Contribution of myosin II activity to cell spreading dynamics†

Noam Nisenholz, Aishwarya Paknikar, Sarah Köster, and Assaf Zemel

Myosin II activity and actin polymerization at the leading edge of the cell are known to be essential sources of cellular stress. However, a quantitative account of their separate contributions is still lacking; so is the influence of the coupling between the two phenomena on cell spreading dynamics. We present a simple analytic elastic theory of cell spreading dynamics that quantitatively demonstrates how actin polymerization and myosin activity cooperate in the generation of cellular stress during spreading. Consistent with experiments, myosin activity is assumed to polarize in response to the stresses generated during spreading. The characteristic response time and the overall spreading time are predicted to determine different evolution profiles of cell spreading dynamics. These include, a (regular) monotonic increase of cell projected area with time, a non-monotonic (overshooting) profile with a maximum, and damped oscillatory modes. In addition, two populations of myosin II motors are distinguished based on their location in the lamella; those located above the major adhesion zone at the cell periphery are shown to facilitate spreading whereas those in deeper regions of the lamella are shown to oppose spreading. We demonstrate that the attenuation of myosin activity in the two regions may result in reciprocal effects on spreading. These findings provide important new insight into the function of myosin II motors in the course of spreading.

I. Introduction

Actin polymerization at the leading edge of a cell provides the driving force for cell spreading and locomotion. Facilitated by cell–substrate friction, actin polymerization drives the extension of thin lamellipodia protrusions that act as motor units at the leading edge of the cell and elastically pull on the cell body forward. A few lines of evidence support this elastic picture. Experiments consistently show that both the cell area and force are monotonically increasing functions of substrate rigidity, and the total (steady-state) force exerted into the matrix has been reported to linearly increase with cell projected area.

Furthermore, we have recently demonstrated with endothelial cells that the development of cellular stress occurs concurrently with the increase in cell area during spreading. The origin of this apparent long-term elasticity of the cytoskeleton on the time scale of spreading [see also ref. 9] is however not fully understood; an attempt to explain it has recently been proposed.

Myosin II activity is known to be a major factor in cellular force generation, however, the impact of this activity on the ability of a cell to spread is not fully understood. Experiments monitoring the consequences of myosin inhibition using various drug treatments (e.g., Rho kinase inhibitor, ML-7, blebbistatin and others) yielded controversial results; in some experiments myosin II forces were found to facilitate spreading (inhibition caused impaired spreading),15–18 in others to inhibit it14,19–21 and some showed no effect.21 Recent investigations have indicated that the two myosin isoforms, non-muscle myosin IIA and IIB, may have distinct functions during spreading. Experiments on fibroblasts,11,14 and similarly on T-cells,22 have shown that the contractile activity of myosin IIA inhibits spreading since the depletion of this protein resulted in faster spreading and larger cell spread areas. In addition, the non-muscle myosin IIA has been shown to be essential for maintaining the integrity of the cytoskeleton and for transmitting forces from one cell end to the other.11,14 In contrast, inhibition of the myosin IIB isoform had little effect on fibroblast spreading.11,14 Similar conclusions have been reached by Betapudi et al.23,24 with breast cancer cells and other cell types who found myosin IIA to inhibit spreading and myosin IIB to facilitate spreading. The measured differences in the spatial distribution of the two myosin isoforms have shown to be consistent with the observed behavior but accurate interpretation of these data is difficult in these cells due to the significant degree of overlap in the spatial...
distributions of these proteins. Interestingly, the contrasting effects of the two myosin isoforms are consistent with the reported roles of the myosin IIB isoform in neurite outgrowth, and of the myosin IIA isoform in neurite retraction. In this cell type myosin IIB concentrates mainly in the peripheral domain of the growth cone while the myosin IIA is found more posteriorly in the growth cone and along the axon.

These studies exemplify the complexity of myosin functions during spreading and the need for a quantitative theoretical investigation of this phenomenon. A number of computational and analytical models have been focused on the role of myosin in force generation and cytoskeleton organization. However, to the best of our knowledge, no previous study has theoretically investigated the contribution of myosin forces to the dynamics and the extent of spreading, nor is there a quantitative analysis of the consequences of the spatial distribution of myosin motors on the regulation of cell area and force. In addition, it is unknown how the intrinsic dynamics of myosin II activity (that generally, are governed both by signaling mechanisms as well as by the physical characteristics of acto–myosin interactions) influence the dynamics of spreading. In a recent study, we presented a minimal elastic theory of cell spreading dynamics that quantitatively demonstrated how the dynamics of spreading on elastic substrates is limited by the rate of actin polymerization at the cell front and cell–substrate adhesion kinetics. Treating the cytoskeleton as an elastic network has shown to be essential for explaining the universal dependence of the cell area and force on substrate rigidity. We nevertheless deliberately did not make explicit account of myosin forces and their dynamics since our goal has been to derive a minimal model of spreading dynamics; myosin activity has only been implicitly accounted for via its contribution to the effective elastic moduli of the cell.

