RESEARCH ARTICLE

The spectraplakin Short stop is an essential microtubule regulator involved in epithelial closure in Drosophila

Zsanett Takács1,*, Ferenc Jankovics1,*,†, Péter Vilmos1, Péter Lénárt2, Katja Röper3 and Miklós Erdélyi1

ABSTRACT

Dorsal closure of the Drosophila embryonic epithelium provides an excellent model system for the in vivo analysis of molecular mechanisms regulating cytoskeletal rearrangements. In this study, we investigated the function of the Drosophila spectraplakin Short stop (Shot), a conserved cytoskeletal structural protein, during closure of the dorsal embryonic epithelium. We show that Shot is essential for the efficient final zipper of the opposing epithelial margins. By using isoform-specific mutant alleles and genetic rescue experiments with truncated Shot variants, we demonstrate that Shot functions as an actin–microtubule cross-linker in mediating zipperping. At the leading edge of epithelial cells, Shot regulates protrusion dynamics by promoting filopodia formation. Fluorescence recovery after photobleaching (FRAP) analysis and in vivo imaging of microtubule growth revealed that Shot stabilizes dynamic microtubules.

The actin- and microtubule-binding activities of Shot are simultaneously required in the same molecule, indicating that Shot is engaged as a physical crosslinker in this process. We propose that Shot-mediated interactions between microtubules and actin filaments facilitate filopodia formation, which promotes zipperping by initiating contact between opposing epithelial cells.

KEY WORDS: Drosophila, Spectraplakin, Short stop, Dorsal closure, Microtubule, Actin

INTRODUCTION

Towards the end of Drosophila embryogenesis, an epithelial discontinuity is formed at the dorsal surface of the embryo, which is covered by amnioserosa cells. During dorsal closure, this gap is closed by the dorsal epithelium while the amnioserosa disintegrates. Dorsal closure involves two distinct mechanisms: convergence of two opposed epithelial sheets towards the dorsal side, and subsequent zipperping of the hole at the dorsal midline of the embryo (Jacinto et al., 2002). At the onset of the closure, cells in the first row of the embryonic epithelium differentiate into dorsal-most epithelial (DME) cells and establish a straight movement front, which initiates dorsal-ward migration. Migrating epithelial sheets first meet at the anterior- and posterior-most ends of the dorsal opening where they fuse by a zipper-like mechanism. Fusion of the opposing epithelial cells spreads from the two zippering corners towards the center of the opening, resulting in a typical lens-shaped outline of the dorsal hole throughout the entire process. During zipperping, interacting surfaces of opposing DME cells form lamellar overlaps, which are resolved by shortening and concomitant thickening to achieve seamless closure of the dorsal epithelium (Eltsov et al., 2015).

Genetic and biophysical investigations revealed that the dorsal opening has to be closed in a tightly regulated and efficient manner (Hutson et al., 2003). Several forces provided by various tissues contribute to the closure process, and loss of one of these forces can be compensated for by the others. In these cases, the dorsal opening is sealed, but the dynamics of the closure is abnormal. Mutations leading to abnormal closure dynamics – although not necessarily causing morphological abnormalities – might have evolutionarily relevance.

Efficient dorsal closure requires the dynamic rearrangement of the cytoskeleton in epithelial cells (Martin and Parkhurst, 2004). DME cells form a leading edge facing towards the dorsal opening, where they accumulate an actomyosin cable. In addition, DME cells extend actin-rich cellular protrusions, such as filopodia and lamellipodia, mediating the initial contact between the opposing DME cells. Dynamic filopodia are essential both for the mechanics of epithelial adhesion during dorsal closure and for the correct ‘matching’ of opposing cells (Hakeda-Suzuki et al., 2002; Harden et al., 1999; Jacinto et al., 2000; Jankovics and Brunner, 2006; Woolner et al., 2005).

The microtubule (MT) network has also been demonstrated to rearrange during dorsal closure (Jankovics and Brunner, 2006; Kalschmidt et al., 2002). At the onset of closure, DME cells display an irregularly distributed network of MTs. During closure, MTs reorganize to form acentrosomal bundles that are aligned along the dorsal–ventral cell axis. Although the bundles are stable, individual MTs remain highly dynamic, and at the leading edge they grow into cell protrusions (Jankovics and Brunner, 2006). MTs are dispensable for dorsal-ward migration of the epithelium but contribute to efficient zipperping of the epithelial hole by two distinct mechanisms at two consecutive steps. In the early zipperping phase, MTs at the leading edge regulate protrusion dynamics to promote initial interactions between DME cells. During later stages of zipperping, shrinking MTs are attached with their plus-ends to newly formed cell adhesions, where they are thought to provide a MT motor-based force to resolve areas of the opposing DME cells that have overlapping lamellae (Eltsov et al., 2015).

