Active Mechanics Reveal Molecular-Scale Force Kinetics in Living Oocytes

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ABSTRACT Active diffusion of intracellular components is emerging as an important process in cell biology. This process is mediated by complex assemblies of molecular motors and cytoskeletal filaments that drive force generation in the cytoplasm and facilitate enhanced motion. The kinetics of molecular motors have been precisely characterized in vitro by single molecule approaches, but their in vivo behavior remains elusive. Here, we study the active diffusion of vesicles in mouse oocytes, where this process plays a key role in nuclear positioning during development, and combine an experimental and theoretical framework to extract molecular-scale force kinetics (force, power stroke, and velocity) of the in vivo active process. Assuming a single dominant process, we find that the nonequilibrium activity induces rapid kicks of duration $\tau \sim 300 \mu$s resulting in an average force of $F \sim 0.4$ pN on vesicles in in vivo oocytes, remarkably similar to the kinetics of in vitro myosin-V. Our results reveal that measuring in vivo active fluctuations allows extraction of the molecular-scale activity in agreement with single-molecule studies and demonstrates a mesoscopic framework to access force kinetics.

INTRODUCTION

Living cells utilize motor proteins to actively generate force at the molecular scale to drive motion and organization in the crowded intracellular environment (1–4). For example, force generation is critical to facilitate the basic tasks of living cells such as spatial organization, motility, division, and organelle transport. Recently, active diffusion has emerged as an important process in cell biology (5–9). Active diffusion is the random motion of intracellular components that is driven by active metabolically powered forces (and not by thermal fluctuations). An interesting example occurs during early vertebrate development, namely, oocyte meiosis, where active diffusion drives nucleus centering (5).

Oocytes are immature female gametes that are destined to be fertilized and grow into a fully functioning organism. They are large cells (80 $\mu$m in diameter) of spherical geometry with a thick actin-rich cortex and a well-separated cytoplasmic skeleton (Fig. 1 A). The cytoplasmic skeleton of a mouse oocyte is a composite material that includes actin filaments, microtubules, and intermediate filaments. The actin network is composed of long unbranched filaments polymerized from the surface of vesicles that harbor actin nucleating factors (10). These vesicles act as nodes to create an interconnected actin network that is uniform in density throughout the cytoplasm and unpolarized (11), with microtubules that are organized in small seeds and do not form long filaments during prophase-I (12); and although it is probable that intermediate filaments are present (13), little is known about their structure or function in mouse oocytes.

Correct nuclear positioning during prophase-I requires precise spatial and temporal coordination of the cytoskeleton and molecular motors (11,14). Recent work has shown the importance of mechanical processes in oocyte development (5,11,14). Proper spindle positioning requires the oocyte cortex to soften and exhibit plastic deformation (14), and motion of the cytoplasmic actin network is closely regulated by myosin-V motors (11). Motion of large objects, such as positioning of the spindle (14) or nucleus (5), must be driven by a physical force generated by nonequilibrium...
processes. For instance, we have recently shown that in prophase-I, myosin-V-driven vesicle motion maintains a soft cytoskeleton and generates a force to center the nucleus (5). Here we focus on the physical understanding of the active forces that generate vesicle motion, and how the detailed quantification and modeling of this activity can successfully predict characteristic parameters of the active forces, such as timescales, velocities, and effective forces.

To investigate the nonequilibrium mechanical activity in living cells, it is necessary to independently measure the active force generation and the local mechanical properties to understand how objects move within the complex intracellular environment (15–24). This approach was introduced to study active processes in hair bundles (15) and stress fluctuations in cells (16). Subsequent studies utilized this concept to investigate cross-linked actin-myosin-II networks in reconstituted systems (17), beads embedded in living cells (18–21), and red blood cell membranes (22). Alternative methods of estimating forces in vivo include FRET force sensors, droplet-based sensors, and force inference as recently reviewed (23,24).

In this study, we measure the actively driven motion of endogenous vesicles in the largely unexplored cytoplasmic skeleton of in vivo mouse oocytes and demonstrate a framework to quantify the underlying driving forces. Endogenous vesicles embedded in the oocyte cytoplasm experience local elastic caging. The vesicles undergo actively driven motion via a sparse actin-myosin-V force-generating network that remodels the environment. The actin-myosin-V network is the key element in generating force for nuclear positioning (5), yet the underlying physical processes related to active force generation are not well understood.

To investigate cytoplasmic force generation, we combine optical tweezer-based active microrheology (AMR) and laser-tracking interferometry with our theoretical framework to quantify the in vivo active mechanical processes and relate them to the underlying stochastic force kinetics (amplitude and temporal correlations). To study the activity

**FIGURE 1** Intracellular mechanics surrounding endogenous vesicles in living oocytes. (A) Mouse oocytes are large spherical cells that have a well separated cortex and cytoplasmic-skeleton composed of biopolymer filaments (bright-field image, left; scale bars, 20 μm). During prophase-I it contains a dynamic actin-myosin-V meshwork that drives vesicle motion (schematic shown in center and fluorescent image of actin filaments on right; scale bars, 5 μm). Endogenous vesicles embedded in the cytoplasmic-skeleton are trapped using optical tweezers (zoomed inset). (B) Once a vesicle is trapped, the mechanical properties of the local environment can be measured by AMR where a sinusoidal oscillating force is applied to the vesicle (blue) and the resulting displacement of the vesicle is measured (green). The viscoelastic shear modulus ($G^*$) is calculated from this force-displacement measurement via the generalized Stokes-Einstein relation. (C) The mechanical properties surrounding vesicles in the cytoplasmic-skeleton of oocytes exhibit power-law behavior with frequency scaling $G^* \propto f^{0.75}$. This shows that the cytoplasmic-skeleton in oocytes can be modeled as a semiflexible polymer network. To see this figure in color, go online.
in mouse oocytes, we examine the cytoplasmic-skeleton during prophase-I, when the actin-myosin-V network is the dominant source of activity (5). By using statistical mesoscopic measurements on endogenous vesicles, and theoretical modeling, we extract the molecular-scale force kinetics (average power-stroke duration, displacement, force, and velocity) of in vivo myosin-V.