In this paper, we extend our theory to explicitly investigate the consequences of myosin activity on cell spreading dynamics and mechanics. Because cell–substrate adhesions and the tractions exerted onto the substrate often tend to localize at the cell periphery, we adopt a coarse-grained model and theoretically distinguish two populations of myosin II molecular motors: those that locate above the adhesion zone at the cell periphery and facilitate the pulling forward of the cell, and those located more centrally in the cytoskeleton and generally inhibit spreading. Biochemically, these two populations may be identical (e.g., be the same isoform) and could therefore be indistinguishable by various drug treatments, however, we show theoretically that myosin activity in the two spatial locations possesses distinct functions during spreading. We demonstrate that the steady-state radius of the cell is dictated by a balance between myosin exerted forces in the two spatial locations; those at the lamella front facilitate spreading, and those in the lamella bulk oppose spreading. The steady-state force is shown to be dictated by myosin activity in the lamellae front that augments the frictional interaction between the cell and the substrate. These conclusions shed light on the ambiguity revolving the effects of myosin inhibition on cell spreading area, and are consistent with the generic tendency of cellular forces to decrease with myosin inhibition.

In addition, we demonstrate that the dynamics of myosin activity in the lamellar bulk may dictate different spreading profiles. These include a “regular” monotonic increase of cell area with time, an over-damped mode in which the cell area first rises to a maximum and then decreases with time to a steady state, and damped oscillations. Such slow modulations of cell area with time have been discussed in the literature in terms of late myosin contraction (and the reciprocal effects of Rac and Rho activity) but have not been quantitatively and systematically studied. Our analysis thus provides important new insight into the function of myosin activity during spreading.

II. Model

Our starting point is a linear theory of cell spreading dynamics described in detail in ref. 8. Accordingly, we model the cell lamella as a homogeneous and isotropic elastic disc of thickness $h$ and radius $R$ that is actively stretched by the propulsion of the cell front forward (see Fig. 1). The cell adheres to the substrate via multiple (transient) adhesion contacts that are assumed to concentrate in a narrow rim between $R$ and $R + L$ (thus neglecting the limited distribution of adhesions underneath the cell body). This coarse-grained approximation separates the lamella into two distinct zones: the adhesion zone at $R < r < R + L$, and an adhesion-free zone at $r < R$. The width $L$ is taken to be fixed during spreading and smaller than

![Schematic illustration of major mechanical elements at the cell front.](https://example.com/image.png)
the cell size, $L \ll R$. The radial extension speed of the cell front is dictated by two oppositely oriented velocities:

$$R(t) = v_{pol} - v_F(t)$$

(1)

here, $v_{pol}$ is the mean radial actin polymerization speed and $v_F(t)$ is the simultaneous retrograde flow of the polymerizing network (measured relative to the substrate frame of reference). In writing this equation we assume that the actin polymerization speed, $v_{pol}$, remains constant during spreading, while the retrograde flow, $v_F(t)$, continuously increases in the course of spreading. This is motivated by recent studies in which the retrograde flow was shown to increase while cell spreading decelerated, and the sum $R(t) + v_F(t)$ remained constant; similar behavior has also been reported for growth cone advancement.

The retrograde flow arises from the forces generated in the cytoskeleton, which in turn are sustained by the frictional interaction between the lamellar network and the substrate, see Fig. 1. This frictional interaction arises from the distribution of transmembrane adhesion receptors (e.g., integrins) that physically and transiently link the actin network to ligands on the substrate. The retrograde flow of the actin network induces local elastic deformations in the vicinity of each adhesion contact. We denote as $s_t$ the local tangential displacement of the lamella, and as $s_m$ the corresponding local displacement of the substrate; $s = s_t + s_m$ is the total local displacement, see the box in Fig. 1. The lipid membrane is assumed to be stationary with respect to the substrate (although lipids can still flow in the plane of the membrane). Using a simple stochastic model for the binding kinetics of these adhesion contacts one may relate the mean displacement and the mean number of contacts to the drag velocity, $v_F$, as follows (see ref. 8 and 44 and Section III in the ESI for a detailed derivation):

$$v_F = \frac{\langle s_t \rangle}{\tau_b} + \frac{\langle s_m \rangle}{\tau_b}$$

and

$$N_b = \frac{\tau_b N}{\tau_b + \tau_{ub}}$$

(2)

here, $N$ is the total density of integrin linkers per unit area of the cell membrane, $N_b/N$ is the fraction of bound integrins, $\tau_b$ is the mean adhesion life time and $\tau_{ub}$ is the mean time of the unbound state. While cell–substrate adhesions have been shown to behave as catch bonds, namely to stabilize when acted upon by force, we here adopt the simplifying assumption that both $\tau_b$ and $\tau_{ub}$ are force and displacement independent. The consequences of this assumption on the dependence of cell spreading speed on the forces generated during spreading have been tested experimentally and this showed the assumption to be a reasonable approximation.