During morphogenesis, not only the proper organization of the MT and actin networks but also the coordination of their interactions is essential for dynamic cell shape changes. In the past years, an expanding number of proteins and protein complexes mediating structural connections between actin and MTs have been described, among which spectraplakins represent an important class of evolutionarily conserved actin–MT crosslinkers (Coles and Bradke, 2015; Suozzi et al., 2012).

Spectraplakins can interact with MTs through two distinct domains localized at their C-terminus. The C-tail and the GAS2
domains mediate the association of spectraplakins with growing plus-ends and along the MT lattice, respectively (Alves-Silva et al., 2012; Applewhite et al., 2010). The C-tail domain contains three MT tip localization signals (MLS), which target spectraplakin to the MT plus-end by linking it to EB1 (also known as MAPRE1 in mammals), a core component of the plus-end tip complex. At the N-terminal end of spectraplakins, two calponin homology (CH) domains facilitate binding to the actin network. The actin- and MT-interacting domains are separated by a flexible rod domain consisting of a series of spectrin repeats. This structural arrangement of the binding domains enables crosslinking of actin with the growing MT tip or with the MT lattice. Adjacent to the GAS2 domain, spectraplakins possess EF-hand motifs, which regulate autoinhibition of the protein established through an interaction between its N-terminal CH and C-terminal GAS2 domains (Applewhite et al., 2013). In addition to their actin- and MT-interacting domains, spectraplakins harbor a plakin domain enabling their direct binding to adhesive junctions (Röper and Brown, 2003). Through alternative splicing and alternative promoter usage, an enormous number of different protein isoforms, with a wide variety of domain compositions, can be created from a single spectraplakin gene, which enables the spectraplakins to fulfill a wide array of functions in various biological processes (Hahn et al., 2016; Röper and Brown, 2003; Röper et al., 2002).

While vertebrates have two spectraplakins, ACF7 (also known as MACF1) and BPAG1 (also known as DST), the Drosophila genome contains a single spectraplakin homolog short stop (shot) providing a powerful model for studying spectraplakin function (Hahn et al., 2016). In Drosophila, Shot has been shown to be involved in a wide variety of developmental processes, such as neuronal cell growth, tracheal cell fusion and attachment of tendon cells to the muscles. In each case, Shot mediates cytoskeletal rearrangements; however, the exact requirements of its individual functional activities in a given cell type are defined by the cellular context (Bottenberg et al., 2009; Lee and Kolodziej, 2002a; Sanchez-Soriano et al., 2009; Subramanian et al., 2003).

Despite the substantial progress achieved in understanding the functions of spectraplakins, the way these activities are regulated is still not completely clear. The analysis of spectraplakins is complicated by the variability of domain composition, the complex interactions between the domains and the enormous differences in the cytoskeletal organization of the cell types in which spectraplakin function has been studied. Therefore, even functions of individual domains seem to vary between experimental conditions, which means conclusions in one system cannot be applied to other model systems. In addition, analysis of cytoskeletal processes in the intact animal in an unperturbed developmental context is not always possible at an appropriate spatial and temporal resolution. Dorsal closure of the embryonic epithelium in Drosophila, however, provides an excellent model to investigate spectraplakin function. In this study, we demonstrate that Shot is required for the final sealing of the epithelial sheets during dorsal closure. In DME cells, Shot acts as a MT–actin crosslinker to stabilize MTs and regulate proper formation of the MT network. At the leading edge, Shot-mediated MT–actin interactions promote formation of filopodia required for the initial contact between opposing DME cells during zipperung.

RESULTS
Shot is required for efficient zipperung of the epithelial sheets during dorsal closure
To uncover new genes involved in cytoskeletal reorganization and function during dorsal closure, various cytoskeletal regulator genes were silenced by microinjection of embryos with in vitro synthesized double-stranded (ds)RNAs (Table S1). The EGFP signal of the ZASP52ZCL42 protein trap line was used to highlight the outline of the dorsal opening. In vivo imaging of the closure process revealed that, of the genes tested, only the silencing of shot results in abnormal dorsal closure (Fig. 1A; Movie 1). In embryos with reduced shot function, the opening is closed completely, but the dynamics of the closure is abnormal. In the shot RNA interference (RNAi) embryos, the dorsal opening was abnormally narrow, and not the typical wild-type teardrop-shaped dorsal hole. To confirm the RNAi results, the shotf20 mutant allele was used, which behaves genetically as a null allele and has been suggested to abolish all shot functions (Fig. S1). shotf20 homozygous embryos displayed the same abnormal closure phenotype as that induced by silencing shot (Fig. 1A; Fig. S2). To quantify the abnormal shape of the dorsal hole, the length-to-width ratio of the opening was used as a numeric parameter. The phenotype was most obvious towards the middle of the closure; therefore, the closure stage at 30 μm opening width was arbitrarily chosen to quantitatively characterize closure defects. In embryos injected with shot dsRNA and in shotf20 null mutants, the length-to-width ratio significantly increased indicating that shot plays an essential role during the closure of the dorsal hole (Fig. 1B).