**MATERIALS AND METHODS**

**Fmn2−/− and myosin-V dominant negative mouse oocytes**

To study the effect of the actin-myosin-V network, we utilize two conditions to independently perturb the actin cytoskeleton (Fmn2−/−) and the myosin-V activity (MyoV−/−). Fmn2−/− mouse oocytes have no detectable cytoplasmic actin filaments, as confirmed by several independent studies (14,25–27). The reduced cytoplasmic actin has also been confirmed via cytochalasin-D-treated mouse oocytes, which exhibit the same phenotype and mechanical properties as Fmn2−/− (28) (Fig. S1). For myosin-V dominant negative experiments (MyoV−/−), oocytes were injected with cRNAs using a Femtotjet microinjector (Eppendorf, Hamburg, Germany) as published previously (5). Oocytes were kept in prophase-I arrest for 1–3 h to allow expression of fusion proteins. The myosin-Vb dominant negative construct corresponds to a portion of the coiled-coil region of the myosin-Vb that mediates dimerization of the motor (10). We believe this construct binds to the coiled-coil region of endogenous myosin-Vb, resulting in impaired motor dimerization. Thus, myosin-Vb is still able to bind to vesicles and actin filaments but does not actively generate force. It is specific to myosin-Vb and has been shown to stop vesicle motion in mouse oocytes as efficiently as brefeldin A, which is a general traffic inhibitor (5). (Note that MyoV−/− oocytes were referred to as “WT + MyoVb tail oocytes” in a previous study (5).)

**Oocyte preparation**

Oocytes were collected from 11-week-old mice OF1, 13-week-old C57BL6 (wild-type), or 15-week-old Fmn2−/− female mice as previously described (29) and maintained in prophase-I arrest in M2 + BSA medium supplemented with 1 μM Milrinone (30). Live oocytes were embedded in a collagen gel to stop movement of the overall cell during measurements. Collagen gel was made by mixing M2 medium (33.5 μL), 5× PBS (10 μL), NaOH (1 M, 0.9 μL), and collagen (3.6 mg/mL, 55.6 μL) to obtain 100 μL of the final collagen solution at 2 mg/mL with a pH ∼ 7.4. Twenty microliters of the collagen solution were deposited on a coverslip and live oocytes were added. The droplet was covered with another glass coverslip using vacuum grease (Dow Corning, Midland, MI) to minimize evaporation. The collagen gel containing oocytes was polymerized in a humid environment at 37°C for 30 min.

**Optical tweezer setup**

The optical tweezer system utilizes a near-infrared fiber laser (λ = 1064 nm, YLM-1-1064-LP; IPG Photonics, Oxford, MA) that passes through a pair of acoustooptical modulators (AA-Optoelectronics, Orsay, France) to control the intensity and deflection of the trapping beam. The laser is coupled into the beam path via dichroic mirrors (ThorLabs, Newton, NJ) and focused into the object plane by a water immersion objective (40×, 1.2 NA; Olympus, Tokyo, Japan). The condenser is replaced by a long-distance water immersion objective (40×, 0.9 NA; Olympus) to collect the light and imaged by a 1:4 telescope on a InGaAs quadrant photodiode (G6849 QPD; Hamamatsu, Hamamatsu City, Japan). The resulting signal is amplified by a custom-built amplifier system (Oeifier Electronics, Heidelberg, Germany) and digitized at a 500-kHz sampling rate, 16 bits, using an analog input card (PCIe-6353; National Instruments, Austin, TX). All control of the experimental hardware is executed using the software LabVIEW (National Instruments). Optical trapping of endogenous vesicles was calibrated similarly as in (31,32), where the high-frequency fluctuations (f > 500 Hz) from the fluctuation and response measurements are aligned to determine the trap stiffness (18,33). This approach assumes that the high-frequency motion is dominated by thermal fluctuations so that the fluctuation dissipation theorem (FDT) holds in this regime. For direct measurement of violation of the FDT, laser tracking interferometry is used first to measure the spontaneous fluctuations of the vesicle, followed immediately by active microrheology to measure the mechanical properties of the local environment surrounding the vesicle.

**Laser tracking interferometry**

The position of the endogenous particle is measured by back-focal-plane-plane interferometry (34). It should be noted that deformable objects (e.g., giant unilamellar vesicles 10–100 μm) undergo shape fluctuations that will manifest in the voltage measured by the QPD (35). For small endogenous particles (~1 μm), it has been confirmed that shape fluctuations are small and laser interferometry can be used to track their position with nanometer precision. This has been verified in mammalian cells (36) and yeast (37). A recent study has also confirmed that laser tracking interferometry and the active-power spectrum calibration (31) allow quantitative force-displacement measurements on vesicles in vivo (38). To confirm the validity of this assumption in mouse oocytes, we investigated the deformability of endogenous vesicles and conclude that the majority of measured fluctuations come from active vesicle displacements (see Figs. S4–S9). Additionally, because the endogenous vesicles serve as nodes integrated in the cytoplasmic meshwork, they are accurate reporters of the network mechanics and fluctuations (39).

**Data analysis**

In the AMR experiments, we apply a known force, Fext, to an endogenous vesicle and measure the resulting displacement, u. Using linear response theory, these are related to the material response function χ(t) as u(t) = u(t) = Fext(t)/c. In Fourier space, we can directly calculate the response function as χ(ω) = (ω)/Fext(ω). To translate the response function into a complex shear modulus G*, we use the generalized Stokes-Einstein relation G* = 1/(6πkT/ω), where k is the average radius of the spherical tracers. It can be split into a real part G and an imaginary part G'', which, respectively, provide information about the elasticlike and viscouslike properties of the medium. For AMR measurements, the laser power exiting the objective was ~120 mW. For the spontaneous fluctuations, we measure the motion, u(t), of endogenous vesicles via laser interferometry (without trapping the vesicle, laser power ~1 mW) and calculate the power spectral density (PSD), C(ω) = (u(t)u(t))exp(iωtdt), as described previously (35). Briefly, we calculate the PSD by using the software MATLAB (The MathWorks, Natick, MA) to take the fast Fourier transform (FFT) of the vesicle position, u(t) = FFT(u). Then the PSD is calculated as PSD = (ω/π) / (ω/π) / (ω/π), where the asterisk denotes complex conjugate, ω is the number of measurement points, and is the sampling frequency. In systems at thermal equilibrium, the response function can be calculated from the PSD via the FDT as is done for passive microrheology (PMR) (40). Due to the particle size and range of the QPD measurement, our PMR measurements are restricted to a particle displacement of ~500 nm during the measurement interval. If the particle left this regime, that part of the data was not analyzed. The limitation to the linear regime of the detector does not imply that we measure only the linear regime of the
force displacement relation. AMR and PMR data analysis is carried out as done previously (22).

