

Robust and automated detection of subcellular morphological motifs in 3D microscopy images

Meghan K. Driscoll^{1,2}, Erik S. Welf^{1,2}, Andrew R. Jamieson^{1,2}, Kevin M. Dean^{1,2}, Tadamoto Isogai^{1,2}, Reto Fiolka^{1,2} and Gaudenz Danuser^{1,2}*

Rapid developments in live-cell three-dimensional (3D) microscopy enable imaging of cell morphology and signaling with unprecedented detail. However, tools to systematically measure and visualize the intricate relationships between intracellular signaling, cytoskeletal organization and downstream cell morphological outputs do not exist. Here, we introduce u-shape3D, a computer graphics and machine-learning pipeline to probe molecular mechanisms underlying 3D cell morphogenesis and to test the intriguing possibility that morphogenesis itself affects intracellular signaling. We demonstrate a generic morphological motif detector that automatically finds lamellipodia, filopodia, blebs and other motifs. Combining motif detection with molecular localization, we measure the differential association of PIP₂ and Kras^{V12} with blebs. Both signals associate with bleb edges, as expected for membrane-localized proteins, but only PIP₂ is enhanced on blebs. This indicates that subcellular signaling processes are differentially modulated by local morphological motifs. Overall, our computational workflow enables the objective, 3D analysis of the coupling of cell shape and signaling.

ell morphogenesis is driven by cytoskeleton-generated forces that are regulated by biochemical signals¹. The cascade from signaling to cytoskeleton to shape control is well established for numerous morphological motifs, including lamellipodia, blebs and filopodia (Fig. 1a–c, Supplementary Video 1 and Supplementary Fig. 1), which depend on well-characterized assemblies of actin filaments (Fig. 1d–f)². How morphology, in turn, may govern signaling is less investigated. Morphology may participate in signal transduction via mechanisms such as preferential protein interaction with membranes of particular curvature³, or modulation of the concentration and diffusion of signaling components⁴,⁵.

The integrated study of signaling and morphology at subcellular length scales has become possible with the recent advent of high-resolution 3D light-sheet microscopy^{6–11}. Using microenvironmental selective plane illumination microscopy (meSPIM)¹⁰ of PIP₂, a membrane-bound phosphoinositide implicated in diverse signaling pathways¹², we found an unexpected formation of PIP₂ clusters in both branched (Fig. 1g,h) and blebbed cells (Fig. 1i,j). 3D renderings of the local concentration of PIP₂ suggest that these clusters tend to colocalize with filopodial tufts (Fig. 1h) and blebs (Fig. 1j). Kras^{V12}, which is a constitutively active GTPase with broad oncogenic functionality¹³, also appears to colocalize with certain morphological structures (Fig. 1k,l and Supplementary Videos 2 and 3). These observations pose the question of whether rugged surface geometries generally associate with elevated signaling, and whether there are differences in how PIP₂ and Kras associate with cell morphologies.

Answering such questions with statistical robustness requires the interpretation of 3D images. Not only is the inspection and quantification of such images exceedingly laborious, the difficulty of representing 3D images in meaningful two-dimensional (2D) perspectives renders the manual annotation of subcellular geometries extremely difficult. Automation by computer vision is essential. However, the tools for subcellular 3D morphometry do not exist¹⁴. Here, we introduce u-shape3D, a pipeline that combines computer graphics and machine-learning approaches to unravel the coupling

between cell surface morphology and subcellular signaling. At its core is the segmentation of any morphological motif a user can provide systematic examples for. We show the robustness of a oncelearned motif classifier to changes in microscopy and cell type. We then apply the method to analyze the differential association of $\rm PIP_2$ and Kras $^{\rm V12}$ with surface blebs. Moving forward, u-shape 3D will be instrumental to furthering our understanding of the feedback interactions between signaling, the cytoskeleton and morphological dynamics in 3D.

Results

Detecting cellular morphological motifs. In designing u-shape3D, we decided to first represent the cell surface as a triangle mesh, and then segment the surface into motifs using machine learning (Fig. 2a-e). An alternative approach would be to segment the motifs directly from the raw image data on a voxel-by-voxel basis, and then generate a surface representation with classified motifs. This would simplify the application of deep learning algorithms, but would require the acquisition of training data in the raw image volume, where manual outlining of interesting motifs can become exceedingly cumbersome. In contrast, the proposed machine-learning pipeline depends on training data that is defined on a surface representation with pre-segmented patches, where a few easy-to-identify examples of the motif of interest are sufficient to constrain a robust classifier. This provides a versatile and efficient approach to quantifying diverse cellular morphologies.

To generate the cell surface, for most cells we automatically extract the mesh as an isosurface of the deconvolved image (Fig. 2f-h and Supplementary Fig. 2). However, some datasets require that surface extraction parameters be tailored to the cell type and fluorescence label (see the user's guide to the software package). For example, we extract the surfaces of actin-labeled dendritic cells as an isosurface of an image that combines the deconvolved image, an image with enhanced planar features, and an image with an enhanced cell interior (Supplementary Fig. 3).

Lyda Hill Department of Bioinformatics, University of Texas Southwestern Medical Center, Dallas, TX, USA. ²Department of Cell Biology, University of Texas Southwestern Medical Center, Dallas, TX, USA. *e-mail: Gaudenz.Danuser@UTSouthwestern.edu

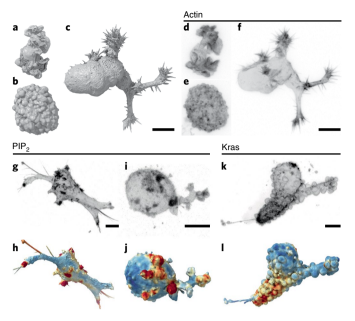


Fig. 1 | Cell morphology and signaling are coupled. a-c, Surface renderings of a dendritic cell expressing Lifeact-GFP (a), an MV3 melanoma cell expressing tractin-GFP (b) and a HBEC expressing tractin-GFP (c). **d-f**, Maximum intensity projections (MIPs) of the cells shown in **a-c**, using an inverse look-up table. a-f are shown at the same scale. Additional views of these cells are shown in Supplementary Fig. 1. g, A MIP of a branched MV3 cells expressing PLCΔ-PH-GFP, a PIP₂ translocation biosensor. **h**, A surface rendering of the same cell. Surface regions with relatively high PIP₂ localization are shown in red, whereas regions of relatively low localization are shown in blue. i,i, A MIP (i) and a surface rendering of a blebbing MV3 cell expressing PLC Δ -PH-GFP (\mathbf{j}). The PLC Δ -PH-GFP images are representative of 23 cells from three experiments. k, A MIP of an MV3 cell expressing GFP-Kras^{V12}. I, A surface rendering of k. Surface regions of relatively high Kras localization are shown in red, whereas regions of relatively low localization are shown in blue. The GFP-Kras^{V12} images are representative of 31 cells from seven experiments. Scale bars, 10 µm.

After cell surface extraction, we decompose the surface into convex patches. People tend to partition 3D surfaces into convex regions 15 , suggesting that canonical protrusions are likely convex or composed of multiple convex regions. Convex decomposition is in general an NP-complete problem 16 , and thus is computationally intractable for large meshes, even with extensive computing resources. We therefore combine several techniques to segment the surface into approximately convex patches. First, we calculate the mean curvature 17 at every face on the mesh, and then break the surface into small patches via a watershed-based segmentation of mean curvature (Fig. 2i-k) 18 . These small patches are computationally manipulated more easily than individual faces and are analogous to superpixels in image segmentation.

Next, we iteratively merge patches using two criteria (Fig. 2l,m and Supplementary Fig. 4). The line-of-sight (LOS)¹⁹ criterion merges patches if the percentage of rays that connect the two patches without exiting the cell is above a certain threshold. Hence, fulfilling this criterion requires only approximate convexity between the patches. The triangle criterion¹⁰ merges adjacent patches whose joint closure surface area, defined as the additional surface area needed to close the mesh composing the patch, is small compared to the sum of their individual closure surface areas. This criterion embodies the short-cut rule²⁰ that the preferred convex shape decomposition has the shortest cuts between segments.

Approximate convex patches are then classified by morphological motif type using a support vector machine (SVM). For each

patch, 23 geometric features are calculated (Supplementary Table 1). Features are automatically selected for each set of training data by successively removing randomly chosen features until prediction quality is hampered. Following SVM training (Fig. 2n), the trained motif model is used to classify each patch by motif type (Fig. 20,p).

The outcomes of machine-learning approaches are critically dependent on training data quality. To generate training data, we built an interface where users can rotate 3D surfaces, zoom in and out and click on patches to identify them as motifs. Presented with the same four randomly chosen cells, three users chose $46\pm6\%$ of the patches when asked to click on blebs and $25\pm4\%$ of the patches when asked to click not on blebs. This discrepancy carried over into SVM models, where for the two training sets 45 ± 7 and $77\pm6\%$ of the patches were identified as blebs. Asking users to click only on patches that are certainly blebs and then only on patches that are certainly not blebs resulted in models that classified an intermediate percentage of patches, $52\pm6\%$, as blebs. To avoid bias, we therefore train models with data where users only choose patches they can confidently classify.