The abnormally narrow shape of the opening can arise when the opposed epithelial sheets converge normally while the final zipperung of the sheets is affected. To characterize the abnormal dynamics phenotype, a mathematical model of dorsal closure was applied (Hutson et al., 2003). In shot mutants, the length and width of the dorsal opening were measured throughout the entire closure process (Fig. 1C,D). The velocity of the epithelial sheet translocation (v), as well as the fractional contribution of zipperung (fz) to the velocity of the closure was calculated. The null mutation of shot resulted in a decrease of fz demonstrating that shot function is essential for efficient zipperung (Fig. 1E,F).

To rescue the shot mutant phenotype, a GFP-tagged long isoform of Shot [Shot-L(A)–GFP] was used, which is the longest available transgenic Shot protein version and has been reported to rescue shot mutant phenotypes during embryogenesis (Bottenberg et al., 2009; Lee and Kolodziej, 2002b). The Shot-L(A)–GFP protein was expressed throughout the dorsally migrating epithelium of shot mutant embryos by the pnr-Gal4 driver, and the outline of the dorsal opening was highlighted by mCherry–Moesin. Targeted expression of the long Shot-L(A)–GFP protein rescued the zipperung defect of the shot mutants, as demonstrated by the restoration of the length-to-width ratio of the dorsal opening to the wild-type level (Fig. 1G).

Shot acts as an actin–MT cross linkser in mediating zipperung
In order to understand the role of the individual protein domains of Shot in dorsal closure, we investigated the mutant phenotypes of various shot mutant alleles abolishing distinct Shot activities. In addition, we carried out a detailed structure–function analysis of Shot using a series of shot transgenes in rescue experiments. Hence, GFP-tagged truncated versions of the Shot protein were expressed in the epithelia of shotf20 null mutant embryos by the pnr-Gal4 driver (Fig. 2C).

To investigate the requirement for the actin-binding activity of Shot, we made use of shotΔCH1 mutant embryos, which express Shot isoforms lacking the actin-binding CH1 domain (Fig. S1) (Gregory and Brown, 1998; Lee et al., 2000; Röper et al., 2002). Loss of the CH1 domain alone has previously been shown to lead to loss of F-actin-binding ability from the N-terminus of Shot (Lee and Kolodziej, 2002b). In shotΔCH1 mutants, zipperung of the dorsal hole
was delayed, indicating the requirement for the actin-binding activity of Shot in zippering (Fig. 2A,B; Movie 1). To further investigate the contribution of the actin-organizing activity of Shot to closure, we used a transgene lacking the CH1 domain, mimicking the shotkakP1 mutant allele. Confirming our results obtained with shotkakP1, the shotΔCH1–GFP transgene failed to rescue the shot mutant zippering defect (Fig. 2D). Therefore, we conclude that association of Shot with actin filaments is essential for efficient zippering of the opposing epithelial sheets.