RESULTS

Intracellular mechanics of the oocyte cytoplasmic-skeleton

To measure the local mechanical properties, we optically trap an endogenous vesicle embedded in the cytoplasmic-skeleton, apply an oscillatory force while measuring its displacement (Fig. 1 B), and calculate the shear modulus via the generalized Stokes-Einstein relation (41). Because vesicles serve as integrated nodes in the cytoplasmic skeleton, they accurately reflect the mechanics and fluctuations of the network (39) (details on vesicle tracking in the Supporting Material). To calibrate the force on the vesicles, we exploit the established observation that high-frequency fluctuations are thermal in origin (18,31,33). AMR directly measures the mechanical response function (\(\chi\)), which allows us to determine the complex modulus (\(G^*\)) characterizing the viscous and elastic resistance of the composite cytoplasmic skeleton including any contribution from actin, intermediate filaments, microtubules, and other structures present. We find that the cytoplasmic skeleton of oocytes has strongly viscoelastic properties similar to semiflexible biopolymer networks (42,43), exhibiting high-frequency power-law behavior (\(G^* \propto f^\alpha\), where \(\alpha \sim 0.75\)) (Fig. 1 C). Thus, the cytoplasmic-skeleton in oocytes is strongly frequency-dependent, which must be accounted for when modeling its mechanical behavior. Extending the theoretical model into the viscoelastic regime with a strong frequency dependence is a necessary advancement to describe the intracellular mechanics.

To dissect the mechanical contributions of cytoskeletal filaments in oocytes, we first perform AMR on formin-2 knockout oocytes (Fmn2\(^{--}\)), which removes cytoplasmic actin filaments nucleated by formin-2 (25), as confirmed via cytochalasin-D experiments as an alternative method of removing actin (Fig. S1). We find that the mechanical properties do not change compared to WT, indicating that the actin network does not provide significant mechanical resistance in the oocyte (Fig. 2, A and B). This behavior is presumably because the actin network in WT oocytes is sparse (Fig. 1 A, right and Fig. S2), with a mesh size of 5.7 \(\pm\) 1.9 \(\mu\)m (26). Depolymerizing the microtubules by nocodazole treatment (1 \(\mu\)M) does not significantly affect the mechanical properties (Fig. 2, A and B), indicating that microtubules also do not provide significant mechanical resistance in the cytoplasmic-skeleton. This result is expected because prophase-I microtubules form small seeds instead of long force-bearing tubule structures (12). These combined results suggest that there are other mechanical scaffolds contributing to the stiffness in the cytoplasm of mouse oocytes.

Although our results show that actin does not contribute significantly to the mechanical resistance, it is known that the actin-myosin-V meshwork activity plays a critical role for self-organization, such as positioning of the meiotic spindle (25,26) and nucleus centering in mouse oocytes (5). To independently probe the activity of this network, we impaired the myosin-V motor force generation by deactivating it via microinjection of a dominant negative construct (MyoV \((-\))\) (5,10) (see Materials and Methods). Under these conditions, the mechanical properties of the cytoplasmic-skeleton stiffens significantly as evidenced by
the shift upwards in the elastic and viscous moduli (Fig. 2, A and B). This increased mechanical stiffness correlates with a higher density actin meshwork that is observed in the absence of myosin-V activity (Fig. S2), and can be explained by increased cross-linking (11). Actin networks are known to stiffen significantly upon addition of cross-links, as long as the inter-cross-link distance is smaller than the persistence length.

The overall behavior of the cytoplasmic skeleton in oocytes is strongly viscoelastic, where neither the elastic nor viscous components dominate by a large margin. At the lowest probed frequencies, the behavior is slightly more elastic, with a crossover (~10–20 Hz) to more viscous at higher frequencies. However, both components of the shear moduli are always within a factor of two at the observed frequencies (Fig. 2 C). This is in contrast to adherent cells, which typically show an overall weaker frequency dependence and are dominantly elastic (18,44).

Quantifying in vivo active force fluctuations

To quantify nonequilibrium activity, we use laser-tracking to precisely measure the spontaneous motion of vesicles in the cytoplasmic skeleton with high spatio-temporal resolution (34,36,37) (see Materials and Methods). Both the spontaneous motion and the local mechanics are measured for each individual vesicle via active microrheology and laser interferometry. This allows direct comparison between the local mechanical environment experienced by the vesicle and the active force that is driving its motion because both measurements are made in situ. This is critical to extract information about the molecular-scale processes. In the absence of biochemical activity, the motion of the vesicles would be due to purely thermal agitation, which is fully determined by the mechanical properties of the system. This basic relation is given by a fundamental theorem of statistical mechanics known as the FDT (40). The FDT relates the small fluctuating motion of the vesicles to the mechanical properties of the surrounding environment by 

$$\chi'' = \pi f \tilde{C}/k_B T$$

where $\chi''$ is the dissipative part of the mechanical response, $\tilde{C}$ is the power spectral density of vesicle motion, $f$ is the frequency, $k_B$ is the Boltzmann constant, and $T$ is the temperature (see Table 1 for a list of symbol definitions). However, in living cells, the presence of biochemical activity gives rise to active forces (e.g., a nonequilibrium process) driven by energy consuming processes (15,16,45). In other words, the force driving the motion of particles in a living cell (e.g., the oocyte) has two contributions: 1) a passive (purely thermal) contribution described by classical equilibrium physics; and 2) an active contribution that is biochemically regulated and cannot be understood via equilibrium physics.

We quantify and explain both the passive and active contributions driving intracellular fluctuations, by independently measuring the response function and power spectral density. We use AMR to measure $\chi$ and laser-tracking to measure $\tilde{C}$, and use this information to check for violation of FDT (15,17,46), which indicates active force generation. In oocytes, at high frequencies the mechanical response ($\chi''$) and the spontaneous vesicle motion ($\pi f \tilde{C}/k_B T$) are related by the FDT, as expected for thermal fluctuations (18,33) (Fig. 3 A). At frequencies <~400 Hz, the observed motion of vesicles is dominated by an active energy-consuming process (highlighted by the pink shaded region between the two curves in Fig. 3 A).