The mean displacements and the retrograde flow are dictated by the forces generated in the cytoskeleton. Denoting as $\sigma_R$ the mean radial stress exerted by the cytoskeleton at $r = R$, averaged over the lamella height, $h$, and as $\langle t_i \rangle$ the mean radial traction acting parallel to the cell basis and averaged over the width of the adhesion layer, $L$, one has (for $L \ll R$) the following force balance condition at the cell front,

$$h \sigma_R = L \langle t_i \rangle = f$$

(3)

$f = F/(2\pi R)$ is the mean radial force per unit length and $F$ is the total radial force exerted into the matrix. The right and left hand sides of eqn (3) can be written in terms of the associated local deformations of the adhesion layer and the cytoskeleton as follows:

$$\langle t_i \rangle = N_b k_m \langle s_m \rangle = \frac{N_b \langle s_l \rangle}{Q} - Q,$$

(4)

and

$$\sigma_R = 2k_c u_{tr} + \zeta_c \dot{u}_{tr} - P$$

(5)

In eqn (4), $k_l$ and $k_m$ denote local effective spring constants that may be related to the bulk moduli of the cell and the substrate, respectively, via Boussinesq Green’s function approach. The quantity $Q = Q(t)$ represents the mean contribution of active actomyosin contractile forces in the lamella front ($R < r < R + L$) to the shear stress at the cell–substrate interface; these forces act in parallel to the passive elasticity of the lamella (see Fig. 1).

In eqn (5), $u_{tr}(t) = [R(t) - R_0]/R_0$ and $\dot{u}_{tr}(t) = d u_{tr}/dt$ denote, respectively, the mean radial strain and strain-rate of the cytoskeleton where $k_c$ is an effective area-expansion modulus and $\zeta_c$ is an effective viscosity coefficient. The quantity $P = P(t)$ represents the mean contribution of acto-myosin contraction in the adhesion-free zone ($r < R$) to the radial stress in the cytoskeleton. The reference length, $R_0$, is used here as a constant for simplicity, although it may have its own dynamics and is generally expected to be increasing in the early stages of cell adhesion as the lamella network assembles at the cell basis.

Since acto-myosin forces are generally contractile, $Q$ and $P$ are taken as negative quantities and the minus sign in eqn (4) and (5) indicates that the stresses they generate are positive (tensile and pointing inward). We take both $Q$ and $P$ to be zero at $t = 0$, hence only the increase in myosin dipolar stress from its baseline level in the suspended cell in solution is included; thus the cross-linking function of myosin motors is included in the value of the elastic moduli $k_l$ and $k_c$. As will be shown below, the variations in $Q$ and $P$ induce opposite effects on cell spreading dynamics. While the enhancement of $Q(t)$ facilitates spreading and results in more extended cell sizes and stronger forces, an increase in $P(t)$ opposes spreading and renders the cytoskeleton stiffer.

A variety of experimental and theoretical studies have indicated that the magnitude and the orientation of acto-myosin forces are regulated by the local stress in the cytoskeleton. Motivated by these studies, we include a simplified account of the dynamics of the acto-myosin polarization response, and study the effect on cell spreading dynamics. Accordingly,
Q(t) and P(t) are assumed to polarize in accordance with the elastic part of the corresponding stress in the particular regime, namely Q \sim N_s k_l (\kappa_l) and P \sim k_m u_{fr}. For simplicity, we associate only one (major) time scale with each response, and we denote them as \tau_q and \tau_p, respectively. We thus write the following phenomenological relaxation response as:\textsuperscript{50,55}

\[ \tau_q Q + Q = -\beta N_s k_l (\kappa_l) \] (6)

\[ \tau_p P + P = -2\kappa_m u_{fr} \] (7)

In these equations, \alpha and \beta reflect an active susceptibility of the acto-myosin dipolar stresses Q(t) and P(t) to the evolving stresses in the two domains. The minus sign in front of \alpha and \beta ensures that Q(t) and P(t) become more negative (more contractile) as the tensile stresses increase in the two domains.

Eqn (1)–(7) constitute a closed set of equations to solve for the evolution of the cell size, \( R(t) \), force, \( f(t) \), and myosin exerted stress \( P(t) \) and \( Q(t) \). The steady-state solution can be examined by setting \( Q = P = R = 0 \) and noting that in this case \( v_{fr} = v_{pol} \) (see eqn (1)). Substituting the steady-state values of \( Q \) and \( P \) in eqn (4) and (5), respectively, and using eqn (2) and (3) one finds the following expressions for the steady-state values of cell radius and force:

\[ R_{ss} = R_0 \left(1 + \frac{\tilde{\varepsilon}_s v_{pol}}{2 k_c (1 + \alpha)} \right) \] (8)

\[ f_{ss} = \tilde{\varepsilon}_s v_{pol} \] (9)

with

\[ \tilde{\varepsilon}_s = L N_0 \tau_b \left( \frac{(1 + \beta) k_l k_m}{(1 + \beta) k_l + k_m} \right) \] (10)