To gain a deeper insight into Shot function during dorsal closure, we studied the requirement for its MT-organizing activity. Crispr/Cas9-based genome editing was applied to generate a novel shot mutant allele specifically impairing its binding to MTs (Table S2). We generated an 1895 bp genomic deletion, which removes the coding sequence of the EF-hand and the Gas2 domains, and, after four additional amino acids, it ends in a premature stop codon. The resulting truncated protein, which we designated ShotΔEGC, lacks the EF-hand, the Gas2 and the C-tail domains, leading to complete loss of the MT-binding activity. Expression of the truncated protein in DME cells was confirmed by immunostaining of shotΔEGC mutant embryos using a polyclonal antibody raised against the spectrin repeats (Fig. S1). In shotΔEGC mutant embryos, we detected the same zippering deficit as in the shotD20 null mutants and the shotkakP1 actin-binding-deficient mutants, suggesting that both the MT-binding and actin-binding activities of Shot are essential for efficient zippering of the epithelial sheets (Fig. 2A,B).
The analysis of shot<sup>ΔEGC</sup> mutant embryos revealed that the C-terminal region composed of the EF-hand, Gas2 and C-tail domains is required for zippering. To further dissect the requirement for the C-terminal region of Shot, we carried out rescue experiments in the shot<sup>f20</sup> null mutant background with transgenes lacking one of these domains (Fig. 2C). Targeted expression of the Shot protein lacking the EF-hand domain (Shot<sup>ΔEF-hand</sup>–GFP) restored the zippering efficiency of shot<sup>f20</sup>.
mutants to the wild-type level, indicating that this domain is not essential for zippering (Fig. 2D). Next, we expressed a truncated Shot protein version lacking the C-tail domain (ShotΔC-tail–GFP), which has been shown to mediate the association of Shot with the growing MT plus-ends by interacting with EB1. In this case, we detected a rescue of the shot mutant zippering phenotype, indicating that interaction of Shot with EB1 and its accumulation at the MT plus-tips is dispensable for dorsal closure (Fig. 2D). The Shot protein variant lacking the MT-stabilizing Gas2 domain (ShotΔGas2–GFP), however, failed to rescue the zippering defect (Fig. 2D). These findings are consistent with the data obtained with the isoform-specific mutant allele shotΔECG, and indicate that it is not the EF-hand or the C-tail domain but rather Gas2 that is required in this cellular context. Expression of a transgenic protein lacking both the CH1 and the Gas2 domains (ShotΔCH1ΔGas2–GFP) failed to rescue dorsal closure defects of the null mutant, confirming the requirement for the actin- and MT-binding activities of Shot (Fig. 2D).

In shotΔECG/shotkakP1 embryos, Shot variants contain exclusively either CH1 or Gas2, but none of them has actin- and MT-binding activities simultaneously. These embryos exhibited the zippering defect, indicating that both the actin- and the MT-binding domains have to be present in the same Shot molecule for proper dorsal closure (Fig. 2A,B; Movie 1).

Taken together, these results demonstrate that both the actin- and MT-binding activities of Shot are required for dorsal closure, and suggest that Shot acts as an actin–MT cross-linker to mediate the zippering step of dorsal closure.

Shot regulates MT organization in epithelial cells during dorsal closure

The MT network has been shown to reorganize during dorsal closure, and its function is required for the zippering of the epithelial sheets (Eltsov et al., 2015; Jankovics and Brunner, 2006; Kaltschmidt et al., 2002). To study the involvement of shot in MT network organization, we examined MT distribution in shot mutant epithelial cells via immunohistochemical labeling. In the cell body of shotΔG null mutants, the overall morphology of the MT network appeared to be slightly disorganized (Fig. 3A,B). At the leading edge of the mutant DME cells, MTs intruded into filopodia-like protrusions; however, the MTs were frequently abnormally long and bent, indicating that shot regulates their proper organization in the cell body and at the leading edge.

To gain a more detailed insight into the MT organization of the shot mutant DME cells, live imaging of embryos expressing Tubulin–EGFP was performed. In the cell body of shotΔG null mutants, sudden bending of MTs was detected (Fig. 3D,D′). Moreover, MTs frequently protruded at the lateral surface of the epithelial cells (Fig. 3E,E′; Movie 2). At the leading edge of the mutant DME cells, long and bent MTs protruded from the cell body over the amnioserosa cells (Fig. 3D). The emergence of abnormal MTs coincided with the onset of the zippering stage, and they extended throughout the entire leading front of the dorsal epithelium (Movie 3). Time-lapse analysis revealed that the maximum length of the protruding MTs was higher in the shotΔG mutants than in wild type (3.8±1.1 µm in wild type, n=108, versus 6.8±3.5 µm in shotΔG mutants, n=109; mean±s.d.;
distribution of acetylated MTs in wild-type and shot–/– mutant DME cells in a representative FRAP experiment. White boxes and white arrows indicate photobleached regions. Scale bar: 5 µm. (B) FRAP recovery curves show the relative GFP fluorescence intensities within the photobleached regions in wild-type (black, n=9) and shot–/– mutant (gray, n=7) DME cells. (C,D) Scatter dot plots of fluorescence recovery half-times (t1/2) and mobile fractions of Tubulin–EGFP measured in wild-type and shot–/– mutants. In shot–/– mutants, t1/2 was increased, whereas the mobile fraction of Tubulin–EGFP was unaffected. (E,F) Scatter dot plots of MT growth rate and lifetime of EB1 comets in wild-type (n=56 comets in two cells from two embryos) and shot–/– mutant (55 comets in two cells from two embryos) DME cells expressing EB1–EGFP. Displacement of EB1–EGFP comets was faster in shot–/– mutants than in wild-type DME cells, whereas the life-time of comets was unaffected. In C–F, results are mean±s.d.; ***P<0.001; ns, not significant (t-test). (G) Projections of ten consecutive time frames of a movie showing EB1–EGFP tracks in wild-type and shot–/– mutant DME cells. The projected time-lapse spans 11 s. (H) Windrose plots of MT growth tracks in wild-type (n=181 in four cells from two embryos) and shot–/– mutant DME cells (n=269 in four cells from two embryos) expressing EB1–EGFP.