Although many morphological motifs, including blebs and filopodia, are described by a single convex surface patch, some motifs, such as lamellipodia, are composites of multiple convex patches. To detect these motifs, we merge convex patches before patch classification using a machine-learning framework (Fig. 2q-u). Thirty-six geometric features are calculated for each pair of patches (Supplementary Table 2), and training data is generated by asking users to click on adjacent patches that should certainly or certainly not be merged. Following sequential feature selection, an SVM is used to merge patches.

We trained models to detect blebs, filopodia and lamellipodia (Fig. 3, Supplementary Video 4 and Supplementary Fig. 5). Most cells in our diverse dataset showed predominately one protrusion type. However, as a proof-of-concept, we also built a multiclass detector using a collection of melanoma cells that exhibited extensive blebs and small numbers of filopodia (Fig. 4a). To do so, we generated multiple SVM models in a one-versus-one framework in which separate models were used to distinguish each pair of morphological motifs.

Validation and robustness of motif detection. To validate the protrusion classification, we calculated the F_1 score using patches selected by the trainer as certainly or certainly not a protrusion. For four randomly chosen blebby melanoma cells, the F_1 score calculated via leave-one-out-cross-validation over cells and averaged across three trainers was 0.986 ± 0.006 , corresponding to $1.3 \pm 0.6\%$ incorrectly classified patches. This score is high, in part, because only patches users were certain about were included. Calculating F_1 scores for the models where users clicked on all the blebs or all the non-blebs yielded 0.77 ± 0.03 and 0.76 ± 0.04 , respectively. However, as discussed above, these training data are biased toward selecting too few and too many blebs, respectively. Indeed, using these training data to validate our model, we find a $16 \pm 1\%$ false positive rate $(5\pm1\%$ false negative rate) when users are asked to click on all the blebs and a $30 \pm 6\%$ false negative rate ($2 \pm 1\%$ false positive rate) when users are asked to click on all the non-blebs. Validating over a larger number of cells with a single user, we measured an F_1 score of 0.99 for 19 MV3 cells with blebs, 0.94 for 13 human bronchial epithelial cells (HBECs) with filopodia and 0.88 for 13 dendritic cells with lamellipodia (Fig. 4b and Supplementary Fig. 6a,b).

We also tested other machine-learning algorithms. We anticipated that the classifier performance would be primarily feature driven, rather than algorithm driven. Indeed, using random forests, linear SVMs and radial SVMs to detect filopodia and varying the number of rounds of feature selection, we calculated F_1 scores of between 0.934 and 0.944 for almost all algorithms (Supplementary

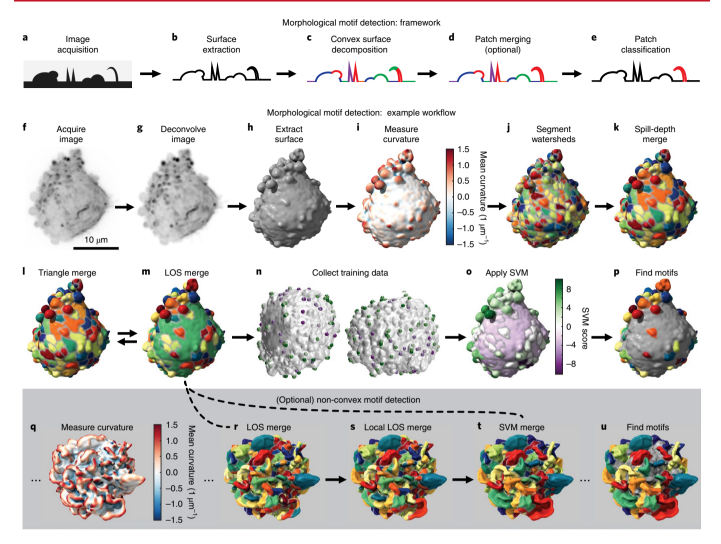


Fig. 2 | Morphological motif detection framework and example workflow. a-e, To detect morphological motifs, following image acquisition (a), we extract the cell surface (b), decompose that surface into convex patches (c), optionally merge those patches (d) and finally classify the patches by morphological motif (e). f-p, Our detection framework applied to a blebbed cell. f, MIP of a 3D image of an MV3 melanoma cell expressing tractin-GFP. g, MIP of the deconvolved image of the same cell. h, The surface of the cell extracted from the deconvolved image as a triangle mesh. i, The mean surface curvature of the cell. Regions of large positive curvature are shown in red, flat regions are shown in white and regions of large negative curvature are shown in blue. j, A watershed segmentation of mean surface curvature. Segmented patches are shown in different colors. k, A spill depth-based merging of the segmented patches. I, A triangle-rule based merging of the patches. m, A LOS-based merging of the patches. The triangle and LOS rules are applied iteratively. n, User generated training data for two different cells. Patches identified as 'certainly a bleb' are shown in green, whereas patches identified as 'certainly not a bleb' are shown in purple. o, A SVM classifier trained on user data applied to the cell. Patches shown in green have high SVM scores and a high inferred likelihood of being a bleb, whereas patches shown in purple have low SVM scores and a low inferred likelihood. p, Detected blebs are shown as randomly colored and non-blebs are shown in gray. To detect non-convex motifs, such as lamellipodia, convex patches are merged as shown in q-u. q, The mean surface curvature of a lamellipodial dendritic cell expressing Lifeact-GFP. r, Convex surface patches for the same cell. s, A local LOS-based merging of these patches. t, An SVM-based merging of the patches. The SVM was trained on user-supplied examples of adjacent patches that should certainly be merged and adjacent patches that should certainly not be merged.

Table 4). This suggests that in our workflow linear SVMs perform as well as a broad class of machine-learning algorithms.

Conversely, a carefully chosen feature set might be able to distinguish motifs from non-motifs using even an unsupervised algorithm that does not require training. We hierarchically clustered all convex surface patches on a set of seven blebby cells into two clusters using such an algorithm (Supplementary Fig. 7). Although one of the clusters substantially overlaps with the bleb detection, the supervised algorithm clearly performs better.

Because of the selective power of the geometric feature set, our workflow requires relatively little training data. One user training on just one cell in a dataset of 19 blebby cells, yields an F_1 score

of 0.94 ± 0.04 (mean \pm s.d.) on the remaining cells (Fig. 4c and Supplementary Fig. 6c). Additional training data improves the model accuracy marginally, suggesting that models generated by a single user on different datasets would be similar. Indeed, models trained by a single user on distinct sets of four MV3 cells show $95.9\pm0.7\%$ overlap, as measured by the Sorrenson–Dice index²¹. This compares to an $88\pm3\%$ overlap between models generated by different users (Supplementary Fig. 6d). To maximize reproducibility, our classifiers therefore incorporate training data from multiple users via majority voting.

Motif models from one cell type can be extended to dissimilar cell types, enabling objective comparisons across biological systems.

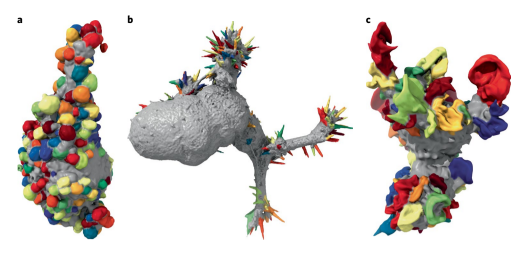


Fig. 3 | Detected blebs, filopodia and lamellipodia. a, Blebs detected on an MV3 melanoma cell (representative of 19 cells). **b**, Filopodia detected on an HBEC cell (representative of 13 cells). **c**, Lamellipodia detected on a dendritic cell (representative of 13 cells). Additional example detections are shown in Supplementary Fig. 5.

Applying a bleb model generated from 19 cells originating from a melanoma cell line to 24 cells originating from a human melanoma xenografted into mice yields an F_1 score of 0.97 (Fig. 4d). Likewise, applying a filopodia model generated from nine melanoma cells to 13 transformed HBEC cells yields an F_1 score of 0.90 (Fig. 4e).

The classifier can also be used to compare perturbed and non-perturbed cell populations. To test whether greatly different cell morphologies confound the detection of particular motifs, we used identical analysis parameters to measure the fraction of the cell surface that is blebby in wild-type U2OS cells and cells where the actin regulatory proteins cofilin-1 and Wave2 were knocked out with clustered regularly interspaced short palindromic repeats (CRISPR) (Fig. 4f,g). Compared to wild-type cells, *CFL1* knockout (cofilinKO) cells exhibit greater cell-to-cell heterogeneity in their bleb surface fraction as well as a larger mean bleb fraction. *WASF2* knockout (WaveKO) cells exhibit even greater heterogeneity and a yet larger mean fraction.

So far, we have presented data acquired via meSPIM, a high-resolution light-sheet microscope with nearly isotropic resolution 10 . On microscopes with anisotropic resolution, the motif structure varies with orientation relative to the microscope. To test if resolution anisotropy impedes motif detection, we analyzed blebby cells imaged via a laser scanning confocal microscope (Fig. 5a,b). Blebs appeared stretched in the axial (z) direction (Supplementary Fig. 8), however, the workflow still achieved an F_1 score of 0.94 (Supplementary Fig. 9). Standard light-sheet microscopy also has reduced axial resolution compared to meSPIM. Analyzing microglial cells imaged in vivo within a zebrafish via a lower resolution commercial light-sheet microscope, we successfully detected extensions (Fig. 5c,d). These findings demonstrate that our pipeline can analyze data from conventional microscopes with anisotropic resolution.