These equations reveal the dependence of \( R_{ss} \) and \( f_{ss} \) on the myosin susceptibility factors, \( \alpha \) and \( \beta \), that determine the extent of myosin activity in the steady-state. Importantly, eqn (8) illustrates the competing effects that myosin activity in the two lamellae regions has in the determination of the cell spreading area; while inhibition of myosin activity in the lamellar bulk is predicted to facilitate spreading (through the effect of \( \alpha \)), the inhibition of myosin activity at the cell front would impair it (through the effect of \( \beta \)). In addition, while our equations show that \( R_{ss} \) (oppositely) depends on both \( \alpha \) and \( \beta \), the steady-state force, \( f_{ss} \), is seen to only depend on \( \beta \), via the dependence of \( \tilde{\varepsilon}_s \) on \( \beta \). This is the expected result so long as adhesions are transient and the cytoskeleton is free to retrogradely flow relative to the substrate and contract. In this dynamic steady-state the force is dictated by the frictional interaction, \( \tilde{\varepsilon}_s v_{fr} = \tilde{\varepsilon}_s v_{pol} \), between the cell and the substrate, which monotonically increases with myosin activity in the adhesion zone. The inhibition of myosin activity at this state thus leads to a decrease in cellular stress since it impairs the physical coupling of the cell to the substrate, and not necessarily due to the weakening of myosin forces in the lamellar bulk.

The set of eqn (1)–(7) can be reduced to a third-order linear differential equation in \( P(t) \) and consequently solved for all other variables (see ESIT). To demonstrate the essential properties of the solution, however, it is instructive to specialize in the simpler situation where the dynamics expressed in eqn (6) of \( Q(t) \) towards a steady-state are neglected; physically this may arise in cases where the reorganization of actomyosin forces more deeply in the cytoskeleton is slower than that in more peripheral locations. The neglect of \( Q(t) \) dynamics is examined in the ESIT and shown to provide a fairly good approximation to the full solution.

Assuming \( \tau_q \) to be sufficiently small, eqn (6) can be restated as follows:

\[ Q = -\beta N_s k_l (\kappa_l) \] (11)

where \( \tilde{\varepsilon}_s \) and \( f_{ss} \) are given in eqn (9) and (10).

Combining eqn (5) and (11), using eqn (7) and noting that \( v_{fr} = v_{pol} \) we find the following differential equation for \( R(t) \):

\[ \frac{\tau_p}{1 + \alpha} \frac{R}{R_{ss}} + \frac{\tau_p + \tau}{1 + \alpha} \frac{R}{R_{ss}} + R = R_{ss} \] (12)

This equation can be now solved for explicit expressions for the evolution of the cell size and force; assuming \( P(0) = 0 \) and \( R(0) = R_0 \) one finds:

\[ \frac{R(t) - R_0}{R_{ss} - R_0} = 1 - \frac{(1 + \alpha - \omega_+ \tau) e^{-\omega_+ \tau} - (1 + \alpha - \omega_+ \tau) e^{-\omega_+ \tau}}{\tau (\omega_+ - \omega_-)} \] (13)

\[ \frac{f(t) - f_0}{f_{ss} - f_0} = 1 - \frac{(\tau_p \omega_+ - 1) e^{-\omega_+ \tau} - (\tau_p \omega_+ - 1) e^{-\omega_+ \tau}}{\tau_p (\omega_+ - \omega_-)} \] (14)

with

\[ \omega_\pm = \pm \frac{\tau_p + \tau}{2 \tau_p} \pm \frac{\sqrt{(\tau - \tau_p)^2 - 4 \tau_\tau \tau_p}}{2 \tau_p} \] (15)

In these equations, \( f_0 = \tilde{\varepsilon}_s (v_{pol} - v_0) \) is the early time force calculated from eqn (11), where the initial spreading velocity, \( v_0 = \frac{\tilde{\varepsilon}_s v_{pol} - \tilde{\varepsilon}_s}{(h / R_0) \tilde{\varepsilon}_s + \tilde{\varepsilon}_s} \), is obtained from eqn (5) by setting \( u_{fr} = P = 0 \).

An explicit expression for the evolution of the myosin stress, \( P(t) \), can readily be derived by the usage of eqn (5) and (11). Analysis of these equations is presented next.