Fig. 4. Shot stabilizes dynamic MTs of DME cells by regulating their dynamic properties. (A) Movies showing recovery of Tubulin–EGFP fluorescence in wild-type and shot–/– mutant DME cells. (B) FRAP recovery curves show the relative GFP fluorescence intensities within the photobleached regions in wild-type (black, n=9) and shot–/– mutant (gray, n=7) DME cells. (C,D) Scatter dot plots of fluorescence recovery half-times (t1/2) and mobile fractions of Tubulin–EGFP measured in wild-type and shot–/– mutants. In shot–/– mutants, t1/2 was increased, whereas the mobile fraction of Tubulin–EGFP was unaffected. (E,F) Scatter dot plots of MT growth rate and lifetime of EB1 comets in wild-type (n=56 comets in two cells from two embryos) and shot–/– mutant (55 comets in two cells from two embryos) DME cells expressing EB1–EGFP. Displacement of EB1–EGFP comets was faster in shot–/– mutants than in wild-type DME cells, whereas the life-time of comets was unaffected. In C–F, results are mean±s.d.; ***P<0.001; ns, not significant (t-test). (G) Projections of ten consecutive time frames of a movie showing EB1–EGFP tracks in wild-type and shot–/– mutant DME cells. The projected time-lapse spans 11 s. (H) Windrose plots of MT growth tracks in wild-type (n=181 in four cells from two embryos) and shot–/– mutant DME cells (n=269 in four cells from two embryos) expressing EB1–EGFP.

P<0.001, t-test), indicating that Shot plays a role in regulating MT stability.

In mutant cells, altered MT dynamics, such as faster polymerization rate or lower catastrophe frequency, can lead to longer MTs which, having reached the cell cortex, continue to grow and push out the cell cortex or bend backward. To study MT stability in epithelial cells, we investigated the stabilization state of the MTs by immunolabeling acetylated tubulin. Acetylation of tubulin is a post-translational modification found mainly on long-lived MTs and serves as a hallmark of stable MTs. Consistent with previous findings, we detected acetylated tubulin incorporated into parallel MT bundles in the cell body of wild-type DME cells (Fig. 3F) (Wolf et al., 1988). Using immunohistochemical labeling of acetylated tubulin, we were unable to detect any abnormal distribution of acetylated MTs in shot–/– mutant DME cells (Fig. 3G). Abnormal MTs present at the leading edge of shot mutants were not labeled by anti-acetylated tubulin antibody indicating that Shot regulates the dynamic pool of MTs.

Fluorescence recovery after photobleaching (FRAP) assays were applied to analyze the turnover of tubulin, which reflects the dynamic properties of the MTs. In living embryos, EGFP-tagged β-tubulin was constitutively expressed, which incorporated into the MTs. A 2 µm-wide stripe close to the leading edge of the DME cells was bleached, and the recovery of Tubulin–EGFP fluorescence was imaged by in vivo video microscopy (Fig. 4A). The mobile fraction, reflecting the proportion of proteins undergoing turnover in the MTs, and the recovery half-time, reflecting turnover speed, were determined (Fig. 4B–D). A quantitative analysis of the FRAP curves revealed that the mobile fraction of tubulin was 0.8±0.06 (mean±s.d.) in the wild-type epithelial cells. In mutant DME cells, a similar mobile fraction was detected (0.76±0.13), confirming the finding that Shot does not influence the stabilized subset of the MTs, but rather exerts its function on the dynamic subset (Fig. 4C).

Indeed, the recovery half-time of Tubulin–EGFP (t1/2) decreased to 17.6±3.4 s compared to the wild-type value of 29.3±6.4 s (mean±s. d.) indicating a faster turnover of Tubulin–EGFP between the MTs and the cytosol (Fig. 4D). Thus, MTs in shot mutants are more dynamic, suggesting that Shot affects the MT organization of DME cells by regulating the dynamic properties of MTs.