To determine whether motif models were transferable among similar microscopes, we directly applied meSPIM motif models to cells imaged by other high-resolution light-sheet microscopes. Detecting blebs on cytosolically labeled cells imaged by axially swept light-sheet microscopy (ASLM)⁹, we measured an F_1 score of 0.96 for both meSPIM and ASLM derived models (Fig. 5e,f). Analyzing previously published movies, we used a meSPIM derived model to detect lamellipodia on a T cell imaged by lattice light-sheet microscopy⁸ (Fig. 5g,h), and trained a new model to detect extensions on a human breast cancer cell moving through the vasculature of a zebrafish embryo imaged by adaptive-optics lattice light-sheet microscopy²² (Fig. 5i and Supplementary Video 5). Together, these

test cases show the broad applicability of u-shape3D, allowing objective comparisons between large numbers of diverse datasets.

Kras and PIP, signals associate differently with blebs. Equipped with a computational framework to analyze 3D cell morphology, we set out to identify relationships between morphological motifs and signaling events. We focused on blebs as the predominant morphological feature of melanoma cells in soft 3D environments¹⁰ and sought to measure how PIP, and constitutively active Kras^{V12}, may associate with this motif. Both Kras^{V12} (Fig. 11) and PIP₂ (Fig. 1j) appear to polarize and associate with blebs. To test these hypotheses, we measured the localization of Kras^{V12} within 2 µm of the cell surface for 13 MV3 melanoma cells (Fig. 6a,b). We computed at every mesh face the average fluorescence intensity in a sphere around that face, including only pixels within the cell and correcting for surface curvature-dependent artifacts by depth-normalization²³. In addition to blebs, cells expressing green fluorescent protein (GFP)-Kras^{V12} exhibited retraction fibers and uropods, which could lead to bias. To exclude these structures from the analysis, we built a retraction fiber/uropod detector and subtracted those patches from the set of detected blebs (Supplementary Fig. 10). Next, using spherical statistics we found that the Kras^{V12} distribution on the cell surface was polarized (Fig. 6c). Likewise, blebs and Kras^{V12} surface intensity were directionally correlated (Fig. 6d). However, randomizing the location of blebs on the surface, the directional correlation of Kras^{V12} with blebs was not significantly different from random. This suggests that Kras^{V12} and bleb polarization are correlated partially through their joint coupling to global cell shape. Measuring the mean Kras^{V12} localization on and off detected blebs, we found no statistically significant difference (Fig. 6e). However, Kras^{V12} does localize to bleb edges (Fig. 6f). In contrast, cytosolic GFP showed no localization to bleb edges (Fig. 6f). This confirms that the modulation of Kras^{V12} across the cell surface is related to the distribution of blebs. To further examine the mechanism of this association we measured bleb density locally over a scale less than that of a single bleb by simulating the diffusion on the mesh of the motif classification label (Supplementary Fig. 11a,b). In this representation, high-density values localize to the bleb center, low-density values to areas away from any bleb and intermediate values colocalize with bleb edges. In agreement with our previous conclusion, high Kras^{V12} signal associated with intermediate local bleb densities (Supplementary Fig. 11c). Thus, these analyses suggest that Kras^{V12} may be organized to bleb edges. This may at first seem surprising: Kras^{V12} is a constitutively activated GTPase

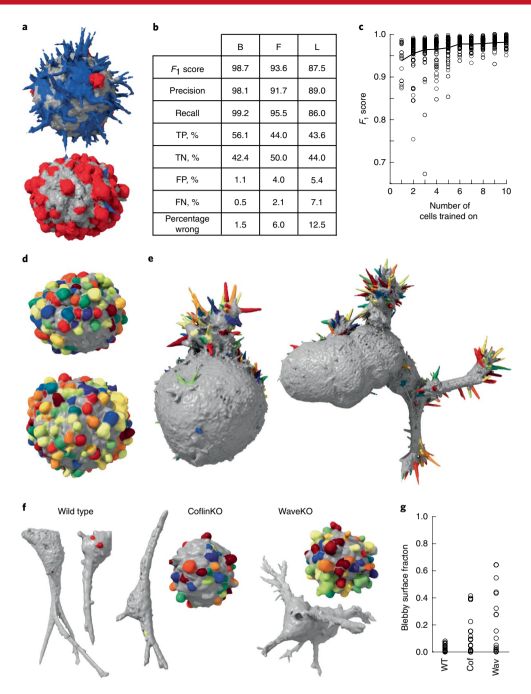


Fig. 4 | Validation and robustness of morphological motif detection. a, A multiclass detector applied to cells derived from a human melanoma xenograft cultured in mice (representative of nine cells). Filopodia are shown in blue, blebs are shown in red and areas with neither filopodia nor blebs are shown in gray. **b**, Validation measures for a bleb detector (B) trained on 19 MV3 melanoma cells, a filopodia detector (F) trained on 13 HBEC cells, and a lamellipodia detector (L) trained on 13 dendritic cells. TP, true positive; TN, true negative; FP, false positive; FN, false negative. **c**, The F_1 score as a function of the number of cells trained on. The black line indicates the mean F_1 score averaged over a maximum of 100 sets of cells, whereas the black circles show individual sets of cells. **d**, A bleb detector trained on the MV3 melanoma cell line applied to cells derived from a human melanoma xenograft cultured in mice (representative of 24 cells). **e**, A filopodia detector trained on xenograft-derived melanoma cells applied to HBEC cells (representative of 13 cells). A filopodia detector trained on HBECs applied to the cell on the left is shown in Supplementary Fig. 5 and applied to the cell on the right is shown in Fig. 3. **f**, Blebs detected on wild type, cofilin-1 knock out and Wave2 knockout U2OS cells in 3D collagen. **g**, For these three cell types, the percentage of the surface that is blebby. We analyzed 19 wild type (WT), 15 cofilin-1 knockout (Cof) and 14 Wave2 knockout (WAV) cells.

without spatially organized interactions with guanine nucleotide exchange factors, GTPase activating proteins and GDP dissociation inhibitors. Accordingly, the Kras^{V12} distribution is expected to be dominated by diffusion within the plasma membrane with an overall uniform steady state. Simulations of uniformly labeled surface distributions in synthetic cells demonstrate an intensity

co-modulation with bleb edges for a variety of surface thicknesses, but not for cytosolically labeled cells (Supplementary Fig. 11e). This shows that the Kras^{V12} localization at bleb edges is consistent with a uniform surface distribution. This discovery also shows how, in 3D, cell morphological motifs alone offer a mechanism for the spatial organization of molecular signals at the subcellular scale.

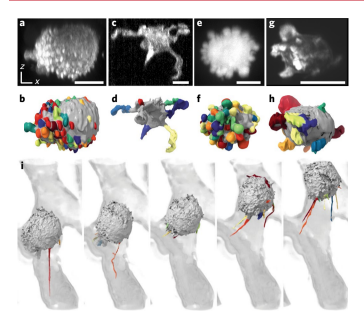


Fig. 5 | Motif detection on images acquired via diverse microscopic techniques. a, A MIP, taken over the xz direction, of an MV3 cell expressing tractin-GFP imaged via laser scanning confocal microscopy (representative of eight cells). **b**, Blebs detected on the same cell using a model derived from eight MV3 cells imaged with this microscope. c, An xz-MIP of a microglia inside a zebrafish embryo imaged using a commercial light-sheet microscope (representative of eight cells). d, Extensions detected on the same cell using a model derived from eight microglia imaged with this microscope. **e**, An xz-MIP of an MV3 cell expressing cytosolic GFP imaged using ASLM, a high-resolution light-sheet microscopy modality (representative of eight cells). f, Blebs detected on the same cell using a model derived from 19 MV3 cells imaged via meSPIM. g, An xz-MIP of a T cell expressing Lifeact-mEmerald imaged using lattice light-sheet microscopy⁸. **h**, Lamellipodia detected on the same cell using a model derived from 13 dendritic cells imaged via meSPIM. i, Extensions detected on an MDA-MB-231 human breast cancer cell moving through the vasculature of a zebrafish embryo imaged via adaptive-optics lattice light-sheet microscopy²². The cell is shown as a surface rendering, whereas the vasculature is shown in gray as a MIP of the deconvolved image. Scale bars, 10 µm.

We next analyzed MV3 cells expressing PLCΔ-PH-GFP (Fig. 6h,i), a PIP, translocation biosensor that reports the activation of PIP₂. Like Kras^{V12}, the surface localization of PIP₂ is polarized (Fig. 6j), and blebs and PIP, are directionally correlated (Fig. 6k). However, unlike Kras^{V12}, the directional correlation of PIP₂ with blebs was significantly different from that of PIP, with randomized bleb distributions. Hence, PIP, polarization is directly correlated with bleb polarization rather than coupled via the overall cell shape. Indeed, PIP₂ localizes to blebs, with each cell exhibiting a higher mean PIP2-intensity on blebs than off (Fig. 6l). Consistent with a surface fluorescence distribution, PIP₂ like Kras^{V12} also associates with bleb edges (Fig. 6m). However, whereas Kras^{V12} localization falls off with increasing distance from a bleb edge both on and off blebs, PIP₂ localization falls off with increasing distance from a bleb edge only in regions that are not identified as blebs. Similarly, high PIP, activity associates with intermediate local bleb densities (Supplementary Fig. 11d), whereas low PIP₂ activity associates with small but not with high local bleb densities. This shows a specific recruitment of active PIP₂ to the entire bleb surface. The mechanism underlying this process remains elusive.