### III. Results and discussion

#### A. Effects of myosin activity on the steady-state cell radius and force

The formulation described above allows us to investigate the effects of myosin II activity on the extent and dynamics of cell spreading and force generation. As has already been mentioned above,
myosin activity plays a dual role in the overall force balance during spreading. While strengthening of myosin forces above the adhesion zone at the cell front facilitates spreading, the enhancement of myosin stress in deeper regions of the lamella opposes spreading. This may also be explained in terms of the effects of myosin activity on the effective rigidity of these two regimes. We note the appearance of the two effective moduli, \( \tilde{k}_c = (1 + \alpha)k_c \) and \( \tilde{k}_l = (1 + \beta)k_l \) in eqn (8) and (10). These provide a model for the effective stiffening of the two lamellar regions by myosin polarization behavior.\(^{14}\) Thus, on the one hand, via the effect of \( \beta \), the inhibition of myosin activity reduces the effective rigidity of the lamella front (\( \tilde{k}_l = (1 + \beta)k_l \)), this reduces the frictional interaction with the substrate and impairs the ability of the cell to spread. The active stiffening of the lamella front by myosin activity is thus seen to provide it with more strength to stretch the interior of the cytoskeleton forward, while inhibition has the opposite effect of loosening the lamella, as indeed often observed by its “floppy” appearance in the microscope.\(^{1,14}\) On the other hand, via the effect of \( \alpha \), the inhibition of myosin II motors loosens deeper regions of the lamella (\( \tilde{k}_c = (1 + \alpha)k_c \)) and this facilitates the elastic stretching of the cell. These two contrasting effects of myosin activity in the lamella are illustrated in panels a and c of Fig. 2.

**Fig. 2** The contrasting effects of myosin II motors in the lamellar bulk and front on cell spreading area and force. Left and right panels, respectively, show the effects of \( \alpha \) and \( \beta \) on the cell radius (a and c) and force (b and d). While strengthening of myosin contractility in the bulk opposes spreading (panel a), myosin motors above the adhesion zone at the cell front facilitate spreading (panel c). Because (and so long as) the lamellar actin network is free to slide in response to myosin tractions in the bulk, the contractile activity expressed by \( P(t) \) only affects the temporal force during spreading but does not alter the steady-state force exerted by the cell into the matrix (as seen in panel b). The only contribution to the steady-state force in this case comes from myosin motors above the adhesion zone that contribute to cell substrate friction. In panels a and b, blue, black and red curves, respectively, correspond to \( \alpha = 0, 0.5, 1 \) and \( \beta = 0, 0.5, 1 \) and \( x = 0 \). Other parameters are \( k_l = k_{xl}, t_0 = 0.6t_0, \) \( t_0 = 0 \), and \( \tilde{\alpha} = \xi_v^c/k_{xl} = 0.17t_0 \) where \( t_0 \) is the value of \( t \) for \( \beta = 0 \) and \( k_l = k_{xl} \). The scaled quantities are defined as follows: \( \tilde{R}(t) = (R(t) - R_0)/(R_m - R_0) \) and \( \tilde{R}(t) = (f(t) - f_0)/C_0 \), where \( R_m \) and \( R_0 \) are the steady-state radius and force in case that \( x = \beta = 0 \).

Interesting is the different way in which the two myosin populations affect the cellular force. Panel 2b shows that with higher values of \( \beta \) larger stresses develop during spreading but all curves converge to the same value at the steady-state. Myosin contractility in the lamellar bulk, \( P(t) \), is thus seen to only affect the temporal value of the cellular stress during spreading but to have no effect on the steady-state stress. The steady-state stress in our model is dictated by the frictional interaction \( z_v = \xi_v/B \) between the cell and the substrate which monotonically increases with myosin activity in the adhesion zone, as shown in panel 2d. As mentioned above, this conclusion is likely to hold so long as the cytoskeleton is free to flow past the substrate and contract; once long-lived stable contacts are established between the actin cytoskeleton and the ECM myosin contractility in the lamellar bulk would naturally also contribute to the generated force. The general increase of steady-state cellular stress with myosin activity is well known.\(^{11,14}\) Assuming \( 1 + \beta \) to inversely scale with blebbastatin concentration or volume ratio, \( B \) as \( 1/\tilde{z}_{bleb} \) one finds \( f_{ss} \sim 1/(k_{xl}/k_l + \tilde{z}_{bleb}) \) which is consistent with the results of Mitrossilis et al.\(^{10,13}\)

Similar structural arguments have been invoked to explain the effects of myosin IIA and IIB on cell spreading dynamics. While the inhibition of both resulted in a reduction of cellular force,\(^{11,14}\) it has been reported in ref. 23 and 24 that the depletion of the myosin IIB isoform resulted in impaired spreading while the depletion of myosin IIA facilitated spreading. Measurements of the spatial distribution of the two isoforms showed that the IIB protein is concentrated slightly more peripherally; while this picture is consistent with our analysis it must be emphasized that the spatial distribution of the two myosin isoforms shows significant overlapping which makes a quantitative interpretation nontrivial. The distribution of the two myosin isoforms is perhaps more apparent in neuronal cells. The myosin IIB isoform mainly concentrates at the peripheral zone of the growth cone whereas the myosin IIA isoform is found in more central locations of the growth cone and along the axon.\(^{29}\) Consistent with this spatial segregation, the inhibition of myosin IIB impedes neurite outgrowth\(^{25,26}\) while the myosin IIA drives neurite retraction.\(^{27,28}\) Our analysis does not distinguish between two myosin isoforms but illustrates that differences in their location may oppositely affect spreading.