Faster turnover of Tubulin–EGFP in shot mutants could be a consequence of alterations in the parameters of plus-end dynamics, such as growth speed or time spent on growth. To test the function of Shot in MT growth regulation, EB1–EGFP was expressed in shot–/– null mutant epithelial cells (Movie 4). EB1 binds to polymerizing MT plus-ends, which enables direct measurement of the dynamic instability parameters by in vivo imaging. In mutant cells, the growth rate of MTs reflected by the speed of EB1 comets increased significantly. Instead of the wild-type growth rate of 17.9±3.5 µm/min (mean±s.d.), we measured 24.8±4.6 µm/min in the mutant cells, supporting the finding that Shot regulates MT dynamics (Fig. 4E).

Quantitative analysis of EB1 tracks revealed that the lifetime of EB1 comets remained unaffected in shot–/– mutants indicating that the catastrophe frequency is not influenced by Shot activity (Fig. 4F).

It has been previously shown that spectraplakins can cross-link growing MT tips to actin filaments, which determine the growth
direction of individual MTs (Kodama et al., 2003). To test this activity of Shot, we tracked EB1 comets in the cell body of epithelial cells. In both wild-type and shot<sup>Δ20</sup> mutant embryos, most of the MTs polymerized parallel to the long axis of DME cells, indicating that Shot is not required for the regulation of MT growth direction along the actin filaments (Fig. 4G,H).

Taken together, these results demonstrate that Shot regulates the morphology of the microtubule network by stabilizing the dynamic microtubules in the DME cells.

**MT-binding activity of Shot is required but is not sufficient for MT stabilization**

In order to better understand the role of Shot in MT regulation, we investigated the MT network of epithelial cells in isoform-specific shot mutants. In shot<sup>ΔEGC</sup> mutants, we detected abnormally long and bent MTs at the leading edge, indicating that the MT-binding activity of Shot is required for correct formation of MTs in DME cells (Fig. 5B). To further investigate the MT regulatory function of Shot in DME cells, we expressed various truncated versions of Shot in shot<sup>Δ20</sup> null mutant embryos using the en-Gal4 driver, which drives the expression of the transgenes in only four-cell-wide stripes of the dorsally migrating epithelial sheets. This experimental design enabled us to compare shot-deficient cells with rescued cells in the same embryo (Fig. S3).

The long transgenic Shot protein version, Shot-L(A)-GFP, predominantly colocalized with actin at the cell cortex and in the protrusions, and accumulated at the dorsal actin cable. Shot-L(A)-GFP was faintly detectable along filamentous structures which may correspond to a subset of MTs (Fig. 6A; Movie 5). Shot-L(A)-GFP failed to accumulate at growing MT plus-tips (Fig. S4; Movie 6). Expression of the Shot-L(A)-GFP protein completely rescued the abnormal MT phenotype of the shot<sup>Δ20</sup> null mutants as indicated by the absence of long and bent MTs protruding from rescued DME cells (Fig. 7; Fig. S3B).

The Shot protein variant lacking the C-tail domain (Shot<sub>ΔC-tail</sub>-GFP) localized diffusely in the DME cells and faintly decorated the cortical actin network and protrusions (Fig. 6B; Movie 7). Shot<sub>ΔC-tail</sub>-GFP rescued abnormal MTs, indicating that plus-tip binding of Shot is not required for MT stabilization (Fig. 7; Fig. S3C). This result was confirmed by the expression of the Shot protein mutated exclusively for the EB1 interaction motifs [Shot-L(A)-3MILS<sup>+</sup>-GFP], which localized similarly to Shot-L(A)-GFP and rescued the MT abnormalities of shot null mutants (Figs 6C, 7; Movie 8, Fig. S3D). Consistent with this finding, the C-terminal MT plus-tip-interacting domain of Shot (Shot-C-tail-GFP) on its own showed a localization in DME cells consistent with its previously reported ability to associate with the MT plus tips (Fig. 6D; Movie 9) (Alves-Silva et al., 2012; Applewhite et al., 2010). The expression of Shot-C-tail-GFP failed to rescue abnormal MTs (Fig. 7; Fig. S3E).