Our workflow supports many other types of analysis relating cell morphology and molecular distributions. For example, underlying the extraction of morphological motifs are geometric properties that can be analyzed. Measuring bleb volume, we found that small and large blebs show a similar association with $Kras^{V12}$ (Fig. 6g), whereas large blebs show greater association with PIP_2 than small blebs (Fig. 6n). Cytosolically labeled cells show no association of intensity with bleb volume (Supplementary Fig. 11f).

Since the study of many signaling pathways benefits from measuring not just morphology, but also morphodynamics, we developed a measure of boundary motion at each mesh face. Figure 60 shows the PIP₂ activation of an MV3 cell, and Fig. 6p shows that cell's boundary motion. Measuring the motion difference over ~30 s, which is on the order of the bleb lifetime²⁴, we found that blebs preferentially associate with regions of protrusive motion (Fig. 6q). We also observed that regions of high PIP₂ tend to be more retractive than regions of low PIP₂ (Fig. 6r), which is consistent with increased PIP₂ localization on blebs because blebs form and retract cyclically. These and other evidence of relations between local surface geometry and PIP₂ activation will be essential to uncovering the mechanism of a bleb-formation and bleb-size dependent organization of PIP₂ signals.

Discussion

High-resolution 3D light-sheet microscopy^{6–11}, has enabled the direct observation of subcellular molecular processes. However, incorporating these observations into a framework for unbiased data exploration, hypothesis testing and ultimately the development of new biological theories remains a challenge.

Most publications describing innovations in 3D microscopy end with the appealing rendering of a few images on a 2D screen. Even this mere visualization task imposes a particular perspective and thus introduces bias¹⁴. Moreover, compared to one- and two-dimensional features, such as length and area, human observers exhibit decreased ability to assess 3D features, such as shape and volume²⁵. Thus, to turn innovation in 3D imaging into biological insight, computing infrastructures are required that minimize the need for human visual interpretation when comparing datasets.

Here, we focused on algorithms that enable the analysis of biological surfaces at the scale of single cells. We developed an algorithm to detect diverse morphological motifs on the cell surface using machine learning. As a demonstration, we trained classifiers for blebs, filopodia and lamellipodia, among other motifs. To detect a new type of morphological motif, users need only click on examples of surface regions that are and are not that motif. This detector is one of the first image analysis tools for cell biology that incorporates techniques from computer graphics. With the rapid rise of 3D microscopy, computer graphics methods will become an important factor in biological discovery.

In addition to a morphological motif detector, we developed an integrated suite of tools for investigating the coupling between morphology, morphology change and intracellular signaling. Since signaling networks are usually highly nonlinear, the spatial distribution of signaling molecules can greatly affect downstream signaling. Cells take advantage of this effect to control signaling via spatial localization in myriad ways including compartmentalization, phase separation and active transport. Cell morphology may also govern signaling. For example, we found that on blebby melanoma cells both Kras^{V12} and PIP₂ polarize with blebs. PIP₂ is enriched on blebs, whereas Kras^{V12} is not. Investigating further, we discovered that Kras^{V12} localizes to bleb edges and that its distribution is consistent with that of a membrane label. Together, these data suggest the possibility that membrane wrinkling alone or enrichment on blebs could modulate nonlinear signaling networks by concentrating membrane-bound proteins. These two examples also illustrate how u-shape3D supports the acquisition of maps and statistics of the spatial modulation of protein concentrations that would be inaccessible by visual inspection, and the numerical treatment of

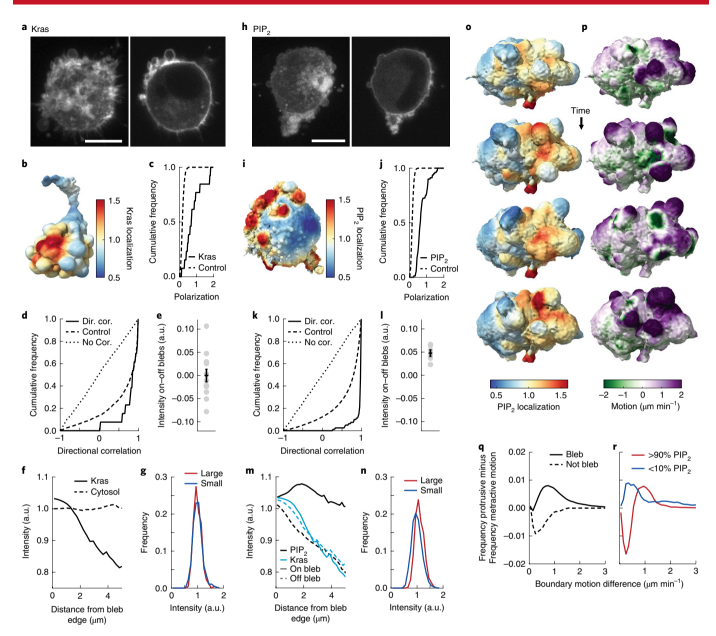


Fig. 6 | Kras and PIP₂ associate with blebs differently. a, An MV3 cell expressing GFP-Kras^{V12} shown as a MIP (left) and an xy-slice (right) (representative of 31 cells). **b**, Kras localization, measured over 2 µm, near the surface of an MV3 cell expressing GFP-Kras^{V12}. **c**, For 13 cells, the cumulative polarization distribution of Kras intensity (solid line) compared to random (dashed line). d, The directional correlation (dir. cor.) of blebs with Kras localization. The cumulative correlation distribution is shown as solid, the control distribution is shown as dashed and the zero correlation (no cor.) distribution is shown as a dotted line. The correlation and control populations are not statistically different (P=0.3; Kolmogorov-Smirnov statistic = 0.3). \mathbf{e} , The differences between the mean Kras intensity on and off blebs (P = 0.5; effect size = -0.006; t-statistic = -0.017). The error bar indicates the standard error of the mean. f, Fluorescence localization versus distance from a bleb edge for 13 GFP-Kras^{V12} labeled cells and 35 GFP cytosolically labeled cells. g. Distributions of Kras intensity for mesh faces on blebs of greater than average volume and on blebs of less than average volume (P = 0.6; effect size = 0.05; Kolmogorov-Smirnov statistic = 0.05; no. of blebs = 1,425). **h**, An MV3 cell expressing PLC Δ -PH-GFP shown as a MIP (left) and an xy-slice (right) (representative of 23 cells). i, PIP₂ localization, measured over 2 µm, near the surface of an MV3 cells expressing PLCA-PH-GFP. i, For six movies of distinct cells, the cumulative polarization distribution of PIP₂ intensity (solid line) compared to random (dashed line). **k**. The directional correlation of blebs with PIP₂ localization. The correlation and control populations are statistically different ($P = 42 \times 10^{-29}$; Kolmogorov-Smirnov statistic = 0.6). I, The differences between the mean PIP₂ intensity on and off blebs for six movies of cells (P = 0.0005; effect size = 1.7; t = 6.9). The error bar indicates the standard error of the mean. **m**, PIP₂ and Kras localization, both on and off blebs, versus distance from a bleb edge. **n**, Distributions of PIP₂ intensity for mesh faces on blebs of greater than average volume and on blebs of less than average volume ($P = 2 \times 10^{-77}$; effect size = 0.5; Kolmogorov-Smirnov statistic = 0.24, no. of blebs = 10,625). **o**, Surface renderings of PIP₂ localization, measured over 2 μm, near the surface of an MV3 cell expressing PLCA-PH-GFP. Cells were imaged every 37 s. p, Surface renderings of the boundary motion of this same cell. Purple indicates regions of high protrusive motion, whereas green indicates regions of high retractive motion. **q**, For six cells, the frequency of protrusive motion minus the frequency of retractive motion on and off blebs as a function of surface speed. r, The same measure shown in f for mesh faces in the top and bottom deciles of PIP₂ localization. Scale bars, 10 μm.

complex geometric arrangements that are at the root of non-intuitive cell behaviors. In future, these features of u-shape3D will enable projects ranging from cell behavioral screens and fluorescence resonance energy transfer measurements linking signaling to morphology to molecularly specific investigations of 3D signaling in vivo.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, statements of code and data availability and associated accession codes are available at https://doi.org/10.1038/s41592-019-0539-z.

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Author contributions

M.K.D. and G.D. conceived and designed the study. M.K.D. wrote most of the software. A.R.J. developed the software's graphical user interface. E.S.W. and M.K.D. performed most of the imaging experiments. K.M.D. and T.I. performed the U2OS experiments. K.M.D. and R.F. provided imaging resources. M.K.D. and G.D. wrote the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Correspondence and requests for materials should be addressed to G.D.

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Methods

Cell culture and genetic engineering. Cells were cultured at 5% CO₂ and 21% O₂. MV3 melanoma cells (a gift from P. Friedl at the MD Anderson Cancer Center) were cultured using DMEM (Gibco) supplemented with 10% fetal bovine serum. Primary melanoma cells (a gift from S. Morrison at UT Southwestern Medical Center) were cultured using the Primary Melanocyte Growth Kit (ATCC). HBEC (a gift from J. Minna at UT Southwestern Medical Center), immortalized with Cdk4 and hTERT expression and transformed with p53 knockdown, Kras V12 and cMyc expression were cultured in keratinocyte serum-free medium (Gibco) supplemented with 50 mg ml $^{-1}$ of bovine pituitary extract (Gibco), 5 ng ml $^{-1}$ of enhanced growth factor (Gibco) and 1% Anti-Anti (Gibco). U2OS osteosarcoma cells (a gift from R. McIntosh at the University of Colorado, Boulder) were cultured using high-glucose DMEM (Gibco) supplemented with pyruvate, stable glutamine and 10% fetal bovine serum. Conditionally immortalized hematopoietic precursors to dendritic cells That express Lifeact-GFP28 (a gift from M. Sixt, IST Austria) were cultured and differentiated as previously described 23 .

Unless stated otherwise, fluorescent constructs were introduced into cells using the pLVX lentiviral system (Clontech) and selected using antibiotic resistance to either puromycin or geniticin. The GFP-tractin construct contains residues 9–52 of the enzyme IPTKA ³⁰ fused to GFP³¹. The CyOFP-tractin peptide contains the tractin peptide fused to the CyOFP protein. CyOFP is a cyan-excitable orange fluorescent protein with peak excitation at 505 nm and peak emission at 588 nm³². The GFP-Kras^{V12} plasmid was constructed by cloning a Kras^{V12} fragment from the pLenti-Kras^{V12} construct²⁶ into the pLVX-GFP vector. The biosensor for PIP₂, PLCA-PH-GFP, encodes a PI(4,5)P2 lipid selective PH domain that can be used as a fluorescent translocation biosensor to monitor changes in the concentration of plasma membrane PI(4,5)P2 lipids³³. Some MV3 cells expressing GFP in the cytosol and imaged via meSPIM, appeared in a previous publication and were analyzed here as a control population¹⁰.

For the CRISPR knockouts, U2OS cells were transiently transfected with pX458 including gene-specific guide RNAs together with a self-cleaving donor vector to deliver a blasticidin S resistant cassette into the genomic cut site. Cells were selected with 5 µg ml⁻¹ blasticidin S and surviving colonies were isolated using 6 mm Pyrex cloning cylinders (Sigma-Aldrich). The pSpCas9(BB)-2A-GFP (pX458) was a gift from F. Zhang (Addgene plasmid no. 48138). The self-cleaving donor vector pMA-tial1 was a kind gift from T. Buerckstuemmer (Horizon Genomics). Guide RNA sequences were cloned into pX458 by Golden Gate cloning using the BbsI cut site. Guide RNA sequences targeting Wave2 (WASF2, exon 3) and cofilin-1 (CFL1, exon 2) were 5′-TGAGAGGGTCGACCGACTAC-3′, and 5′-CGTAGGGGTCGACCACTAC-3′, respectively. Gene knockout was verified by western blotting using rabbit anti-cofilin-1 (Cell Signaling, D3F9 XP no. 5175) and rabbit anti-Wave2 (Cell Signaling, D2C8 XP no. 3659) antibodies (Supplementary Fig. 12).

Imaging. Unless stated otherwise, imaging was performed via microenvironmental selective plane illumination microscopy 10 , a type of two-photon Bessel beam light-sheet microscopy that confers near-isotropic resolution (300 nm lateral, 340 nm axial) and permits recording of cell behavior several millimeters from mechanically perturbing hard surfaces. Images were acquired at 37 °C in a non-descanned image capture mode with an axial step size of 160 or 200 nm and an excitation wavelength of 900 nm. Melanoma cells were imaged in cell culture medium supplemented with HEPES buffer to maintain the pH during imaging.

Confocal imaging was performed using a Zeiss LSM 780 with a \times 40 (1.4 numerical aperture (NA)) objective. Microglia were imaged within zebrafish using a Zeiss Lightsheet Z.1 with \times 20 detection (1.0 NA) and \times 5 illumination (0.1 NA) objectives. The zebrafish line was P2Y12::P2Y12-GFP and was 3.5 days post-fertilization. ASLM imaging was performed using a custom-built microscope as previously described°.

U2OS cells were allowed to spread overnight in pH-neutralized rat-tail collagen (3 mg ml $^{-1}$) before imaging. All other cells, except for those imaged by the Peri and Betzig laboratories, were imaged in collagen gels created by mixing bovine collagen I (Advanced Biomatrix) with concentrated PBS and water to a collagen density of 2.0 mg ml $^{-1}$. This collagen solution was then neutralized with 1 M NaOH and mixed with cells just before incubation at 37 $^{\circ}$ C to induce collagen polymerization. U2OS cells and MV3 cells imaged via confocal microscopy were fixed in 4% paraformaldehyde before imaging.

Image deconvolution. All microscopy images shown are raw, non-deconvolved images. However, as a first analysis step, we deconvolved each 3D image. Most images acquired via meSPIM were Wiener deconvolved as previously described 10. The microscope's point spread function was measured using fluorescent beads. The Wiener parameter, which is the inverse of the signal-to-noise ratio, was usually set to 0.018. However, to better detect the dim ends of filopodia, it was set to 0.015. For cytosolically labeled cells, we automatically estimated the Wiener parameter in each frame by defining the signal as the average fluorescence intensity within the cell and the noise as the standard deviation of the fluorescence intensity outside the cell. Supplementary Fig. 2 shows the effect of varying the Wiener and other deconvolution parameters, and Supplementary Table 5 shows the deconvolution and surface extraction parameters for all datasets presented in this paper. Since

Wiener deconvolution is sensitive to point spread function quality, for images acquired via microscopy modalities other than meSPIM, we used the Richardson–Lucy deconvolution algorithm built-in to MATLAB. The movie of the MDA-MB-231 human breast cancer cell was deconvolved as previously described²².

Following deconvolution, an apodization filter was applied to the optical transfer function of the image in the spatial frequency domain. This filter had a value of 1 at the origin and decayed linearly to 0 at the edge of the filter support, which is set by the user as a percentage of the maximum optical transfer function value. This threshold value, here termed the apodization height, was usually adjusted according to the homogeneity of the fluorescence label and the fineness of the morphological motif being detected. Higher apodization heights smooth the image more and allow for more robust detection of large objects, whereas lower apodization heights allow for the detection of finer structures but also admit more noise.

Cell surface extraction. The deconvolved images were further processed before cell surface extraction. For most datasets, an Otsu threshold was first calculated from the 3D image³⁴, holes were filled using a 3D grayscale flood-fill operation and objects disconnected from the main cell were removed. We also optionally smoothed the image with a 3D Gaussian kernel and applied a gamma correction. MATLAB's *isosurface* function was then used to create a triangle mesh at the intensity value specified by the Otsu threshold. Finally, the triangle mesh was smoothed using curvature flow smoothing³⁵.

For some datasets, this procedure does not segment the nucleus along with the cytoplasm. In these cases, we therefore combined the output image of the procedure described above with an 'inside' image that segmented the cell interior. To create the 'inside' image from the gamma corrected image, we applied an additional gamma correction, smoothed the image with a 3D Gaussian kernel of standard deviation 2 pixels, Otsu thresholded the image, morphologically dilated the image, filled holes in each *xy*-slice, morphologically eroded the image by a radius greater than the morphological dilation and finally smoothed the binary image with a 3D Gaussian kernel of 1 pixel width. Since this process shrinks the cell, if the parameters are chosen correctly the edges of the morphological motifs should mostly lay outside the 'inside' image. To combine the 'inside' image with the image outputted by the procedure above, we normalized this image by its Otsu threshold value, took the pixel-by-pixel maximum of this image and the 'inside' image, and extracted a triangle mesh as an isosurface at an intensity level of one.

The ends of the long, thin lamellipodia of dendritic cells fail to segment using the techniques described above. To better segment lamellipodia, we combined the 'inside' and normalized deconvolved images described above for PIP2 labeled cells with a 'surface filtered' image that enhances planar features, such as lamellipodia (Supplementary Fig. 3). The surface filter, which was developed by Elliott et al. 23, uses multiscale Gaussian second order partial-derivative kernels of the form

$$s(x)_{i,\omega} = \frac{1}{s_{i,\omega}\sqrt{2\pi}} \sum_{x'}^{x' \in \Omega_k} \frac{\partial^2 e^{\frac{-|x-x'|^2}{2\sigma_{i,\omega}^2}}}{\partial x_i^2} I(x')$$

$$\tag{1}$$

where I(x) is the image intensity, $\sigma_{i,\omega}$ is the half width of the Gaussian in dimension i at scale ω , Ω_k is the filter kernel support and $s(x)_{\omega}$ is the filter response at scale ω . The total filter response, S(x), is merged across scales via

$$S(x) = \max(\{|s(x)_{\omega}| \ \sigma_{\omega}|\omega = 1, ..., n\})$$
 (2)

$$\sigma_{\omega} = 2^{\omega - 1} \tag{3}$$

We used filter scales 1.5, 2 and 4 pixels to segment lamellipodia of various thicknesses. To combine the response of the surface filter with the 'inside' and normalized deconvolved images, we normalized the response by subtracting both the mean image intensity and twice the standard deviation of the image intensity before dividing by the standard deviation of the image intensity.

Although not used in this paper, our software also includes the option to segment cells by combining a normalized deconvolved image with a steerable filtered image. Steerable filters are computationally efficient edge detectors that, depending on the parameters chosen, enhance linear or planar structures at specified scales 36,37.

Segmentations were spot checked by thresholding the 3D image at the isosurface intensity value immediately before mesh extraction and examining the overlaid raw and thresholded images as 3D image stacks in Image]³⁸ (Supplementary Fig. 13). For analyses where internal mesh cavities could alter results, meshes were also exported to ChimeraX³⁹ for further examination. Segmentations that were found to be inaccurate or had cavities were excluded from further analysis.

Decomposition of the cell surface into convex patches. Although the image deconvolution and cell surface extraction parameters require customization for different cell types, the remainder of the workflow does not, and its parameters were kept constant throughout the paper.

To decompose the cell surface into convex patches, we first performed a watershed segmentation of surface mean curvature, as previously described10. This oversegments the cell surface into small patches, which are analogous to superpixels in image analysis, which we later merge to create convex patches. First, we calculated the mean and Gaussian curvature at every triangle face^{17,23}. Next, we constructed an adjacency graph of faces where each face is a node that is connected to exactly three other spatially adjacent faces. MATLAB's isosurface function does not always produce triangle meshes with sufficient topological consistency to create such a graph. Our software fixes common topological inconsistencies, such as triangular edges that are only connected to one face. Rarely, however, a face graph cannot be constructed. In these situations, very slightly changing the image deconvolution parameters usually solves the problem, although we did not need to do so here. Since curvature can be noisy, we next smoothed mean curvature in two different ways. First, we used a kd-tree to median filter curvature in 3D space over two pixels. The meSPIM is Nyquist sampled, and so 2 pixels, which is 320 nm, is approximately the microscope's spatial resolution. Second, to reduce spurious curvature fluctuations, we diffused mean curvature on the mesh using a diffusion kernel40,41 according to the equation

$$\mathbf{S} = \bar{A}^k \mathbf{R} \tag{4}$$

for 20 iterations, where $\bf R$ is the curvature, $\bar A$ is a normalized, weighted adjacency matrix of the faces graph, k is the number of iterations and $\bf S$ is the smoothed curvature. We defined A as

$$A_{ij} = \begin{cases} 1, & \text{if } i = j \\ \frac{1}{d_{ij}}, & \text{if } i \text{ is adjacent to } j \\ 0, & \text{otherwise} \end{cases}$$
 (5)

where d_{ij} is the distance between faces i and j. To normalize A, we multiplied it by a diagonal matrix, where each diagonal element was the inverse of the sum of that row. Next, we performed a watershed segmentation of the smoothed curvature over the cell surface¹⁸. Watershed segmentations are often performed on 2D images, where each pixel is adjacent to exactly four other pixels. Here, we similarly performed a watershed segmentation over the adjacency graph of faces, where each face is adjacent to exactly three faces.

We next merged adjacent patches using a spill depth criterion ¹⁸. Here, the spill depth between two adjacent patches was defined as the maximum curvature of the two patches minus the maximum curvature at the patch–patch interface. This is analogous to the depth of water that the patch can hold before spilling into the neighboring patch. Starting with the smallest spill depth, we merged patches until no spill depth was below a cutoff of 0.6 times the Otsu threshold of mean curvature for the cell. Supplementary Fig. 4 shows the effect of altering the spill depth cutoff and other patch-merging parameters.

Finally, we decomposed the surface into approximately convex patches by iteratively applying the triangle and LOS criteria. To apply the triangle criterion to the first calculated the closure surface area for each patch and pair of adjacent patches. We defined the closure surface area as the minimum additional surface area needed to create a closed polyhedron from a surface patch. We then merged adjacent patches if they meet the criterion

$$\frac{\sigma_A + \sigma_B - \sigma_{AB}}{\sqrt{\sigma_A \sigma_B}} > \rho \tag{6}$$

where σ_A and σ_B are the closure surface areas of the two patches, σ_{AB} is the closure surface area of the merged patch and ρ is the triangle cutoff parameter, which we here set to 0.7. The triangle criterion can be thought of as similar to the law of cosines and intuitively seeks to merge patches that meet at small angles. Starting with the largest ρ , we merged all pairs of patches that met the triangle criterion before applying the LOS criterion.

The LOS criterion merges adjacent patches with high mutual visibility^{19,42}. We defined the mutual visibility of patches A and B as the percentage of line segments that connect a face in A with a face in B that are lines of sight, where a LOS is a line segment that falls entirely within the mesh. We calculated mutual visibility by randomly selecting a face on each patch, and using a triangle-ray intersection 43 algorithm to determine whether a line segment connecting the two faces exited and reentered the mesh. A small patch and an adjacent very large patch may have a large mutual visibility because of lines of sight that extend across the width of the cell, even if these two patches should not be merged. When merging two patches, we therefore discarded line segments that were longer than twice the smaller patch size. Supplementary Fig. 14a shows the convergence of mutual visibility as a function of the number of line segments tested. We calculated mutual visibility from 20 line segments per pair of patches. In an exact convex decomposition, any two points within any patch could be connected by a LOS. However, because of biological variation and image noise, requiring a mutual visibility of one is too strict a requirement for cell images. We instead merge patches if their mutual visibility is greater than 0.7. Starting with the largest mutual visibility between patch pairs, we merged all patch pairs meeting the LOS criterion, before again applying the triangle criterion.

Having three patch-merging criteria for convex surface decomposition allows us to balance accuracy, speed and robustness to noise. The spill depth criterion is fast but potentially inaccurate, whereas the LOS criterion is relatively slow, but exact. The triangle criterion implements the short-cut rule²⁰, which biases merging toward certain types of convex decomposition. By adjusting the three merging parameters, users can control which criteria dominate in their analysis.

Classification of morphological motifs. To classify each patch by morphological motif, we first performed feature selection on the geometric patch features listed in Supplementary Table 1. Implemented by the MATLAB built-in function sequentialfs(), our sequential feature selection randomly successively removed features as long as doing so reduced the misclassification rate. The misclassification rate was measured using ten-fold cross validation. The geometric features selected can vary considerably from dataset to dataset even for similar training sets, presumably because of correlations between features, randomness and dataset differences. For example, Supplementary Table 6 shows the features selected for bleb detection models generated by three different users training on the same four cells. In this example, no feature was selected by all three models and no two models shared more than two selected features. Once features were selected, features were normalized to have the same mean and standard deviation, and a linear SVM44 was used to classify patches. Since SVM models vary from user to user, to analyze actin, Kras and PIP2 localization, we had models created by three different users vote on the classification of each bleb.

We also validated our workflow with the linear SVM replaced with a radial SVM or a random forest 45 . Supplementary Table 6 shows the precision, recall and F_1 score of these algorithms for various iterations of feature selection. For the radial SVM, we used the Gaussian kernel,

$$K(\mathbf{x}_{s},\mathbf{x}_{k}) = e^{-\|\mathbf{x}_{j} - \mathbf{x}_{k}\|^{2}}$$

$$\tag{7}$$

To implement the random forest, we used the treeBagger() function in MATLAB. Measuring the out-of-bag classification error as a function of the number of trees grown, we observed that the error plateaued at approximately ten trees, which is well below the 30 and 200 tree forests that we tested.

To compare our workflow, which employs a supervised machine-learning algorithm, to an unsupervised algorithm, we performed an agglomerative hierarchical clustering on all the patches and the patches classified by the supervised algorithm as motifs of interest (Supplementary Fig. 7), respectively. We used the correlation as a distance metric and measured the distance between a pair of clusters as the average distance between any two pairs of patches in these clusters. To avoiding biasing the algorithm, we only clustered on statistics defined at the patch scale, and did not include cell scale statistics, such as cell volume.

Characterization of patches. To classify patches by morphological motif, we calculated geometric descriptions of each patch. The full list of 23 features used by the SVM classifier is provided in Supplementary Table 1. In calculating these features, mean curvature was smoothed as described above, but Gaussian curvature was not. We defined the average patch position as the mean location of the faces in the patch, and we similarly defined the weighted average patch position as the mean location of the faces weighted by curvature. The feature 'variation from a sphere' was defined by the standard deviation of the distances from a patch's faces to the average patch position divided by the mean distance of those faces to the average patch position. We defined the closure surface area as described above. The closure center was also defined as the mean position of the mesh vertices at the patch edge. We defined the patch radius as the mean distance of the patch's faces from the closure center.

The volume, V, was calculated using the equation

$$V = \frac{1}{6} \sum_{i}^{N} \mathbf{v}_{1,i} \cdot (\mathbf{v}_{2,i} \times \mathbf{v}_{3,i})$$
 (8)

where N is the number of faces, and $\mathbf{v}_{1,p}$ $\mathbf{v}_{2,i}$ and $\mathbf{v}_{3,i}$ are the vertices of face i. The vertices must be ordered such that the face normal extends outward from the cell. To derive this equation, the mesh can be thought of as decomposed into tetrahedrons where the vertices of each tetrahedron are those of a face combined with the origin⁴⁶. The signed volumes of the tetrahedrons sum to the volume of the mesh. Patches were closed before calculating their volumes.

We calculated the shape diameter function similarly to Shapira et al.⁴⁷. For each patch, we randomly picked 20 mesh faces on the patch and extended a ray inward from the mesh face at a randomly chosen angle within $\pi/3$ of the direction opposite to the face's normal. We calculated the distance each ray traveled before intersecting the opposite side of the mesh. The shape diameter function of the patch was then defined as the mean travel distance within one standard deviation of the median distance.

Features selected for patch classification. The feature selection algorithm selected different geometric features to detect the three morphologies. To determine which geometric features best distinguished morphologies, starting from no features, we successively added the most discriminative feature to the model (Supplementary

Table 3). The features that best distinguished blebs from non-blebs were volume/(closure surface area)^{3/2} and mean curvature on the protrusion edge. Closure surface area is the minimum amount of additional surface area needed to create a closed polygon from the mesh of the patch. The features that best distinguished filopodia from non-filopodia were the distance from the center of the closure surface area to the mean face position, a measure of morphological feature length and patch surface area. This same measure of morphological feature length as well as patch volume were the best features for distinguishing lamellipodia from non-lamellipodia.

Optional merging of convex patches. Some morphological motifs, such as lamellipodia and flagella, are not convex but are composed of multiple convex regions. To detect such motifs, we optionally merge convex patches into patch composites. Since adjacent patches that compose a larger structure often have smooth curvature at their interface, we first merge patches using a modified LOS criterion with line segment length capped at 10 pixels and a mutual visibility cutoff of 0.7. The LOS criterion is described above. This step is not required for convex patch merging and can be disabled by the user. We next employed a more versatile machine learning-based framework to merge adjacent patches. Using the geometric features for pairs of adjacent patches listed in Supplementary Table 2, as well as user provided training data, we trained an SVM to automatically merge patches. We used the same feature selection procedure and SVM parameters as for patch classification.

Characterization of adjacent patches. To merge adjacent patches into patch composites using an SVM, we calculated geometric characterizations of each pair of adjacent patches. The full list of 36 features used by the SVM is provided in Supplementary Table 2. Some measures of patch pairs incorporate individual patch measures, which are described above. Unless otherwise specified, mean curvature was smoothed as described above, but Gaussian curvature was not.

To better describe the surface geometry at patch-patch interfaces, we calculated the two principal curvatures, κ_1 and κ_2 , everywhere on the cell surface,

$$\kappa_1 = H + \sqrt{H^2 - K} \tag{9}$$

$$\kappa_2 = H - \sqrt{H^2 - K} \tag{10}$$

where H is the unsmoothed mean curvature and K is the unsmoothed Gaussian curvature. For various geometries defined by principal curvature values, we then calculated the fraction of the interface that had that geometry. As a noise threshold, we used the standard deviation of the smoothed mean curvature. Principal curvatures above this threshold or below the negative of this threshold were defined as large, and those more than four times above or below it as very large. We defined a ridged geometry as a large positive κ_1 and a small κ_2 , a very ridged geometry as a very large positive κ_1 and a small κ_2 , a valley-like geometry as a small κ_1 and a large negative κ_2 , a very valley-like geometry as a small κ_1 and a very large negative κ_2 , a domed geometry as a large positive κ_1 and a large positive κ_2 , a cratered geometry as a large negative κ_2 , and a saddle-like geometry as a large negative κ_1 , and a large negative κ_2 , and a saddle-like geometry as a large negative κ_3 , and a large negative κ_4 .

Generation of training data. We designed a graphical user interface to enable the collection of training data necessary for motif classification. Users are shown a surface rendering of a cell with surface patches outlined and can interact with the cell by rotating and moving it, and zooming in and out on regions of interest. To generate data for patch classification, we asked users to click on patches that are certainly the morphological motif of interest and then subsequently asked them to click on patches that are certainly not that motif. Similarly, to generate data for the optional step of convex patch merging, we asked users to click on pairs of patches that should certainly be merged and then asked them to click on pairs of patches that should certainly not be merged. Pairs of patches that were not adjacent were automatically excluded from the training set. We have successfully tested this interface in MATLAB v.R2017b and v.R2013b. However, since in MATLAB user interface functionality can vary from version to version, it may not work in some versions of MATLAB.

Validation. To validate the protrusion classification, we calculated the F_1 score, which is the harmonic mean of precision and recall. Here, precision is defined as the ratio of patches correctly classified as protrusions to the total number of patches classified as protrusions, whereas recall is defined as the ratio of patches correctly classified as protrusions to the total number of patches that are protrusions. Unless otherwise specified, in calculating the F_1 score, we only used patches selected by the trainer as certainly a protrusion or certainly not a protrusion.

Generation and analysis of synthetic images. For algorithm validation, we created synthetic spherical cells of radius 48 pixels. The cell size was chosen to mimic the pixel spacing on the meSPIM of $0.16\,\mu m$ per pixel for a cell $7.6\,\mu m$ in radius. Placed randomly on the cells' surfaces were spherical blebs that ranged in radius from 2 to 32 pixels and in number from 4 to 256 per cell (See Supplementary Fig. 15 for an

example of synthetic cells). Since pixelation at the cell edge could hamper the cell surface extraction and subsequent analysis, edge pixels were subdivided into a finer 3D grid to calculate the percentage of the pixel occupied by the synthetic cell. The final synthetic images were saved with 32 grayscale intensity values. Synthetic cells were not deconvolved, but the remainder of the analysis workflow was identical to that used for microscopic data. The same surface extraction parameters were used as for bleb detection on tractin and cytosolically labeled cells.

An F_1 score does not measure whether or not the workflow preferentially detects certain subtypes of protrusion. Since patch-merging algorithms could be sensitive to protrusion size, we used synthetic data to test the algorithm's sensitivity to bleb size (Supplementary Fig. 15). On synthetic cells of radius 7.6 µm (48 pixels) we simulated blebs ranging in radius from 0.32 µm (2 pixels) and 0.64 µm (4 pixels) to 5.1 µm (32 pixels). Although only 70% of the smallest 0.32 µm radii blebs were decomposed as convex surface patches, almost all of the larger blebs were decomposed. A bleb detector trained on synthetic data correctly classified all blebs that were decomposed as convex surfaces.

Mapping fluorescence intensity to the cell surface. To measure the fluorescence intensity local to each mesh face, we used the raw, non-deconvolved, fluorescence image. At each mesh face, we used a kd-tree to measure the average pixel intensity within the cell and within a sampling radius of the mesh face. To correct for surface curvature-dependent artifacts, we depth normalized²³ the image before measuring intensity localization by normalizing each pixel by the average pixel intensity at that distance interior to the cell surface. Before analysis, we also normalized each cell's surface intensity localization to a mean of one.

Calculation of distance from a bleb edge. On the adjacency graph of faces, we calculated the distance from each face to the nearest bleb edge measured in number of faces traversed. To convert this distance to micrometers, we multiplied by the average distance between faces for each cell in each frame. Since the distance in micrometers between adjacent faces varies, our calculation of distance is an estimate rather than exact.

Calculation of local bleb density. To calculate bleb density, we first assigned the value one to each mesh face on a bleb and the value zero to each mesh face not on a bleb (Supplementary Fig. 11a). We then diffused these values on the mesh surface using equation (4) over 600 iterations (Supplementary Fig. 11b). We choose the number of iterations such that the bleb density would be calculated over a short distance on the order of a bleb length. Equation (4) does not allow an exact measurement of bleb density and may be unstable over distances on the order of many bleb lengths.

Spherical statistics. The von Mises–Fisher distribution is defined on an \mathbb{R}^{d-1} sphere within \mathbb{R}^d space⁴⁸. For d=2 dimensions it approximates a wrapped normal distribution on a circle and, similar to the normal distribution, for any d is parameterized by a mean and an inverse spread. For d=3 dimensions, the von Mises–Fisher distribution is

$$p(\mathbf{x}; \boldsymbol{\mu}, \kappa) = \frac{\kappa}{2\pi (e^{\kappa} - e^{-\kappa})} \exp(\kappa \boldsymbol{\mu}' \mathbf{x})$$
(11)

where μ is the mean direction parameter and κ is the concentration parameter, which is inversely related to the data spread. The maximum likelihood estimate of the mean direction is simply

$$\mu = \frac{\sum_{i}^{N} \mathbf{x}_{i}}{\left|\sum_{i}^{N} \mathbf{x}_{i}\right|} \tag{12}$$

A Newton's method approximation for κ , κ_2 , in three dimensions is

$$\overline{R} = \frac{\left| \sum_{i}^{N} \mathbf{x}_{i} \right|}{N} \tag{13}$$

$$A(\kappa) = \frac{I_{\frac{3}{2}}(\kappa)}{I_{\frac{1}{2}}(\kappa)}$$
 (14)

$$\hat{\kappa} = \frac{\overline{R} (3 - \overline{R}^2)}{1 - \overline{R}^2} \tag{15}$$

$$\hat{\kappa}_1 = \hat{\kappa} - \frac{A(\hat{\kappa}) - \overline{R}}{1 - A(\hat{\kappa})^2 - \frac{2}{c}A(\hat{\kappa})}$$
(16)

$$\hat{\kappa}_2 = \hat{\kappa}_1 - \frac{A(\hat{\kappa}_1) - \overline{R}}{1 - A(\hat{\kappa}_1)^2 - \frac{2}{\hat{\kappa}_1} A(\hat{\kappa}_1)}$$
(17)

where N is the number of data vectors and I are Bessel functions of the first kind48.

In Fig. 6c,j, we measured the magnitude of PIP2 and Kras V12 polarization by mapping the intensity values defined on each surface mesh onto a unit sphere and then using spherical statistics to calculate κ . To map the intensities onto the unit sphere, we calculated a set of unit vectors, $\mathbf{x}_{\text{intensity}}$ that extended in the direction from the cell center to every mesh face. The cell center was defined as the location within the cell farthest from the cell edge. Since we measured intensity at every mesh face over a radius of $2\,\mu\text{m}$, to avoid spatially oversampling, we used only every 1000th mesh face. We defined $w_{\text{intensity}}$ as the measured intensity value associated with each unit vector and then discretized the range of intensity values into 32 bins. Finally, we replaced every vector $\mathbf{x}_{\text{intensity}}$ with w copies of that vector and calculated κ from this set of unit vectors. As a control, we also measured κ from a set of $\mathbf{x}_{\text{intensity}}$ with randomized directions.

In Fig. 6d,k we computed the directional correlation of morphological motifs, here blebs, with intensity localization. In each frame, we defined the directional correlation as $\mu_{\text{blebs}} \cdot \mu_{\text{intensity}}$. To measure μ_{blebs} , we calculated a set of unit vectors, x_{blebs} , that extended in the direction from the cell center to each mesh face on a bleb. To measure $\mu_{\text{intensity}}$, we calculated $x_{\text{intensity}}$ and in equation (12) we weighted x_i by the intensity localization. Since the cell is not a sphere and most cells have polarized shapes, the surface itself is expected to have a nonrandom μ and a small κ . To account for this, we created a control distribution of directional correlations $\mu_{\text{blebsRand}} \cdot \mu_{\text{intensity}}$, where $\mu_{\text{blebsRand}}$ was calculated from a set of vectors where the patch classification was randomly permuted. In each frame, we created 200 such permutations by randomly assigning patches to be a bleb or not a bleb.

Measurement of boundary motion. To measure boundary motion, for each face we found the closest face in the previous frame using a kd-tree. We then defined the boundary motion as

$$m_i = -\operatorname{sign}(\mathbf{d}_i \cdot \mathbf{n}_i) |\mathbf{d}_i|$$
 (18)

where m_i is the boundary motion at face i, \mathbf{d}_i is the vector from face i to the closest point in the previous frame, and \mathbf{n}_i is the normal to the surface at face i.

This is not an ideal measure of boundary motion since the mapping vectors \mathbf{d}_i may cluster on select faces of the previous frame's surface, or even alter the topology among faces, in a physically unrealistic manner (see Machacek et al.⁴⁹ for an illustration of these problems with 2D boundaries). As a control, we also calculated the boundary motion for each face by finding the closest point in the next frame. Supplementary Fig. 14b shows the protrusive and retractive motion of six cells using both definitions of boundary motion. Here, backward motion is the mapping of points from each frame to the previous frame and is the definition used elsewhere, and forward motion is the mapping of points from each frame to the subsequent frame. Even though the backward and forward motions of the cells are different, in both cases blebs are more protrusive than non-blebs. This measure is also not a subpixel measure of motion, and should not be used to measure subpixel motions. Because we map each face to the closest face rather than the closest surface point in the previous frame, motions that are less than the average distance between faces will be undersampled in the motion distribution.

Statistical hypothesis testing. For each Kras and PIP_2 labeled cell, we measured the mean intensity localization of faces on and off blebs and then performed a one-sided t-test on the differences of the means after testing for normality using a Kolmogorov–Smirnov test. The Cohen's d effect size was measured.

Unless otherwise indicated, all errors and error bars show the standard error of the mean.

Surface rendering. Most triangle meshes were rendered in ChimeraX³⁹. Colored triangle meshes were exported from MATLAB as Collada.dae files using custom-written code and were rendered using full lighting mode. Lighting intensity and ambient intensity were adjusted. Colormaps were modified from colorBrewer⁵⁰. The surfaces in Supplementary Figs. 11 and 15 were rendered within MATLAB. Our software is capable of rendering all meshes shown in the paper within MATLAB, as well as creating Collada files for export to ChimeraX.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

Data are available from the corresponding author upon reasonable request.

Code availability

The latest version of the software described here, as well as a user's guide, is available from https://github.com/DanuserLab/u-shape3D.

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Corresponding author(s):	Gaudenz Danuser
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Reporting Summary

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X	For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
	Estimates of effect sizes (e.g. Cohen's d , Pearson's r), indicating how they were calculated
	Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

The meSPIM and ASLM microscopes are operated via custom-written software as detailed in the below publications describing their development.

Erik Welf*, Meghan Driscoll*, Kevin Dean, Claudia Schäfer, Jun Chu, Michael W. Davidson, Michael Z. Lin, Gaudenz Danuser, Reto Fiolka. *Co-first-authors. Quantitative multiscale cell imaging in controlled 3D microenvironments. Development Cell, 2016.

Kevin M. Dean, Philippe Roudot, Erik S. Welf, Gaudenz Danuser, and Reto Fiolka. Deconvolution-free Subcellular Imaging with Axially Swept Light Sheet Microscopy. Biophysical Journal, 2015.

Data analysis

Custom code, which will be published with the manuscript, was written in Matlab. Surface renderings were performed by Matlab and UCSF ChimeraX version 0.9 or earlier. Most versions of Matlab released in 2013 or later were used, but the software was most thoroughly tested in MatlabR2018a.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

Policy information about <u>availability of data</u>

Data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The datasets generated for the current study are available from the corresponding author on reasonable request.

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For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-flat.pdf

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size Sample sizes were chosen to be sufficiently large to demonstrate the feasibility of the method. More specifically, for software validation we used large enough samples such that the F1 score converged as a function of the number of cells.

Data exclusions
Only single cells that were not in contact with other cells were analyzed. For morphological motif detection, cells not exhibiting the motif of interest were excluded. Unless stated otherwise, cells where surface extraction failed were also excluded.

Replication We validated the software on multiple and diverse datasets as described in the manuscript.

Randomization Cells were chosen for imaging based on apparent cell health, fluorescence expression, and optical clarity.

Blinding The analysis was automated and the same analysis parameters were used for populations being compared.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Ν	1ateria	ls &	experimental	l systems
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n/a Involved in the study Antibodies

Eukaryotic cell lines

Palaeontology

Animals and other organisms

Human research participants

🔀 🔲 Clinical data

Methods

n/a Involved in the study

ChIP-seq
Flow cytometry

MRI-based neuroimaging

Antibodies

Antibodies used

rabbit anti-cofilin-1 (Cell Signaling; D3F9 XP® # 5175 used at 1:2000 dilution), rabbit anti-Wave2 (Cell Signaling; D2C8 XP® #3659 used at 1:2000 dilution), rabbit anti-GFP (Rockland Inc; 600-401-215 used at 1:2000 dilution), and mouse anti-GAPDH (GeneTex, GTX627408 used at 1:5000 dilution)

Validation

- Anti-Cofilin-1 antibodies were reported by the manufacturer to react with endogenous levels of Cofilin-1 from Human, Mouse, Rat, Monkey, Dog. We have further confirmed the specificity of the antibodies by blotting for cells knocked out for cofilin-1.
- Anti-Wave2 antibodies were reported by the manufacturer to react with endogenous levels of Wave2 from Human, Mouse, Rat, Monkey. We have further confirmed the specificity of the antibodies by blotting for cells knocked for Wave2.
- Anti-GFP antibodies were reported by the manufacturer to react with wt and all variants of GFP.
- Anti-GAPDH antibodies were reported by the manufacturer to react with GAPDH from Human, Mouse, Rat, Zebrafish, Yeast, Drosophila, Pig, Monkey, E. coli.

Eukaryotic cell lines

Policy information about <u>cell lines</u>

Cell line source(s)

The MV3 melanoma cells were a gift from Peter Friedl (MD Anderson), the transformed HBEC cells were a gift from John Minna and Jerry Shay (UT Southwestern), the U2OS osteosarcoma cells were a gift from Dick McIntosh (University of Colorado, Boulder) and the dendritic cell precursors were a gift from Michael Sixt (IST Austria).

Authentication

None of the cell lines used were authenticated.

Mycoplasma contamination

The MV3, HBEC, and U2OS cell lines tested negative for mycoplasma contamination. The dendritic cell precursors were not tested.

Commonly misidentified lines (See ICLAC register)

No commonly misidentified cell lines were used.