To quantify the nonequilibrium activity in an active soft material, it is instructive to consider the effective energy (15,47–49), $E_{\text{eff}} = \pi f \tilde{C}/\chi''$, which is a measure of how far the system is from thermal equilibrium. Note that the measurement of $E_{\text{eff}}$ requires two separate measurements: rheology to get $\chi''$ and laser interferometry to get $\tilde{C}$. WT oocytes exhibit a strong departure from equilibrium due mainly to the actin-myosin-V network activity. Accordingly, the deviation is reduced when either actin is absent (Fmn2−/−) or myosin-V is inactivated (MyoV −/−) (Fig. 3 B). This is quantitative confirmation that the dynamic actin-myosin-V meshwork drives vesicle dynamics out-of-equilibrium in the cytoplasmic skeleton of mouse oocytes (5,11). Note that in Fmn2−/− and MyoV −/− not all activity is abolished, which is presumably due to residual

| Symbol | Description | Units |
|--------|-------------|-------|
| $\chi$ | response function | m/N |
| $\omega$ | frequency | rad/s |
| $\gamma(t)$ | memory kernel | N/m |
| $G^*(\omega)$ | complex shear modulus | N/m² |
| $G'(\omega)$ | elastic modulus | N/m² |
| $G''(\omega)$ | viscous modulus | N/m² |
| $R$ | vesicle radius | m |
| $\epsilon$ | local cage stiffness | N/m |
| $\tau_{\text{p}}$ | passive timescale | s |
| $\alpha$ | power-law exponent | |
| $\lambda(t)$ | vesicle position | m |
| $\lambda_0(\tau)$ | local cage position | m |
| $\xi(t)$ | thermal noise | m/s |
| $\nu_0(t)$ | active burst noise | m/s |
| $T$ | bath temperature | K |
| $T_{\lambda}$ | active temperature | K |
| $\tau_0$ | active persistence time | s |
| $v$ | active waiting time | m/s |
| $\tilde{C}$ | active burst amplitude | m²/s |
| $E_{\text{eff}}(\omega)$ | position power spectral density | N-m |
| $S_{\text{eff}}(\omega)$ | effective energy | N²-s |
| $S_{\text{thermal}}(\omega)$ | total cell force | N²-s |
| $S_{\text{active}}(\omega)$ | cell force spectrum | N²-s |
| $F$ | thermal force spectrum | N |
| $v_{\text{vesicle}}$ | active force spectrum | m/s |
| $\Delta \epsilon$ | active force amplitude | m |
| $\Delta \epsilon$ | average vesicle velocity | m/s |

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FIGURE 3 Active force generation by myosin-V drives the cytoplasmic-skeleton out-of-equilibrium. (A) At frequencies below 400 Hz the spontaneous motion of vesicles in WT oocytes (green circles) is larger than expected for thermal equilibrium (blue circles). This shows that active forces are contributing to vesicle motion in this regime as highlighted by the red shaded region. This is direct evidence of nonequilibrium behavior in the cytoplasmic-skeleton (via violation of the FDT). At high frequencies the observed vesicle motion resembles thermal motion indicated by the blue-shaded region. (B) WT oocytes (blue) are the furthest from equilibrium as shown by their higher effective energy. In the absence of actin (red) or when myosin-V is inactivated (black) the dynamic actin-myosin-V meshwork is compromised, and oocytes have lower effective energy. Solid lines are theoretical fits (see equations in the Supporting Material; error bars = mean ± SE). (C) The cell force spectrum (S_{cell}) experienced by vesicles is the sum of thermal forces (S_{therm}) and active forces (S_{active}). At high frequencies S_{cell} (green) is dominated by S_{therm} (blue) and the two spectra coincide (blue shaded region). At lower frequencies S_{cell} is larger than S_{therm} showing the existence of additional active forces. Solid and dashed lines are theoretical predictions (see equations in the Supporting Material).

low-frequency deviation is due to simple power-law model. Standard error of experimental data is within marker size. (D) When a representative trajectory (black) is filtered to remove the high-frequency thermal fluctuations the result is a smoothed trajectory (green) that represents actively driven motion. The difference between the true trajectory (gray) and the smoothed trajectory (green) recovers the high-frequency thermal fluctuations (blue) shown in (E). (F) The MSD of vesicles, calculated from trajectory data, indicates they undergo random-confined motion in the oocyte cytoplasmic-skeleton at short timescales. This behavior transitions to active diffusion at longer times (C), and is reminiscent of cytoplasmic stirring (33). To see this figure in color, go online.

actin-myosin-V that is active or other remaining sources of nonthermal activity in the oocyte (e.g., enzymes, polymerization, myosin-II in the cortex (11)).

To develop a more intuitive picture of the activity, we quantify the forces generated in the cell by calculating the cell force spectrum (S_{cell}). The S_{cell} directly represents the average total force on a vesicle from all stochastic sources (active and thermal) inside the cell, and has been used by several previous studies to quantify activity (16,18,19,46). The force spectrum is a convenient and common way to quantify the total intracellular forces. In analogy to the force on a simple spring, where the force is the stiffness multiplied by displacement (F = κΔx, where κ is stiffness and Δx is displacement), we calculate S_{cell} = \frac{6πR^2}{\kappa} |G''|^2 \tilde{C}, where |G''|^2 represents the stiffness of the cytoplasmic-skeleton (18,19,50). In our framework we separate the total force spectrum in the cell to be the sum of thermal forces and active forces (S_{cell} = S_{therm} + S_{active}) (46,51). The thermal force spectrum is calculated directly from the AMR measurements as S_{therm} = 12πRG''k_BT/ω, where R is the vesicle radius, G'' is the dissipative component of the shear modulus, and ω = 2πf is frequency. S_{therm} represents the forces present at thermal equilibrium and is a direct consequence of the FDT (40). The resulting force spectrum is shown in Fig. 3 C, where at high frequencies the cell force spectrum (S_{cell}) is dominated by thermal forces (where S_{active} << S_{thermal} as expected, but at lower frequencies (f < 400 Hz) active forces dominate (S_{active} >> S_{thermal}), which results in higher total force experienced by vesicles in the cytoplasm over thermal equilibrium (lines represent the theoretical model (52) introduced in the next section). To visualize the effect of thermal and active forces, we show a representative trajectory of a vesicle (Fig. 3 D, black). Based on our measured violation of FDT (Fig. 3 A), we apply a filter to isolate the low-frequency (<400 Hz) vesicle motion that is mainly due to active processes (Fig. 3 D, green), and the high-frequency motion that is dominated by thermal fluctuations (Fig. 3 E).

Modeling actively driven vesicle motion

Mouse oocytes in prophase-I exhibit nondirected motion of vesicles in a viscoelastic environment (Fig. 3 F, blue) due to a unique cytoplasmic network (Fig. 4 A). We use this picture to motivate our theoretical model of generic active forces in the cell cytoplasm and do not impose any physical assumptions on the origin of that force. Therefore, our model lumps all active forces together as a single dominant process, regardless of their origin. To gain access to the molecular-scale kinetics of the dominant active process driving the
cytoplasmic interior of oocytes, we developed a quantitative model describing vesicles embedded in a viscoelastic environment that are subjected to thermal and active forces. Here, the nonequilibrium processes provide the active forces that reorganize the polymer cytoskeleton and drive motion of the vesicles. Our theoretical framework extends previous approaches for near-elastic networks (46,50,53) to include the complex mechanical properties measured experimentally (Fig. 2).

We begin by using the traditional Langevin equation to describe a vesicle freely fluctuating in a viscoelastic continuum where the left side of Eq. 1 is the drag force on the vesicle and the right side includes the elastic caging force from the cytoskeleton (with stiffness $k$) and the thermal force. It is an application of Newton’s second law on the fluctuating vesicle. Equation 1 describes how a vesicle thermally fluctuates randomly about its equilibrium position, but on average will not move from its original position due to elastic caging of the cytoskeleton:

$$\int \gamma(t - t') \dot{x}(t') = -k(x - x_0) + \xi,$$

where $x$ is the position of the vesicle; $x_0$ is the position of the elastic cage; $\xi$ is thermal force taking the form of Gaussian colored noise with correlations $\langle \xi(t)\xi(t') \rangle = k_BT \Gamma(1 - \alpha)$, as provided by the FDT (40); $k_B$ is the Boltzmann constant; and $T$ is the bath temperature of the environment. The viscoelastic material properties are modeled as a soft glassy material, where $\kappa$ is the harmonic trap stiffness centered at $x_0$, and the memory kernel is $\gamma(t) = \kappa(\xi_0/t)^{\alpha}/\Gamma(1 - \alpha)$.

**FIGURE 4** A theoretical model of active mechanics connects in vivo measurements to molecular force kinetics. (A) Vesicles (dark gray) are embedded in the complex mechanical continuum (gray background) of the oocyte interior. Actin filaments emanate from the surface of vesicles creating an entangled network (green, left). Myosin-V motors generate force on actin (magenta, right) giving rise to forces throughout the network driving random motion of vesicles. (B) We model vesicles embedded in a mechanical continuum with local cage stiffness ($k$) represented by the blue harmonic potential, viscoelastic dissipation ($\zeta_\alpha$), and thermal fluctuations ($\xi$) (left). Active processes rearrange the network through bursts of motion ($v_A$), resulting in displacement of the local cage (blue harmonic), which generates the active force ($k\zeta_\alpha v$, red arrow) that drives vesicle motion toward the local minimum (right) (inset indicates active force kinetics). (C) The active force spectrum ($S_{active}$) quantifies the forces on vesicles due to only active processes. Combined with our quantitative model we find that the vesicles are subjected to 0.4 pN of force, during a power stroke of length $\Delta x \sim 20$ nm and duration $300 \mu s$, resulting in a vesicle velocity of $320$ nm/s, which is strikingly similar to the kinetics measured for single molecule myosin-V in vitro and the in vivo vesicle velocity. The solid line is the theoretical fit (Eq. 7), and the dotted line is a $-4/3$ power law consistent with the cytoplasmic-skeleton mechanics. (D) Simulated vesicle motion (green squares) agrees with experimental data (blue circles) for a wide range of timescales as shown by the probability distribution of displacements. This includes long timescale ($t_{pdf} = 1$ s) Gaussian behavior and short timescale ($t_{pdf} = 0.01$ s) non-Gaussian tails (indicated by red-shaded regions) that suggest molecular motor behavior, where $t_{pdf}$ is the time-lag for calculation of the displacement correlations. Gaussian distributions shown in black. These results show that molecular level kinetics of active processes can be extracted from mesoscopic in vivo measurements. To see this figure in color, go online.
where $\Gamma$ is the Gamma function, $\alpha$ is the viscoelastic power-law frequency dependence, and $\zeta_\alpha$ is a passive timescale of the material (see Table 1 for symbol definitions and the Supporting Material for details). In an attempt to capture the active vesicle dynamics, we introduce Eq. 2, which phenomenologically describes how nonthermal forces remodel the cytoplasmic-skeleton and determine the dynamics of the elastic caging, as

$$\int dt' \gamma(t-t')x_0(t') = \kappa \zeta_\alpha v_A,$$

where the left side is the drag force on the cage and the right side is the active nonthermal force that drives motion of the cage. Equation 2 is an application of Newton’s second law on the elastic cage, which is a geometric constraint, and fundamentally does not have mass. In this case, the cage represents the confining effects of the cytoplasmic skeleton and thus is endowed with the mass of this surrounding material and its center of mass corresponds to the center of the cage. The equilibrium position of the cage is $x_0$ and the active force is $\kappa \zeta_\alpha v_A$, where $v_A$ is the velocity of the cage motion. For completeness, Eq. 2 should include the force applied on the cage by the vesicle (the action-reaction principle) and the thermal force on the cage itself. However, when each increment of vesicle motion is small compared to the cage size, these forces do not affect the cage dynamics to leading order and previous work has suggested they can be neglected (see Eq. 3, a and b, in (50)). The overall dynamics describe a vesicle in the cytoplasmic-skeleton that experiences an elastic caging (Eq. 1), until active forces act to remodel the cytoskeleton and move the cage (Eq. 2), resulting in a vesicle motion driven by both thermal and active forces. This can be considered a coarse-grained approach, where the dynamics are divided into two equations of motion describing the thermal fluctuations (Eq. 1) and the active motion (Eq. 2) to simplify the mathematics and subsequent analysis. Therefore, this phenomenological model can describe diffusive-like motion driven by nonthermal forces in a viscoelastic medium, also known as “active diffusion.” Interestingly, a recent study used a coarse-grained two-fluid continuum model (54) and arrived at the same conclusion (46).

The parameters characterizing the mechanical properties are obtained from AMR measurements. The memory kernel quantifies the mechanical resistance of the cytoplasmic-skeleton and influences the dynamics of both the vesicle, $x$, and the cage, $x_0$. In principle, one would expect the memory kernel in Eqs. 1 and 2 to be different because Eq. 1 describes motion of the vesicle, and Eq. 2 describes motion of the cage. Provided that the dissipation is mediated by the surrounding medium in both cases, then the memory kernels would only differ by a prefactor. In this model, the prefactor ends up on both sides of Eq. 2 and mathematically cancel, leaving the memory kernels in Eqs. 1 and 2 identical (see previous derivation (50)).

The microscopic interpretation of $\kappa$ is the local caging due to the cytoskeleton (as evident from the low-frequency plateau in the measured shear modulus Fig. 2), where $x_0$ is the location of this cage. In the absence of active forces, the position of this local cage does not move ($\dot{x}_0 = 0$). When active forces do exist, as in living cells, they rearrange the cytoskeleton and drive motion of the cage ($\dot{x}_0 \neq 0$). The active force experienced by the cage results from the active burst velocity, $v_A$, which is a stochastic process consisting of alternating active and quiescent periods. During an active phase, the active burst velocity, $v_A$, takes on a constant nonzero value, $r$, for a random exponentially distributed time of average value $\tau$. During a quiescent phase, the active burst velocity is 0 for an exponentially distributed random time of average value $\tau_0$ (46,48,50). A schematic example is shown in Fig. 4 B (right inset). The active burst velocity is a zero-mean stochastic process with correlations $\langle v_A(t)v_A(0) \rangle = k_B T_A \exp(-|t|/\tau)/(|\zeta_\alpha\tau|)$, where $k_B T_A = k \zeta_\alpha (\tau)^2/[3(\tau + \tau_0)]$ defines an active energy scale. Note that the dynamics of the joint process $\{x,x_0\}$ is not Markovian because of the memory kernel in the equations of motion.

From the generalized Stokes-Einstein relation, the memory kernel can be associated with a complex shear modulus $G' = G' + i G''$ of the form

$$G'(\omega) = \frac{k}{6\pi \mu} \left[ (\omega \zeta_\alpha)^2 \frac{\cos(\pi \alpha/2)}{2} + i \right],$$

$$G''(\omega) = \frac{k}{6\pi \mu} (\omega \zeta_\alpha)^2 \sin(\pi \alpha/2).$$

This model separates the passive forces that originate from the thermal fluctuations of the medium and the active forces that depend on the energy-consuming processes in the active material. Note that if the active force on the cage is zero ($\zeta_\alpha v_A = 0$), then Eq. 2 is zero, and the equation of motion simplifies to confined viscoelastic Brownian motion (Eq. 1). If the active force is not zero, then it originates from an active process that has steplike velocity kinetics (Fig. 4 B, right inset). These force kicks drive the nonequilibrium fluctuations in our model.

The nonequilibrium properties of the active force are quantified by the deviation from the FDT defined by a frequency-dependent effective energy, $E_{\text{eff}}$, as discussed earlier. We compute it in terms of the microscopic ingredients of our theoretical model,

$$E_{\text{eff}}(\omega) = k_B T + \frac{1}{(\omega \zeta_\alpha)^2 \sin(\pi \alpha/2)} \frac{k_B T_A}{1 + (\omega \tau)^2/4}.$$
where $k_B T_A$ is the energy scale associated with the active process. The dynamics of the tracer can be written as: $x(t) = \int dt' x(t - t')F_{\text{cell}}(t')$, where $F_{\text{cell}} = \xi + \kappa x_0$ is the cell force that describes the thermal forces arising in the cell and the effect of the active forces via $x_0$. Explicitly, we calculate the cell force spectrum as $S_{\text{cell}} = \langle |\tilde{F}_{\text{cell}}|^2 \rangle = \langle |\tilde{\xi}|^2 \rangle + \kappa^2 \langle |\tilde{x}_0|^2 \rangle$, which corresponds to the spectrum calculated previously as $S_{\text{cell}} = \tilde{C}/|\tilde{x}|^2 = (6\pi R)^2 G^2 \tilde{C}$ from experiments. Because we separate the thermal and active contributions to the force spectrum in our model, we compute the explicit expressions for each as

$$S_{\text{therm}}(\omega) = 2k_B T \kappa \zeta_x \omega^2 \sin \left( \frac{\pi \alpha}{2} \right), \quad (6)$$

$$S_{\text{active}}(\omega) = \frac{2k_B T \kappa \zeta_x}{(\omega \zeta_x)^2 \kappa T} \left[ \frac{k_B T \zeta_x}{1 + (\omega \tau)^2} \right]. \quad (7)$$

The active force spectrum (Eq. 7) can now be used to extract the kinetics of the active force. Details of the derivation can be found in the Supporting Material.

### Extracting in vivo molecular-scale force kinetics

By combining our analytical model and experimental measurements, we extract the force kinetics driving active mechanical processes in the oocyte. First, the mechanical measurements from AMR were fit to Eqs. 3 and 4 to determine the mechanical parameters: $\alpha$, $\zeta_x$, and $\kappa$ from the data shown in Fig. 2A and B. Once the mechanics is determined, the remaining equations are largely constrained. The best fit of the data for effective energy (Fig. 3B) is used to determine $k_B T_A$ in Eq. 5, where the fit is independent of $\tau$ when $\tau < 10$ ms. From fitting the active force spectrum, we find that $\tau = 0.3$ ms (the only free parameter) to capture the high-frequency dropoff (Fig. 4C). The fit parameters for the theoretical model applied to all experimental conditions are shown in Table 2. To summarize: $\alpha$, $\zeta_x$, and $\kappa$ are obtained from fitting AMR measurements of the mechanics ($G^2, G''$), $T_A/\kappa$ is obtained from fitting the effective energy data ($E_{\text{cell}}$), and $\tau$ is obtained from fitting the active force spectrum ($S_{\text{active}}$). Our model extracts the molecular-scale force kinetics directly from fitting the active force spectrum.

### DISCUSSION

**Myosin-V activity maintains a soft cytoplasmic-skeleton**

Our mechanical measurements (AMR) indicate that when myosin-V activity is present, the cytoplasmic-skeleton in oocytes is softer. This observation was first reported by Almonacid et al. (5), and the new data in this study extends this
observation across a wide range of frequencies. Additionally, the combined measurements of viscoelastic properties and spontaneous vesicle motion provides access to an independent measurement of mechanics and activity. The mechanism of how softening occurs is not yet clear, but may be related to the structural connectivity of the actin-myosin-V network. In our experiments, it is possible that when myosin-V is inactivated (MyoV ($-$)), it serves as a passive cross-linker that stiffens the environment or cross-links the vesicle to the network. Because cross-linking leads to highly nonlinear stiffening (59) (if cross-link distance is not large compared to the persistence length), a small change in cross-linking could effectively increase the stiffness of the cytoplasmic-skeleton. Alternatively, the presence of myosin motors may reduce the mechanical contribution of the actin cytoskeleton by causing fluidization (see (60)) so that other intracellular structures (e.g., intermediate filaments) dominate the mechanics in the WT condition. This interpretation is also consistent with the finding that the removal of the cytoplasmic actin has no effect on the shear modulus. However, in the absence of experimental tools to dissect the mechanical contribution of all cytoplasmic components in the mouse oocyte, the answer to this question is currently out of reach.

Active force spectrum reveals molecular-scale kinetics of the active process

Myosin-V is typically considered a directed transport motor. However, in prophase-I mouse oocytes, myosin-V drives vesicle motion in a nondirected fashion as evidenced by its sublinear scaling of the MSD (Fig. 3 F, blue). Previous work has also shown that myosin-V drives random motion of vesicles (active diffusion) on longer timescales of minutes (5). To further illustrate this point, Fig. 4 A (left) shows a schematic illustration of vesicles and actin filaments embedded in the oocyte interior. The complex mechanical environment is represented by a generic continuum (gray background). When myosin-V is attached to a vesicle and applies force on an actin filament, this force is transduced through the actin filament to a neighboring vesicle where it is bound by formin-2 (e.g., black arrows in Fig. 4 A, right). This process occurs frequently throughout the oocyte interior, resulting in randomly distributed force-dipoles (similar to myosin-II in some systems (17,33)). Thus, each vesicle experiences forces in random directions due to the action of myosin-V throughout the network. This is represented by many actin filaments under force, as shown in magenta in Fig. 4 A (right). Therefore, due to the unique network connectivity in the cytoplasmic-skeleton, myosin-V motors drive active random motion in prophase-I mouse oocytes reminiscent of cytoplasmic stirring observed previously that was driven by myosin-II (33). This actin-myosin-V activity is expected to be the dominant driver of active diffusion in our system (5).

By looking at the active force spectrum in WT oocytes, we can characterize the kinetics of the dominant active process. The first factor in Eq. 7 depends on the mechanical properties of the cytoplasmic-skeleton whereas the second factor is related to the kinetic properties of the active processes. This illustrates that the active force spectrum is dependent on the active force generation as well as the mechanics of the environment it must push against. The power-law scaling of the active force spectrum contains information about the underlying physics. At lower frequencies the active force spectrum scales as $f^{-2}$, reflecting the active forces pushing against the viscoelastic environment. At higher frequencies the active force spectrum is dominated by the kinetics of the active force generation process, which scales as $f^{(-2\alpha+2)}$. This is illustrated in Fig. S3, where at low frequencies there is a power-law dependence of $-4/3$ consistent with the mechanics ($\alpha \approx 2/3$) and at high frequencies the power-law dependence is $\sim -10/3$ consistent with kinetics and mechanics. The crossover between these two regimes occurs at $1/\tau$ as shown approximately by the purple dotted line (Fig. S3).

Our predicted active force spectrum exhibits power-law scaling that is dependent on the mechanical properties of the system ($\alpha$), in contrast to previous studies (16,53,61). Previous theoretical developments predict an active force spectrum that scales as $f^{-2}$, which is independent of the mechanical properties (16), and a plateau ($f_0$) below a critical frequency (Eq. 2 in (33) and Eq. S1 in (18)). It is worth noting that in the near-elastic case ($\alpha \approx 0$), our model recovers the behavior observed in previous work (18,33). The divergence of the active force spectrum at low frequencies is consistent with our experimental measurements, and others, where a low-frequency plateau is not observed (18,19,21,62).

The experimental measurements for the active force spectrum ($S_{active} = S_{cell} - S_{term}$) and theoretical model are compared in Fig. 4 C. The low-frequency power-law behavior is clearly seen, whereas at higher frequencies the active forces drop off rapidly, which may reflect molecular motor statistics (63). This allows us to calculate other kinetic parameters, such as the apparent active force amplitude, $F$, which is the typical force amplitude that an endogenous vesicle feels that drives its nonthermal motion:

$$F = \left[3k_BT_A(\tau + \tau_0)/\zeta_a\right]^{1/2},$$

where $\tau_0$ is deduced from the single-molecule myosin-V duty cycle (64) (see details in the Supporting Material). Here, $F$, lumps all activity into one value to quantify the active force. Once this force is known it is also straightforward to calculate the typical vesicle velocity,

$$v_{vesicle} = \frac{F}{k_B \zeta_a},$$
which is a ratio of the driving force, and the resistance provided by the surrounding environment, $\kappa_S^\omega$. Using this argument, the typical vesicle displacement induced by one force kick can be computed as $\Delta x = F/\kappa$. Thus, once the timescale of the force kick ($\tau$) is extracted from Eq. 7, the apparent force felt by the vesicle ($F$), the expected vesicle velocity ($v$), and the step-size $\Delta x$ can be deduced.

Our combined experimental and theoretical framework allows access to the molecular-scale kinetics via mesoscopic measurements of the active force spectrum (Fig. 4 C). The active force spectrum, $S_{\text{active}}$, has only one free fitting parameter, $\tau$, which corresponds to the molecular-scale activity. All other model parameters are fixed previously. In living oocytes we find that endogenous vesicles experience a force of $F \sim 0.4$ pN during a power stroke of $\tau \sim 300$ $\mu$s duration, with a step-size of $\Delta x \sim 20$ nm. This is strikingly similar to single molecule myosin-V kinetics measured in vitro (see Table 3) (63–69). In addition, we find that the predicted average vesicle velocity due to active forces is $v_{\text{vesicle}} \sim 320$ nm/s, which is in agreement with myosin-V velocity in vitro (65–67), as well as the velocity of myosin-V driven vesicles measured in in vivo oocytes via video microscopy (5,10). Whereas our model does not assume the action of a single motor acting on the vesicle, the extracted kinetics are in agreement with single myosin-V. Because it is likely there are several motors affecting the vesicle dynamics, our results suggest that the myosin-V motors in prophase-I oocytes do not act simultaneously, thus allowing individual motor kicks to be observed. Interestingly, previous work has estimated the number of motors involved based on quantification of energy dissipation (52). It is important to note that although our extracted kinetics are in agreement with myosin-V measurements, this is not sufficient to prove that we are accessing single-molecule kinetics. Together, our results suggest that force kinetics of in vitro myosin-V is remarkably similar to the active force fluctuations in in vivo oocytes, where they drive the composite cytoplasmic-skeleton out-of-equilibrium (Table 3).

### Success and limitations of this theory and experiments

Our phenomenological model is designed to capture the nonequilibrium fluctuations of vesicles in the oocyte cytoplasm. Specifically, this model is able to reproduce the deviation from equilibrium as measured by the effective energy and active force spectrum. In that respect, the agreement between model and experiments supports the underlying phenomenological picture that: 1) active fluctuations of the vesicles are mediated by the surrounding medium; 2) the complexity of the surrounding medium can be reduced to a coarse-grained representation described by local caging, 3) the damping force acting on the vesicle and the cage exhibit similar memory effects. Although this minimal approach is completely phenomenological, several studies have shown that introducing a fictitious cage coordinate has been a powerful framework for describing intermittent dynamics of a particle in a damped medium (70). And in the case of mouse oocytes, it is able to capture the nonequilibrium fluctuations of vesicles and predict force kinetics that are in agreement with single-molecule experiments on myosin-V motors. Nevertheless, this coarse-grained approach inherently neglects the microscopic details of the underlying processes.

The main drawbacks of our minimal model are: 1) We neglect any collective effects that may be occurring due to multiple motor interactions. If several motors are interacting to generate a force kick, we only capture their combined action and no information about individual motors. 2) All material properties are taken as an input to the model (as measured from experiments). Thus, the model is not able to capture how the motors may affect the material properties (e.g., cannot explain softening/stiffening (71)). Therefore it is critical that the material properties of the medium are measured in situ, and the quality of the measurements will strongly affect the model. 3) Our coarse-grained interpretation involves representing the cytoskeleton as a local cage constraining vesicle motion. Active motion of the local cage obeys Newton’s second law, which typically describes the motion of a specific material point and its associated mass. In this case, because the local cage is not an actual physical object, it’s mass is associated with the material that is surrounding the vesicle. This interpretation is necessary to apply Newton’s second law to the active motion in our phenomenological model, and similar approaches inspired by the itinerant oscillator model have been used previously (70). Together, these drawbacks mask the underlying microscopic processes and future work will focus on gaining access to these details.

To spur future developments and improve this type of analysis we would like to point out the current limitations of the theory and experiments. In theory, our phenomenological approach to describe the cage dynamics (Eq. 2) does not track a specific material point in the cytoskeleton, making it difficult to interpret the physical basis of the memory kernel; the active process driving nonequilibrium motion is dominated by a single timescale, which may not be true in other cell types; and the number of motors driving the vesicle dynamics is not specified, because this information is typically not known in vivo. Overcoming these limitations in the theory is challenging, because tracking a

### TABLE 3 Molecular-Scale Force Kinetics

|                  | In Vivo Oocytes | Single-Molecule | Reference |
|------------------|-----------------|-----------------|-----------|
| $F$ (pN)         | 0.4             | 0–4             | (63–66)   |
| $\tau$ (ms)      | 300             | 160–1000        | (64,67,68)|
| $\Delta x$ (nm)  | 20              | 15–25           | (64,67–69)|
| $v_{\text{vesicle}}$ (nm/s) | 320            | 270–480         | (65–67)   |
specific material point in an active material (that is constantly changing) may not be possible, and introducing multiple timescales and details about motors requires more specific information about the experimental system at hand. A useful first step could be to develop a new metric to characterize the mechanical response of active materials that does not rely on the material itself remaining invariant. In experiments, force calibration relies on the high-frequency collapse of fluctuation and response, which requires precise measurement that is often not accessible in many systems; measurements of vesicle motion are limited to a relatively small range (≤400 nm), due to the linear range of the QPD; and rheological measurements are limited to frequencies greater than ~1 Hz, due to large-scale motion in the cell. These experimental limitations could be overcome by implementing new force calibration techniques and a feedback system that continually refocuses the optical trap on the vesicle.

CONCLUSIONS

We quantify the molecular-scale force kinetics of active diffusion in mouse oocytes via experiments, theory, and simulation. We find that active forces in in vivo oocytes are remarkably similar to myosin-V kinetics in vitro, and that the in vivo kinetics can be extracted from cytoplasmic fluctuations. Our results demonstrate a framework for connecting cellular scale phenomena to their underlying molecular force kinetics in vivo and provide insight on the kinetic origin of active diffusion in mouse oocytes.

SUPPORTING MATERIAL

Supporting Materials and Methods and nine figures are available at http://www.biophysj.org/biophys/supplemental/S0006-3495(18)30212-1.

AUTHOR CONTRIBUTIONS

W.W.A. and T.B. conceived and supervised the project. W.W.A., M.B., M.A., and M.-H.V. performed experiments. É.F., N.S.G., P.V., and F.v.W. developed the theoretical model and performed simulations. W.W.A. and É.F. integrated experiment and theory. All authors contributed to data analysis and/or interpretation. W.W.A. and T.B. wrote the manuscript, which was seen and corrected by all authors.

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