To reveal additional domains regulating MT organization, Shot protein lacking the Gas2 domain was expressed in epithelial cells. ShotGas2-GFP lost its ability to bind along the MT lattice and localized to the MT plus-tips and to the actin-rich cell cortex, indicating that the Gas2 domain inhibits the C-tail-mediated MT-tip binding of Shot (Fig. 6E; Movie 10). Expression of the ShotGas2 protein rescued the shot<sup>Δ20</sup> null mutant phenotype, suggesting that the Gas2 domain is dispensable for MT stabilization (Fig. 7; Fig. S3F). The Shot protein version lacking the EF-hand domain (Shot<sub>ΔEF-hand</sub>-GFP) localized similarly to the long Shot-L(A)-GFP and rescued abnormal MTs found in the null mutant cells (Figs 6F, 7; Movie 11, Fig. S3G). The transgenic shot construct composed of the EF-hand, Gas2 and C-tail domains alone (ShotEGC-GFP) displayed a strong localization along the MT lattice but failed to rescue the abnormal MT phenotype (Fig. 6G, 7; Movie 12, Fig. S3H). Taken together, none of the three C-terminal domains seemed to be individually essential for MT stabilization, apparently contrasting with the results obtained with the shot<sup>ΔEGC</sup> mutant embryos. A possible explanation for this phenomenon could be that, at the leading edge, the Gas2 and C-tail domains of Shot mutually substitute for each other in MT regulation, suggesting that these domains have redundant functions in DME cells. In summary, we conclude that the MT-binding activity of Shot is required but is not sufficient for MT stabilization.

**Actin- and MT-binding domains of Shot organize the epithelial MT network**

Therefore, to identify additional domains involved in MT stabilization, we tested the requirement for Shot’s actin-binding activity by using shot<sup>ΔC-actin</sup>, an isoform-specific mutant allele. In shot<sup>ΔC-actin</sup> mutant DME cells, abnormally long and bent MTs were found, revealing that the actin-binding activity of Shot is required for MT stabilization (Fig. 5C). To further investigate the role of the actin-binding activity of Shot, Shot<sub>ΔC-H1</sub>–GFP, a Shot isoform lacking the CH1 domain and therefore actin-binding ability, was expressed in DME cells. The truncated protein lost its association with actin and localized intensely along the MTs in the cell body, indicating that the CH1 domain exerts an inhibitory effect on the ability of Shot to bind MTs (Fig. 6H; Movie 13). MTs decorated with Shot<sub>ΔC-H1</sub>–GFP were abnormally bundled and curved, raising the possibility that Gal4-driven overexpression of the truncated Shot<sub>ΔC-H1</sub>–GFP protein has a dominant gain-of-function effect on MT organization. No abnormally protruding MTs were observed at the leading edge of shot<sup>Δ20</sup> null mutant in DME cells expressing Shot<sub>ΔC-H1</sub>–GFP (Fig. 7; Fig. S3I). The results obtained with the transgenic Shot<sub>ΔC-H1</sub>–GFP protein contrast with our previous observations on shot<sup>ΔC-actin</sup> mutants, which display abnormal MTs...
protruding from the leading edge. The apparent capacity of the transgenic protein to rescue the phenotypes could arise from an artificial dominant effect caused by the overexpression of $\text{Shot}^{\Delta \text{CH1}}$–GFP. Consistent with this idea, neuronal overexpression or a C-terminal truncation of Shot has been shown to dominantly cause alteration in MT network organization (Lee et al., 2016; Sánchez-Soriano et al., 2010). In DME cells, upon overexpression of $\text{Shot}^{\Delta \text{CH1}}$–GFP, the truncated protein inhibits the formation of protruding MTs at the leading edge; however, it does not restore wild-type MT organization. Taken together, based on the $\text{shot}^{\Delta \text{ECG}}$ mutant phenotype, we conclude that the actin-binding activity of Shot is required for MT stabilization.

The involvement of the actin- and MT-binding activities of Shot was further investigated by analyzing the $\text{Shot}^{\Delta \text{CH1}\Delta \text{Gas2}}$–GFP variant lacking both the CH1 and the Gas2 domains. $\text{Shot}^{\Delta \text{CH1}\Delta \text{Gas2}}$–GFP localized along the MT lattice but it was not capable of rescuing MT organization defects at the leading edge (Figs 6I, 7; Movie 14, Fig. S3J). Lack of rescue indicates that the actin-binding and the Gas2-mediated MT-binding activities of Shot are required for proper MT organization in the epithelial cells. Furthermore, these activities have to be simultaneously present in the same Shot molecule, as indicated by the abnormal MTs found at the leading edge of DME cells in $\text{shot}^{\Delta \text{ECG}}$/$\text{shot}^{\Delta \text{kakP1}}$ embryos (Fig. 5D). In summary, Shot functions as an actin–MT crosslinker to ensure proper MT regulation at the leading edge of DME cells.