Our theory also predicts the effects of matrix rigidity on the extent and dynamics of cell spreading. Consistent with experiments, eqn (8)–(10) predict that both the cell radius and force monotonically increase with matrix rigidity and combining eqn (5) and (11) reveals that the initial spreading speed is given by \( v_0 = \xi_v B \left( \left( k_{pol}/k_c \right) z_v + \xi_v \right) \), which in agreement with experiment is also an increasing function of matrix rigidity. A detailed examination of these predictions in comparison to experimental data is presented in ref. 8. Our extension here to account for myosin polarization response allows us to predict how the matrix rigidity might affect the responsiveness of cell spreading to myosin inhibition. On very stiff substrates eqn (10) predicts that \( z_v \sim 1 + \beta k_l \) on very soft substrates \( z_v \) is independent of \( \beta \). One consequently finds \( R_{ss} \sim (1 + \beta)/(1 + \beta) \) for very stiff substrates, and \( R_{ss} \sim 1/(1 + \beta) \) for very soft substrates. Thus in
the case of stiff substrates, myosin inhibition is expected to have a reduced effect on cell spreading in comparison to soft substrates where spreading is predicted to be facilitated. This is consistent with the recent observations of Dillard et al.\textsuperscript{21} with T cells spreading on mobile and fixed ligands on a supported lipid bilayer.

**B. Effects of myosin activity on cell spreading dynamics**

Spreading dynamics and force generation have been described by two major characteristic times, the spreading characteristic time, \( \tau \), and the polarization response time, \( \tau_p \), of myosin motors in the lamellar bulk; the less prominent consequences of myosin dynamics in the lamellar front are examined in the ESL.\textsuperscript{1} The frequencies \( \omega_\alpha \) and \( \omega_\beta \) are combinations of the two time scales that also depend on the polarizability factors \( x \) and \( \beta \); the spreading time, \( \tau(\beta) \), is generally an increasing function of \( \beta \) (via the dependence of \( \zeta_\alpha \) on \( \beta \)). While spreading experiments reveal that \( \tau \) is on the order of minutes (e.g., as in blood platelets) to tens of minutes (other cell types), we are still uncertain about the characteristic time of acto-myosin polarization; measurements of force generation in clamped cells held between cantilevers reveal an active polarization of force that occurs on a time scale of minutes to tens of minutes.\textsuperscript{13,26} This suggests a time scale for \( P(t) \), however, even in those experiments it is unclear to what extent do spreading forces contribute to the actual measurement. Our formalism here allows us to investigate how the ratio between \( \tau_p \) and \( \tau \) may be reflected in cell spreading dynamics.

We first examine the predicted behavior in the two extreme cases where: (i) cell spreading is faster than the polarization response, \( \tau < \tau_p \) and (ii) polarization response is faster than spreading, \( \tau_p < \tau \). In the former case (i) actomyosin polarization occurs with a delay relative to spreading. For sufficiently strong polarization response we find that \( R(t) \) and \( f(t) \) may evolve non-monotonically with time, showing a characteristic “overshoot” in early spreading, as exhibited by the black and red curves in Fig. 3a and b. Similar behavior has been observed, for instance, in ref. 22 (see Fig. 1c in that paper) as well as in our experiments on platelets (data not shown). Mathematically one finds that for long \( \tau_p \) and small enough \( x \) early spreading behaves as: \( R(t) \approx R_0 + (R_{ss} - R_0)(1 - e^{-\tau t}) \) and \( f(t) \approx f_0 + (f_{ss} - f_0)(1 - e^{-\tau t}) \). Thus, in early spreading the cell extends with minor (opposing) polarization forces, as if it tries to reach the steady-state cell size, \( R_{ss} \), that would be obtained with \( x = \beta = 0 \). However, at longer times, when \( t > \tau \), a gradual decrease in cell size may be observed once myosin motors start pulling on the cytoskeleton. Interestingly, this behavior may also be interpreted as an effective stiffening of the cytoskeleton from the initial value \( \kappa_\epsilon \) to the final value \( \kappa_\epsilon(1 + x) \). The gradual decrease in cell size may result in the overall decrease in cell force (e.g., red curve in panel 3b).

In the opposite case (ii) where acto-myosin forces polarize more quickly than cell spreading one finds: \( R(t) \approx R_0 + (R_{ss} - R_0)(1 - e^{-\tau_p t}) \). This is similar to the previous case, but here both the steady-state size \( R_{ss} \) and the time scale \( \tau = \tau_p(1 + x) \) are reduced by the cytoskeletal stiffening factors, \( 1 + x \) and \( 1 + \beta \) (cf. panel 3c). For the force we find the following behavior: \( f(t) \approx f_0 + (f_{ss} - f_0)[1 - (1 + (1/2 \tau_p) e^{-t/\tau_p} + (1/2 \tau_p e^{-t/\tau_p})^2] \). At early times when \( f(t) \) and its elastic contribution are still small, the acceleration of \( P(t) \) may dominate the behavior of \( f(t) \) (with large enough \( x \)) and give rise to an apparent sigmoidal appearance of \( f(t) \) (see for instance Fig. 3b and e, ESI).

