first bound state is not in equilibrium with the detached state. A related
in the filament could show some sideways motion in the 2-D model. In the 2-D model, which describes the filament underneath the actin filament, and which assumes that motors are distributed one-dimensionally and have no distinct target zones. Any rotation resulting from an effect of the kind we describe here would be negligible. Therefore, as suggested in [22], the rotation caused by kinesins has to result from a lateral (off-axis) component of a power stroke, or from an asymmetry in the binding rate.

**Appendix**

In the following we will discuss how the simplified model, which assumes that motors are distributed one-dimensionally underneath the actin filament, and which we use throughout the main text, relates to the full two-dimensional model. In the 2-D model, which describes the actual situation in a gliding assay, myosin heads are distributed all over the glass surface and their positions are described with two coordinates, \((x_M, y_M)\). The position of the actin filament is described with the coordinates \((X, Y)\) and the angle \(\Theta\). We assume that the filament keeps its direction parallel to the X-axis. The elastic distortion of the head \(j\) binding to the site \(i\) can then be written as

\[
U_j = \frac{1}{2} k (X + ia - x_M^j)^2 + U_A(\Theta + i\vartheta_0, Y - y_M^j) \tag{20}
\]

where \(U_A\) is an unknown function of the azimuth angle of binding site \(i\) and of the lateral position of the filament relative to the motor.

The total binding rate to site \(i\) of all motors located at longitudinal position \(x_M\) is then

\[
k_A(x_M) = \int_{-\infty}^{\infty} k_A \exp \left[ - \frac{U_i(X + ia - x_M, \Theta + i\vartheta_0, Y - y_M)}{k_BT} \right] \times \rho(D(x_M, y_M)) dy_M \tag{21}
\]

where \(\rho\) is the 2-D surface density of myosin motors and \(D(x_M, y_M)\) the probability that a motor at that position is in the detached state. If we assume that \(D(x_M, y_M) = D(x_M, 2Y - y_M)\), i.e., that the distribution of unbound heads is symmetric with respect to the filament, the resulting function \(k_A(x_M)\) has to be symmetric in \(\Theta + i\vartheta_0\). We can therefore approximate it with the expression used in Eqs. [112].

The average torque generated by a head that binds to site \(i\) can be calculated as

\[
\langle F_l \rangle = -\frac{\int_{-\infty}^{\infty} \exp (-U_i/k_BT) D(x_M, y_M) \partial U_i/\partial Y dy_M}{\int_{-\infty}^{\infty} \exp (-U_i/k_BT) D(x_M, y_M) dy_M} \tag{23}
\]

If \(D(x_M, y_M)\) is independent of \(y_M\), or, more generally, if it has a dependence that can be written as a function of \(U_i\), the integral in the numerator is 0 and there is no lateral force. However, with different distributions \(D(x_M, y_M)\), a small lateral force is possible, so that the filament could show some sideways motion in the 2-D model. While the torque results from a \(D(x_M, y_M)\) which is asymmetric in \(x_M\), a lateral force needs asymmetry in both coordinates and is therefore a higher-order effect.

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