**Shot promotes filopodia formation at the leading edge of the epithelial cells**

Our previous experiments revealed that both the actin and the MT regulatory activities of Shot function in DME cells are required to ensure proper MT organization and dorsal closure. Restoration of wild-type MT organization in $\text{shot}$ mutants, however, was not sufficient for proper zippering, suggesting that an additional activity of Shot might be required for dorsal closure. Therefore, we investigated the actin network of DME cells by analyzing actin accumulation and protrusion formation at their leading edge. In wild-type embryos, actin cables accumulate in epithelial cells at the leading edges. In fixed $\text{shot}^{\Delta \text{ECG}}$ null mutant and $\text{shot}^{\Delta \text{kakP1}}$ isoform specific mutant epithelia, normal actin accumulation was detected by phalloidin labeling (Figs 3B, 5C). In addition to accumulating actin, DME cells extend dynamic actin-based cell protrusions, such as lamellipodia and filopodia, at their leading edges. To identify the activity of Shot required for protrusion-mediated zippering, we analyzed the protrusion dynamics of DME cells by live imaging of $\text{shot}^{\Delta \text{ECG}}$ null mutants. Under the control of an en-Gal4 driver, mutant embryos expressed an actin–EGFP fusion protein, which specifically labeled the actin structures of the
epithelial cells (Fig. 8A; Movie 15). In shot^{G20} embryos, in addition to normal looking filopodia and lamellipodia, we frequently observed long protrusions extending even 10 μm above the amnioserosa (Fig. 8A; Movie 15). These protrusions were faintly decorated with actin–EGFP and might correspond to the cell extensions containing abnormally long and bent MTs. Dynamic parameters such as length and the number of filopodia were measured. Although the filopodia length of shot^{G20} null mutant DME cells was normal, the number of filopodia was reduced, indicating that Shot is involved in regulating filopodia formation (Fig. 8B,C). This phenotype was a direct consequence of the loss of the actin- and MT-binding activities in Shot, as indicated by the reduced number of filopodia in shot^{ECGC} and shot^{kakP1} mutant cells (Fig. 8A,C). Furthermore, and supporting a requirement for crosslinking activity of Shot, the MT- and actin-binding domains have to be simultaneously present in the same Shot molecule as indicated by the reduced number of filopodia found at the leading edge of DME cells in shot^{ECGC}/shot^{kakP1} transheterozygous embryos (Fig. 8A,C).

**DISCUSSION**

Shot is a key structural component of the cytoskeleton and has been demonstrated to be involved in several regulatory processes leading to rapid changes in cellular morphology. Here, we show that the cooperative functioning of actin and MTs is essential for efficient dorsal closure, and their activities are coordinated by Shot. We describe two functions of Shot in organizing the MT network of DME cells and in regulating zippering by promoting filopodia formation.

**Shot regulates MT organization in DME cells**

Based on their differential distribution and stability, we discriminate three subsets of MTs in DME cells. MTs of the first subset contain acetylated tubulin, a post-translational modification commonly
associated with more stable MTs, which turn over slowly (Asthana et al., 2013; Matov et al., 2010; Matsuyama et al., 2002; Tran et al., 2007; Webster and Borisy, 1989; Zilberman et al., 2009). In DME cells, these MTs may correspond to the immobile fraction of tubulin found in FRAP experiments. The low immobile fraction indicates that only a small fraction of the MTs (~one in five) belong to the stable MT subset. Acetylated MTs are concentrated at the apical surface of the cell body and are organized into parallel bundles. This MT arrangement is specific for the closure stage and seems to be independent of Shot activity, as indicated by the proper distribution of acetylated MTs in shot mutants. Furthermore, the immobile fraction of tubulin is not influenced by the shot null mutation. Thus, although Shot has been reported to influence MT stability in several cellular contexts, MT stabilization reflected by tubulin acetylation is insensitive to shot activity in DME cells.

MTs of second subset are the dynamic MTs found in the cell body. These MTs are aligned in an antiparallel manner in stable bundles and grow towards either the dorsal or the ventral periphery of DME cells. A possible role for Shot could be the guidance of MT growth along existing cytoskeletal filaments in a predefined pattern. Indeed, in vitro, parallel actin arrays can globally organize MT growth in a spectraplakin-dependent manner resulting in a parallel alignment of MTs (Peciaco López et al., 2014). In neurons and in vitro assays, the MT guidance activity of Shot requires C-tail-mediated plus-end-binding and MtLS-motif–EB1 interaction (Alves-Silva et al., 2012; Peciaco López et al., 2014). In DME cells, however, these interactions are dispensable for the organization of proper MT architecture, indicating that Shot does not play a prominent role in MT guidance in this cellular context. This conclusion is supported by the wild-type growth tracks observed by direct visualization of growing plus-tips in shot mutant DME cells. Thus, we hypothesize that additional, motor-driven guidance mechanisms are responsible for the establishment of the antiparallel arrangement of dynamic MTs (Akhmanova and Steinmetz, 2015; Chen et al., 2014; Doodhi et al., 2014; