Deciphering dynamics of clathrin-mediated endocytosis in a living organism

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Current understanding of clathrin-mediated endocytosis (CME) dynamics is based on detection and tracking of fluorescently tagged clathrin coat components within cultured cells. Because of technical limitations inherent to detection and tracking of single fluorescent particles, CME dynamics is not characterized in vivo, so the effects of mechanical cues generated during development of multicellular organisms on formation and dissolution of clathrin-coated structures (CCSs) have not been directly observed. Here, we use growth rates of fluorescence signals obtained from short CCS intensity trace fragments to assess CME dynamics. This methodology does not rely on determining the complete lifespan of individual endocytic assemblies. Therefore, it allows for real-time monitoring of spatiotemporal changes in CME dynamics and is less prone to errors associated with particle detection and tracking. We validate the applicability of this approach in vivo systems by demonstrating the reduction of CME dynamics during dorsal closure of Drosophila melanogaster embryos.

Introduction

Clathrin-mediated endocytosis (CME) is the major pathway responsible for internalization of lipids and receptor-bound macromolecules from the plasma membrane of eukaryotic cells (Conner and Schmid, 2003). During internalization of a cargo molecule, clathrin triskelions assemble into submicron-sized polyhedral structures upon their recruitment to the plasma membrane by the endocytic clathrin adaptor protein AP2 (Ehrlich et al., 2004; Saffarian and Kirchhausen, 2008; Boucrot et al., 2010; Cocucci et al., 2012; Hong et al., 2015). Live-cell imaging studies designed for tracking of fluorescently tagged clathrin coat components have revealed the dynamics of formation, internalization, and dissolution of distinct classes of clathrin-coated structures (CCSs; Gaidarov et al., 1999; Merriefield et al., 2002; Ehrlich et al., 2004; Loerke et al., 2009; Mettlen et al., 2010; Taylor et al., 2011; Kural and Kirchhausen, 2012; Aguet et al., 2013). The best-characterized structures are highly curved, cage-like assemblies that deform the plasma membrane into pits and vesicles. In conventional fluorescence time-lapse acquisitions, clathrin-coated pits appear as diffraction-limited spots with mean lifetimes of ~1 min (Ehrlich et al., 2004; Saffarian et al., 2009; Kural et al., 2012; Aguet et al., 2013). CCSs disappearing within the first ~20 s of their initiation are abortive structures that fail to construct bona fide endocytic carriers (Hong et al., 2015). Flat arrays of clathrin, also known as plaques, are larger than coated pits and slower in their internalization dynamics (Saffarian et al., 2009; Grove et al., 2014). Physiological relevance of clathrin-coated plaques has been equivocal, because they only appear at the substrate contact sites of cultured cells and, because of their long lifetimes, they are not effective endocytic carriers.

Dynamics of endocytic pathways are inversely related to plasma membrane tension, because membrane internalization machinery are required to do work against the two major constituents of tension (i.e., in-plane tension and membraneskeleton adhesion) to create invaginations (Dai et al., 1997; Raucher and Sheetz, 1999; Sheetz, 2001; Apodaca, 2002; Gauthier et al., 2012; Diz-Muñoz et al., 2013). Tension regulates formation and curvature of clathrin coats reconstructed on giant unilamellar vesicles (Saleem et al., 2015). Studies in yeast and in polarized and mitotic mammalian cells show that CME is inhibited unless plasma membrane tension is counteracted by actin dynamics (Aghamohammadzadeh and Ayscough, 2009; Boulant et al., 2011; Kaur et al., 2014). Regulation of endocytic rates by mechanical cues has important roles in development; during the early stages of Drosophila melanogaster embryogenesis, increased tension inhibits Fog receptor endocytosis, which is required for completion of ventral furrow formation (Pouille et al., 2009).

Our current understanding of CME dynamics is based on in vitro imaging studies that are limited in their potential to mimic physical properties of tissue microenvironments. In...
Results

CCS growth rates are robust reporters of CME dynamics

In fluorescence imaging assays, endocytic CCS formation is marked by appearance of a diffraction-limited spot that steadily increases in intensity because of accumulation of fluorescently tagged coat components. The internalization of the clathrin-coated vesicle is followed by a relatively fast dimming of the fluorescence caused by dissolution of the coat (Kural and Kirchhausen, 2012; Fig. 1, A and B). Lifetime (i.e., the time required for formation and internalization of CCSs from the plasma membrane) is an extensively used metric for characterizing CME dynamics. Factors that elongate CCS lifetimes reduce the efficacy of ligand endocytosis (Cureton et al., 2010; Boulant et al., 2011). Any factor that affects the rate of formation and/or dissolution of clathrin coats is a potential regulator of CCS lifetimes and, hence, CME dynamics. To establish whether we can use growth rates of CCS signal as metrics for endocytic dynamics, we quantified formation and dissolution (negative growth) rates of individual CCSs by determining the slope of the normalized fluorescence intensity within 12-s temporal windows (Fig. 1 C and online supplemental software).

Because hundreds of CCS traces can be detected within a single cell at a given instant, we were able to assemble distributions of growth rates for each frame of time-lapse acquisitions (Video 1). When applied to cells that produce predominantly clathrin-coated pits, the majority of the growth rates obtained in this way were positive (i.e., corresponding to increasing fluorescence signal; Fig. 2 B). This is consistent with the characteristic intensity profile of pits, which has a steady formation phase followed by a relatively abrupt dissolution (Ehrlich et al., 2004; Massol et al., 2006; Fig. 1 B).

As a means to alter CME efficiency, we used micropipette aspiration to increase in-plane membrane tension in cells (Houk et al., 2012). We detected a significant increase in CCS lifetimes when micropipette aspiration was applied (45.5 ± 27.1 s versus 80.1 ± 86.6 s, N_cells = 7, N_traces= 38,136; Fig. 2 A). Asymmetry of the growth rate distributions is abolished upon aspiration, and a greater fraction of intensity traces were associated with steady levels of fluorescence signal (i.e., the plateau phase). High-magnitude slopes, which represent fast formation and dissolution phases, are diminished with the increasing membrane stiffness.
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FIGURE 2. Using growth rate histograms as reporters of clathrin dynamics. (A) Kymographs are generated from the same BSC-1 cell before and during microaspiration, respectively. Elongated AP2 traces demonstrate longer CCS lifetimes under increased membrane tension. (B) Growth rate distributions are shown for seven different cells before and during aspiration. Change in CCS dynamics induced by micropipette aspiration can be observed in growth rate distributions. In the control experiments (cells before aspiration), CCSs spend more time in the formation phase (i.e., the distribution is inclined toward positive slopes). The asymmetry was abolished upon aspiration and plateau phases got relatively longer ($N_{\text{cells}} = 7$ and $N_{\text{traces}} = 38136$). (C) Growth rate distributions in B are assembled in five bins to better delineate different phases of clathrin-coated vesicle formation (FD, fast dissolution; FF, fast formation; P, plateau; SD, slow dissolution; SF, slow formation). The bars show mean ± standard deviation to illustrate dispersion between cells. P-values were obtained using the two-tailed t test. (D) 2D histograms of normalized intensity traces aligned at different time points (beginning, trace maximum, and end) and superposed as represented by the cartoons. In each alignment, the aspirated cells show a significantly widened distribution, demonstrating a preference for slopes lower in magnitude. Bins corresponding to multiples of 12 s are more populated, as they contain trace data from both 3- and 4-s frame rate acquisitions.

Errors associated with particle detection and tracking result in significant underestimation of CCS lifetimes (Aguet et al., 2013; Mettlen and Danuser, 2014). Slope values populating the growth rate histograms, however, are calculated using trace information within 12-s-long windows and thus do not rely on determining the complete CCS traces. We tested the reproducibility of growth rate distributions on 24-s-long integrated movie segments (IMSS) that are the quadrature sum of four different temporal sections of a movie (Fig. 4, A–C). Faithful assessment of CCS lifetimes from IMSS is impractical because (1) density of CCSs is on average four times greater than the original acquisition, (2) background noise level is increased due to error propagation, and (3) IMSS are significantly shorter than the mean coated pit lifetime (24 s versus 1 min). Despite these impediments, we found that CCS growth rate distributions could be reproduced very accurately from the IMSS (Fig. 4, B and C). We also found that distortion of the growth rate distributions caused by increased tension could be observed in the
Figure 3. CCSs in proximity of substrate adhesion sites have reduced dynamics. (A) Arrowheads mark clathrin-coated plaques at the ventral surface of a BSC-1 cell stably expressing AP2-GFP. Bar, 5 µm. (B) Scatterplot shows maximum intensity versus the axial positions of CCSs detected at the ventral surface. The positions are relative to the lower bound of the confocal volume used for 3D tracking (Kural et al., 2012). CCS traces are divided into plaque (red) and pit (green and blue) populations based on their maximum intensities. Clathrin pits positioned closer to the substrate are in green (N_plaques = 6 and N_traces = 11,482). (C) Scatterplot shows the lifetime versus axial positions of the CCSs shown in B. (D) Growth rate distributions of the pit and plaque populations. The inset shows the distributions as bar plots.

Figure 4. Growth rate analysis does not depend on determining the complete traces of CCSs. (A, left) Thumbnails represent 12 temporal sections (24 s each) of a 288-s-long fluorescence movie of a BSC-1 cell stably expressing AP2-GFP. (right) CCS growth rate distribution of the entire movie. (B, left) Integrated movie segments (IMSs) are the quadrature sum of four 24-s-long segments in which density of CCSs is approximately four times larger than the original movie. Temporal separation is maximized between the segments to minimize the number of self-overlapping clathrin spots. (right) Growth rate distributions obtained from each IMS is plotted in comparison with the distribution obtained from the entire movie (black). (C) Growth rate distributions of the IMSs and the entire movie are displayed as bar plots. (D) We tested the reproducibility of growth rate distributions using the CCS traces produced by cmeAnalysis (Aquet et al., 2013) and TraCKer (homemade particle tracking program; see online supplemental software). Changes in slope distributions upon micropipette aspiration can be observed in both datasets. Numbers of the analyzed IMSs are 25 and 64 for before and during aspiration conditions, respectively (N_cells = 7). Error bars show standard deviations. FD, fast dissolution; FF, fast formation; P, plateau; SD, slow dissolution; SF, slow formation.
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IMSs of microaspirated cells regardless of the algorithm used for tracking CCSs (Fig. 4 D).

Spatiotemporal variations in CME dynamics can be resolved in real time using CCS growth rates

When CCS lifetime distributions are used as metrics for CME efficiency, the mean lifetime of clathrin-coated pits (~1 min) sets the limit of the temporal resolution for distinguishing the changes in clathrin coat dynamics. Consequently, the factors that influence endocytic processes at shorter time scales become indiscernible. Growth rate distributions, however, are constructed before the completion of traces and thus enable real-time monitoring of CCS dynamics in cells. We used growth rate analysis to monitor gradual changes in CCS dynamics upon acute cholesterol depletion. Structural studies show that cholesterol depletion reduces CME efficiency by replacing clathrin-coated pits with flat clathrin arrays (plaques) at the cell membrane (Subtil et al., 1999; Fig. S2 A). When cells are treated with methyl-β-cyclodextrin (MβCD), in line with our micropipette aspiration experiments, we found that impairment of CCS dynamics is coupled with disappearance of high-magnitude slopes in CCS growth rate distributions (Fig. 5, A and B; and Video 2). This result was anticipated, as cholesterol depletion increases the effective membrane tension through escalation of membrane–cytoskeleton adhesion energy (Khatibzadeh et al., 2012; Fig. S2 B). We used standard deviation of regional CCS growth rates to generate a visualization tool for monitoring spatiotemporal changes in clathrin dynamics in real time (Fig. 5 C). In regions with impeded CCS dynamics, standard deviations of the growth rates were reduced because of the disappearance of the high-magnitude slopes (Fig. 5 D). Using this approach, we found that MβCD treatment can affect CME dynamics at different rates in neighboring cells (Fig. 5 D, E). Cell-to-cell variation may stem from cells’ mitotic state or position with respect to cell islet edges, as described earlier (Snijder et al., 2009).

Reduction in CCS formation rates (positive slopes) upon cholesterol depletion could be attributed to depletion of free clathrin coat components in the cytosol caused by increased CCS lifetime, as they are contained within existing coats. In that case, we would expect dissolution rates to disappear the earliest. However, we found that both positive and negative slopes (i.e., fast formation and dissolution phases) diminish simultaneously (Fig. 5 B), which indicates that flat clathrin arrays have slower...

Figure 5. Real-time monitoring of CCS dynamics in cholesterol-depleted cells. (A) Fluorescence images show snapshots of three BSC-1 cells stably expressing AP2-GFP. 10 mM MβCD was applied at the sixth minute (t = 360 s) to initiate cholesterol depletion (Video 2). Dashed lines demarcate the cell boundaries. Bar, 10 µm. (B) For the three cells shown in A, percentage frequency of the five growth phases are plotted as a function of time. The dashed lines denote the time point of MβCD addition. The fastest change in CCS dynamics is observed in cell 1. (C) Figures show standard deviation of local clathrin growth rates for the snapshots shown in A. Each pixel in the image is given the value of standard deviation of the growth rates detected from CCSs found in a radius of 4.8 µm. This representation is used for illustrating temporal and spatial variations in CCS dynamics. Standard deviation is lower in regions where the growth rate distribution is narrow, a signature of impeded CCS dynamics. Cell 1 responds to cholesterol depletion the earliest, as observed at t = 816 s. (D) Growth rate histograms are shown for the three cells at the time points of the snapshots in A and C. Spatial and temporal heterogeneity of CCS dynamics can be detected using the standard deviation of growth rate histograms, which change at different rates for the three cells. (E) Plots show spatial variation in standard deviation of local CCS growth rates over the distances shown by the dashed arrows in C. Positions of the cell boundaries are marked by the dashed red lines.
formation rates than clathrin-coated pits (Fig. S2 A). Therefore, at variance with earlier interpretations (Subtil et al., 1999), our results indicate that flat clathrin arrays found in cholesterol-depleted cells cannot be considered as direct precursors of clathrin-coated pits, as they have distinct formation dynamics.

**CCS intensity profiles can be reproduced from growth rate distributions**

Hierarchical clustering of CCS traces allows us to determine growth rate distributions corresponding to different intensity profiles (Fig. 6 A). An interesting question that arises is whether we can reverse the flow of information in this process (i.e., reproduce the mean intensity profiles using growth rate distributions). Our IMS analyses show that growth rate distributions can be accurately assembled even in the absence of complete CCS intensity profiles (Fig. 6 B). Therefore, whatever the design, such a methodology would be immensely useful for interpreting the CCS growth rates that are obtained from acquisitions in which monitoring complete CCS traces is infeasible. We used a strategy based on determining the analogous growth rate distributions within a library of CCS clusters (Fig. 6 C). We assembled a library of growth rate histograms that are obtained from 361 clusters containing a total count of 6,958 CCS traces. When we used test clusters obtained from various cellular contexts, we were able to make accurate predictions of their intensity profiles solely by determining the most analogous growth rate distribution within the cluster library (Fig. 7 A and online supplemental software). When applied to cells coexpressing fluorescently tagged AP2 and clathrin, we found that the AP2 growth rate distribution has higher frequencies at the low-magnitude slope region, and its predicted intensity profile has a longer plateau before dissolution (Fig. 7 B). This result is in accord with the previous studies, which show that AP2 fluorescence growth halts earlier than the clathrin signal during the completion of the coat (Saffarian and Kirchhausen, 2008).

When cells are treated with MJCD, incremental changes in mean CCS intensity profiles taking place within short intervals could be resolved in real time (Fig. 7 C and Video 2). Upon cholesterol depletion, intensity profiles predicted using growth rate distributions became longer lived and reached higher maximum intensities in time. This was expected, as clathrin-coated pits are gradually replaced by larger and less dynamic CCSs in these cells (Fig. S2 A). Although the extent of the cluster library is a critical determinant of the fidelity of predicted intensity profiles, the presented strategy is a powerful tool for analyzing the changes in CCS growth rates obtained from in vivo datasets.

**CME dynamics slow down during dorsal closure of Drosophila embryos**

Taking advantage of the new set of analytical tools at hand, we investigated the dynamics of CME in amnioserosa tissue of developing Drosophila embryos. Amnioserosa consists of a single layer of polarized cells that covers the dorsal surface of the embryo after germ band retraction. The tissue is entirely curtain by lateral epidermal cells in ~4 h by a process called dorsal closure (Jacinto et al., 2002). CCSs originating at the apical and basal surfaces of the amnioserosa can be detected in embryos expressing clathrin light chain (CLC) fused with GFP, using spinning disk confocal fluorescence imaging (Fig. 8 A). CCSs forming on intracellular organelles appear as bright blobs at the perinuclear regions. To determine individual CCS traces at the amnioserosa, we used a robust 2D particle tracking software (Aguet et al., 2013) and processed the output data using a trace combination algorithm to correct for particle disappearances caused by movements in the axial dimension (Fig. 8 B and Video 3). We used local distributions of the number of CCS traces to determine the axial positions of apical and basal surfaces (Fig. 8 C). We found that lifetime distributions for both of the surfaces are dominated by short CCS traces in comparison with the distributions obtained for clathrin-coated pits originating in cultured cells (Fig. 8 D). We believe the anomaly in lifetime distributions obtained from the amnioserosa has no biological basis but is a consequence of single-particle tracking errors caused by high CCS motility in this tissue and increased ratio of incomplete trace fragments (Fig. S3). When we used CCS growth rates as the alternative approach, we noted rapid changes in the distributions obtained from individual amnioserosa cells. However, we found no correlation between the temporal evolution of the distributions at the apical and basal surfaces (Fig. 9, A and B). We extended the duration of the imaging assays by acquiring CCS growth rates from 30-s-long acquisitions that are separated by temporal gaps of 3 or 10 min (Fig. 9 C). In this experimental scheme, monitoring CCS dynamics of individual cells throughout the entire assay was not achievable because the cellular organization of amnioserosa is not preserved over long durations, as the tissue is gradually replaced by lateral epidermis. When the growth rates are calculated for the entire tissue, we found that the intensity traces predicted using CCS growth rates were significantly longer lived than typical clathrin-coated pits (Fig. 9 E), in striking contrast with the results of the lifetime distributions (Fig. 8 D). In all of the four embryos we have analyzed in this way, we recorded significant change in growth rate distributions at the apical surface, which correspond to slower CCS dynamics and elongated lifetimes over time (Fig. 9, D–F).
We show that clathrin coat growth rates can be used as quantitative reporters of CME dynamics in cellular contexts where errors associated with single-particle tracking are significant. Growth rate distributions obtained from short fragments of CCS traces offer the ability to assess clathrin dynamics from acquisitions shorter than the mean CCS lifetime. Gradual changes in endocytic dynamics can be detected in real time. Mean intensity profiles of the CCS traces that are used to generate growth rate distributions can be reconstructed by finding analogous growth rates in a library of trace clusters. These advantages make growth rate analysis a strong alternative to existing methodologies, which rely on quantifying lifetimes of individual CCSs.

Growth rate analyses reveal that physical factors increasing the energy cost of membrane deformation (e.g., in-plane tension, membrane–cytoskeleton adhesion, and membrane–substrate interactions) slow down formation and dissolution of CCSs. It has been shown that the same energy cost also reduces the curvature of CCSs (Saleem et al., 2015). In-plane tension is assumed to be homogenous in a cell, but membrane–cytoskeleton adhesion and membrane–substrate interactions are not uniform and, therefore, may induce spatial heterogeneities in dynamics and geometries of CCSs. This heterogeneity
may account for the coexistence of clathrin-coated pits and plaques at the ventral surface of certain cell types (Saffarian et al., 2009). Growth rate analyses suggest that flat clathrin arrays and clathrin-coated pits have distinct formation dynamics. Even though flat lattices may acquire curvature over time (Avinoam et al., 2009), our data suggest that this mechanism cannot account for the formation of clathrin-coated pits. Transition of flat clathrin lattices to curved pits requires fourfold reduction in the projected area of clathrin coats on the x–y plane (Fig. S4). Such abrupt shape transformations are not observed in recent super-resolved fluorescence acquisitions displaying the formation of clathrin-coated pits (Li et al., 2015), which suggests that curvature of the clathrin coat is constant during pit formation.

Strategies described here lay the groundwork for other quantitative assays that aim to elucidate endocytic processes in various multicellular contexts. Experimental and analytical deficiencies that distort CCS lifetime distributions are less detrimental to growth rates, as their quantification does not rely on tracing the complete lifetime of CCSs. Our study shows that CME dynamics can be assessed within tissues of developing organisms through quantification of the fluorescence growth rates of individual endocytic assemblies. Lifetime distributions obtained from Drosophila amnioserosa are overpopulated by traces that last 0.2 s, which would be regarded as abortive structures in in vitro assays. The intensity traces reproduced using growth rate distributions extended to durations longer than the mean clathrin-coated pit lifetime. Longer CCS lifetimes may be a result of the intrinsic tension of the amnioserosa tissue (Kiehart et al., 2000). Likewise, gradual increase in CCS lifetimes during dorsal closure may be induced by the changes in tension levels. This assumption is in accord with a recent study showing that reduction in volume of amnioserosa cells leads to increasing tension during dorsal closure (Saias et al., 2015). Drosophila amnioserosa cells undergo periodic but asynchronous shape changes at shorter time scales (Solon et al., 2009). Further research is needed to elucidate if rapid fluctuations of CCS growth rates observed in individual cells are correlated with temporal changes in amnioserosa mechanics. Such studies may also elucidate whether dynamics of CCSs can be used as reporters for mechanical states of cells and tissues.

### Materials and methods

#### Fluorescence imaging

The imaging system was an Eclipse TI-E microscope (Nikon) equipped with a temperature controlled chamber, a CSU-W1 spinning disk confocal unit (Yokogawa Electric Corporation), a 100× objective lens (Nikon CFI Plan-Apochromat Lambda, NA 1.45) and an EMCCD camera (iXon DU897 Ultra; Andor Technology). 2D and 3D time series were obtained using NIS Elements image acquisition software.

S. Boullant (University of Heidelberg, Heidelberg, Germany) provided BSC-1, MDCK, and U373 cells stably expressing σ2-EGFP. Imaging of cultured cells was performed 8–24 h after plating on glass bottom dishes at 37°C ambient temperature (Greiner Bio-One), unless stated otherwise. Cells were imaged at a rate of 0.25–0.5 Hz and laser exposure of 50–300 ms per frame. Imaging medium was phenol red–free L15 (Thermo Fisher Scientific) supplemented with 10% FBS. Serum-free L15 was used for cholesterol depletion experiments. Final concentration of MβCD (Sigma-Aldrich) was 10 mM in Fig. 5, Fig. 7 C, and Video 2 and 4 mM in Fig. S2 B. Latrunculin B (Sigma-Aldrich) was used at the final concentration of 2 μM.
Drosophila embryos were harvested from a cross between male flies homozygous for GAL4-Arm and female flies homozygous for CLC-GFP (Bloomington Drosophila Stock Center). Eggs laid by this cross were collected and allowed to develop for ∼11 h at 25°C (stage 14 of development). The embryos were then mounted on a slide with glue (ventral side up), immersed in halocarbon oil (to prevent drying out), and covered with a coverslip. Amnioserosa tissues were imaged at 22°C using 3D time-lapse acquisitions containing ∼20 planes (400 nm apart) exposed for 50–100 ms. Temporal gap between adjacent z-stacks was 3 s.

Micropipette aspiration
Glass micropipettes (BF100-58-10) were pulled using a micropipette puller (P-97; Sutter Instrument). A custom stage was built to mount a Sutter Instrument BRM/E micromanipulator to the live-cell imaging platform. Suction pressure was controlled with a Sutter Instrument BRE110/E microinjection system. Cell aspiration was performed with a 5- to 10-µm microneedle at the dorsal surface of cells, and the clathrin activity at the ventral surface was recorded with spinning-disk fluorescence imaging. P. Selvin (University of Illinois at Urbana-Champaign, Urbana, IL) provided the micromanipulator and injection system.

Single-particle tracking
2D particle tracking was performed using the cmeAnalysis software (obtained from http://lccb.hms.harvard.edu/software.html) unless stated otherwise. TraCker software is a less sophisticated but faster tracking algorithm that lacks the advanced forward-backward-forward rechecking and intelligent thresholding of cmeAnalysis. It also lacks the 2D Gaussian fitting of point-spread functions and uses integrated pixel intensity as the CCS signal. TraCker uses a simple threshold determined over a Mexican hat filtered image for detection of fluorescent spots. CCS positions are located using the intensity-
weighted center of the point-spread function. In this work, TraCKer was used for the sole purpose of testing growth rate analysis on IMSs. cmeAnalysis software occasionally detects objects that last a single frame or persist consistently in the background without following an intensity path that could be considered as a CCS. We implemented a sorting scheme for these traces, requiring that they be at least three frames long and at some point meet statistical criteria for demonstrating a linear increase or decrease in intensity (corresponding to CCS growth and dissolution, respectively). We went over every three or four consecutive intensity points (three for traces that last for <10 frames and four for longer traces) and performed a least-squares fit. Traces that had no fits with \( r^2 \) value > 0.75 were rejected. Rejected traces were not used in calculation of growth rate distributions. Traces with at least two separate high \( r^2 \) values of both positive slope (growth) and negative slope (dissolution) were valued most highly. Fig. S5 shows the classification of CCS traces obtained from amnioserosa tissues.

3D tracking of CCSs within amnioserosa tissues

Each \( z \)-plane of a 3D time-lapse movie was analyzed using cmeAnalysis software. Detected 2D traces were then run through our trace-rejection scheme. The resulting data were analyzed to combine traces that occur at the same lateral position in two adjacent \( z \)-planes. Coincident traces had to be within one pixel (160 nm) x–y distance for at least three consecutive frames. For each time point in the movie, the resulting trace contains the maximum intensity value of all traces considered for combination. Axial positions were assigned by calculating the intensity-weighted mean \( z \)-position of all traces considered. The algorithm for trace combination ran from the outermost \( z \)-planes to the innermost (alternating between the top and the bottom) to ensure that there was no directional bias and all possible trace combinations were considered. The resulting data structure will have the duplicate traces deleted and contain subplane \( z \)-position data where possible.

Classification of apical and basal CCSs and blobs

Because of the curvature of the dorsal surface, cells in the amnioserosa tissue are not coplanar. Therefore, the field of view was divided into 64 equal square regions, and apical and basal surfaces were determined for each region independently (Fig. 8 B). Axial positions of the traces detected inside each square region were put together within in a two-frame temporal radius. These values were binned into discrete \( z \)-plane positions, and the resulting count-value graph was fit with two Gaussian functions. The fit for the apical surface has a low standard deviation. The basal surface was found at a higher axial position and the corresponding Gaussian fit had a larger standard deviation in general. Traces found at the mean or within one standard deviation of their respective Gaussian fits were classified as apical or basal for each of the 64 squares. The brightest 2% of all traces were classified as blobs and were excluded from the apical and basal populations.

Determining amnioserosa cell boundaries

To determine the positions of amnioserosa nuclei, the middle imaging plane of each \( z \)-stack was filtered with a Gaussian filter. Each frame was then inverted and local maxima were determined using the FastPeakFind function (A. Natan, PULSE Institute, Stanford, CA). After multiple nuclei positions were determined for each frame, they were tracked using the TraCKer algorithm. Once the centers were known and connected for each frame, the boundaries were determined by creating a Voronoi diagram using the cell centers in each frame.

Growth rate distributions

Slopes were extracted from CCS traces that pass the trace rejection scheme. We found that CCS lifetime distributions of obtained from accepted (nonrejected) traces were very similar to those reproduced by others who used different parameters to distinguish genuine endocytic transporters (Fig. S5; Aguet et al., 2013). Each trace was normalized by subtracting a global background and dividing by the new maximum of the trace. From this normalized trace, every 12-s interval was used in a least-squares fit to determine the slope of the trace at all frames (Fig. 1 C). A trace had to be at least 12 s long to be considered for slope extraction. 12 s was chosen because it evenly divides 1-, 2-, 3-, and 4-s frame rates, which are the most common frame rates used. An arbitrary bin width (0.03) was assigned for the histograms displaying the five distinct growth phases.

Hierarchical clustering of intensity profiles and library lookup

Representative traces were created by combining similar intensity traces within a single cell using a standard hierarchical clustering algorithm. The metric used was the mean squared Euclidean distance, with the condition that the lifetimes of the traces were within 10 s of each other. To compare clusters, we used complete linkage clustering; that is, we used the distance between the farthest two component traces. After clustering, clusters with a minimum of eight traces were added to our trace library. For such clusters, the mean of the component traces was calculated. These mean traces were normalized to the mean maximum trace intensity of the cell to compensate for cells’ differing expression levels and imaging conditions. Additionally, for each of these clusters, a single growth rate histogram was generated from the combined data of the constituent traces. The library was assembled from clusters obtained from the following cell types and conditions: BSC-1 cells expressing \( \sigma_2 \)-GFP (before and during aspiration), polarized MDCK cells expressing \( \sigma_2 \)-GFP (apical and basal surfaces), and U373 cells expressing LCa-mCherry.

Growth rate histograms of test clusters were compared with the histograms of library clusters using the squared Euclidean metric. If the input histogram was generated from trace data that were confined spatially (e.g., single cell in a culture or tissue) and/or temporally (e.g., half a minute long acquisition of the whole tissue), predicted traces are the weighted averages of the 10 closest cluster matches from the CCS library (Fig. 7 C, Video 2, and Fig. 9 E). For this procedure, the library clusters were sorted from closest to farthest by calculating the squared Euclidean distance between their growth slope histograms, and each cluster’s contribution to the mean is weighted by the factor \((10 - i)\), where \( i \) is the cluster’s index in the sorted list and the first index is \( i = 0 \).

Online supplemental material

Fig. S1 provides examples of hierarchical clustering in cells before and during micropipette aspiration. Fig. S2 describes the difference between traces found in control cells and in those treated with MβCD and how latrunculin B affects that response. Fig. S3 compares the root mean squared displacement of traces detected in the amnioserosa tissue versus those found in U373 cells. Fig. S4 demonstrates the differences between a flat clathrin array and a curved clathrin pit. Fig. S5 shows the intensity and lifetime distributions of CCS traces detected in amnioserosa cells based on their classification. Video 1 gives a frame-by-frame depiction of CCS growth rates throughout a cell and its corresponding growth rate histogram. Video 2 gives a frame-by-frame depiction of growth rate histograms in three separate cells treated simultaneously with MβCD and displays corresponding intensity traces derived from our hierarchical clustering library. Video 3 shows a 3D maximum projection image of amnioserosa tissue with an overlay marking CCSs classified as blobs. It also shows traces colored to demonstrate the axial position. Video 4 shows the boundaries of individual cells in the amnioserosa over the course of the movie. Finally, the MATLAB-based software used for our analysis is provided to be used or modified as fits the user’s needs. The software package includes a README file, clearly commented code, and example data files. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201604128/DC1.
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