Finally, eqn (15) predicts the occurrence of damped oscillations when the discriminant is negative (red curve in Fig. 3e and f). As the equation shows, the closer the time scales \( \tau \) and \( \tau_p \) are, the more likely are these oscillations to be observed. We find that for \( \tau_p \approx \tau \) the frequency of these oscillations scales as: \( \sqrt{1/x/\tau_\tau} \), i.e., increases with \( x \) and decreases with \( \tau_p \) and \( \tau \) as one expects. This predicted global oscillatory behavior of cell area should be distinguished from the more rapid protrusion–retraction cycles that occur locally at the cell periphery as reported for instance in ref. 15 and 57.

A phase diagram summarizing when damped oscillations of this sort are to be expected is shown in Fig. 4a for the case \( k_1 = k_{mi} \); the ordinate measures the ratio \( \tau_p/\tau_0 \) between the polarization time and the spreading time, where \( \tau_0 = \tau (\beta = 0) \), and the abscissa is the myosin susceptibility \( x = \beta \). Interestingly, the phase diagram predicts that substrate rigidity may oppositely affect the profile of spreading dynamics \( R(t) \) for cells located in the upper or lower halves of the phase diagram. Substrate rigidity is predicted to modulate the spreading time, \( \tau = 1/2 \kappa_\epsilon \left( \frac{\kappa_\epsilon + R_0}{h} \right) \), via the augmenting effect it has on \( \zeta_\alpha \) (see, eqn (10)). Hence, if a cell is located on the lower branch of the diagram (e.g., point a in panel 4a), a lowering of substrate rigidity will increase the ratio \( \tau_p/\tau_0 \) and cause the cells to enter the oscillatory
regime; as seen in panel 4b (dark-red shows regular spreading versus light-red that shows damped oscillatory behavior). The opposite effect is predicted for cells located on the upper branch of the phase diagram, see 4c; in this case an oscillatory mode (dark red curve) is observed on the more rigid substrate. Note that all curves in panels 4a and b were deliberately scaled by the steady-state change in cell radius, $R_{ss} - R_0$, which by eqn (8) is an increasing function of matrix rigidity; hence all curves converge to 1 at $t \to \infty$. Complementary calculations of $R(t)$, $t_p$ and $R_0$ for the dynamics observed in panels b and c are presented in Fig. S2 (ESI†).

Fig. 2–4 show an interesting means of investigating the contribution of myosin activity to cell spreading dynamics. It is predicted that the stimulation of myosin activity would increase the likelihood of observing non-monotonic spreading patterns that are otherwise more rarely observed; this is illustrated by the effect of increasing $\alpha$ or $\beta$ on the evolution of cell area and force. Furthermore, our prediction of a phase diagram that distinguishes different effects of substrate rigidity on the cell spreading profile (panels b and c in Fig. 4) suggests another interesting way to investigate the coupling of myosin polarization dynamics to spreading induced stresses. To this end, however, it is essential for first experimentally quantifying the time scale of actomyosin polarization in the course of spreading ($t_p$). One way to achieve this is by measuring the course of force generation in cells whose spreading area has been restricted using the micro-patterning technique. An experiment in this spirit has been reported by Aratyn-Schaus et al. who used blebbistatin washout to set the initiation of myosin force generation in an already full spread cell; these authors reported a time scale of a few minutes for myosin force generation in U2OS osteosarcoma cells.

IV. Concluding remarks

We presented a theoretical analysis of the contribution of myosin II generated stress to the extent and dynamics of cell spreading, and of the force generation in this process. The theory distinguishes two myosin populations according to their location in the lamella. Since cell-substrate adhesions tend to concentrate near the cell periphery we used a coarse grained picture and distinguished between myosin motors above the adhesion zone and those that locate more deeply in the lamella. The two myosin populations are shown to affect cell spreading in opposite ways. Those located above the adhesion zone facilitate spreading since they effectively strengthen the frictional interaction between the cell and the substrate and more efficiently allow the cell to extend forward. Myosin II motors in the lamella bulk have the opposite effect of pulling the cell inwards, thereby resisting cell spreading. We have demonstrated that the time scale of actomyosin polarization plays a crucial role in shaping the time course of cell spreading and predicted a phase diagram which shows the conditions for the occurrence of global oscillatory changes in cell area. The closed form expressions we derived for the evolution of cell area and force provide a simple quantitative framework for analyzing the contribution of myosin dynamics during spreading.

Acknowledgements

Funding to SK was provided by the Deutsche Forschungsgemeinschaft (DFG) in the framework of the SFB 937/A12 and the Excellence Initiative. AZ thanks the Israel Science Foundation (grant no. 1396/09) and the Niedersachsen, German-Israeli Lower Saxony Cooperation for their support.

References

1 U. S. Schwarz and M. L. Gardel, J. Cell Sci., 2012, 125, 3051.
2 N. Wang, I. M. Tolic-Norreykje, J. X. Chen, S. M. Mijailovich, J. P. Butler, J. J. Fredberg and D. Stamenovic, Am. J. Physiol.: Cell Physiol., 2002, 282, C606.
3 J. P. Califano and C. A. Reinhart-King, Cell. Mol. Bioeng., 2010, 3, 68–75.
4 T. Yeung, P. C. Georges, L. A. Flanagan, B. Marg, M. Ortiz, M. Funaki, N. Zahir, W. Ming, V. Weaver and P. A. Janmey, Cell Motil. Cytoskeleton, 2005, 60, 24.
5 C. Reinhart-King, M. Dembo and D. Hammer, Biophys. J., 2005, 89, 676.
6 L. Trichet, J. Le Digabel, R. J. Hawkins, S. R. K. Vedula, M. Gupta, C. Ribraut, P. Hersen, R. Voituriez and B. Ladoux, Proc. Natl. Acad. Sci. U. S. A., 2012, 109, 6933.
7 S. Schwarz G. Henriques, R. Sandmann, A. Strate and S. Köster, J. Cell Sci., 2012, 125, 3914.
8 N. Nisenholz, K. Rajendran, Q. Dang, H. Chen, R. Kemkemer, R. Krishnan and A. Zemel, Soft Matter, 2014, 10, 7234–7246.
9 S. Yang and T. Saif, Exp. Cell Res., 2005, 305, 42–50.
10 J. Étienne, J. Fouchard, D. Mitroffisli, N. Bufo, P. Durand-Smet and A. Asnacios, Proc. Natl. Acad. Sci. U. S. A., 2015, 112, 2740.
11 Y. Cai, N. Biais, G. Giannone, M. Tanase, G. Jiang, J. M. Hofman, C. H. Wiggins, P. Silberzan, A. Buguin and B. Ladoux, Biophys. J., 2006, 91, 3907.
12 M. L. Gardel, B. Sabass, L. Ji, G. Danuser, U. S. Schwarz and C. M. Waterman, J. Cell Biol., 2008, 183, 999.
13 D. Mitroffisli, J. Fouchard, A. Guirou, N. Desprat, N. Rodriguez, F. Babry and A. Asnacios, Proc. Natl. Acad. Sci. U. S. A., 2009, 106, 18243.
14 Y. Cai, O. Rossier, N. C. Gauthier, N. Biais, M. A. Fardin, X. Zhang, W. W. Miller, B. Ladoux, W. W. Cornish and M. P. Sheetz, J. Cell Sci., 2010, 123, 412.
15 G. Giannone, B. J. Dubin-Thaler, H.-G. Döbereiner, N. Kieffer, A. R. Bresnick and M. P. Sheetz, Cell, 2004, 116, 431.
16 G. Johnson, L. Leis, M. Krumwiede and J. White, J. Thromb. Haemostasis, 2007, 5, 1516.
17 F. Rehfeldt, A. E. Brown, M. Raab, S. Cai, A. L. Jazay, A. Zemel and D. E. Discher, Integr. Biol., 2012, 4, 422.
18 Y. Qiu, A. C. Brown, D. R. Myers, Y. Sakurai, R. G. Mannino, R. Tran, B. Ahn, E. T. Hardy, M. F. Kee and S. Kumar, et al., Proc. Natl. Acad. Sci. U. S. A., 2014, 111, 14430.
19 T. Wakatsuki, R. W. Wyssolmerski and E. L. Elson, Cell Sci., 2003, 116, 1617.
20 J. D. Mih, A. Marinkovic, F. Liu, A. S. Sharif and D. J. Tschumperlin, J. Cell Sci., 2012, 125, 5974.
21 P. Dillard, R. Varma, K. Sengupta and L. Limozin, Biophys. J., 2014, 107, 2629.
22 S. Kumari, S. Vardhana, M. Cammer, S. Curado, L. Santos, M. P. Sheetz and M. L. Dustin, Front. Immunol., 2012, 3, 230.
23 V. Betapudi, L. S. Licate and T. T. Egelhoff, Cancer Res., 2006, 66, 4725.
24 V. Betapudi, PLoS One, 2010, 5, e8560.
25 S. R. Wylie, P.-J. Wu, H. Patel and P. D. Chantler, Proc. Natl. Acad. Sci. U. S. A., 1998, 95, 12967.
26 P. C. Bridgman, S. Dave, C. F. Asnes, A. N. Tullio and R. S. Adelstein, J. Neurosci., 2001, 21, 6159.
27 M. Amano, K. Chihara, N. Nakamura, Y. Fukata, T. Yano, M. Shibata, M. Ikebe and K. Kaibuchi, Genes Cells, 1998, 3, 177.
28 S. R. Wylie and P. D. Chantler, Mol. Biol. Cell, 2003, 14, 4654.
29 D. M. Suter and K. E. Miller, Prog. Neurobiol., 2011, 94, 91.
30 U. S. Schwarz, T. Erdmann and I. B. Bischofs, BioSystems, 2006, 83, 225.
31 P. Marcq, N. Yoshinaga and J. Prost, Biophys. J., 2011, 101, L33.
32 C. Borau, T. Kim, T. Bidone, J. M. Garca-Aznar and R. D. Kamm, PLoS One, 2012, 7, e49174.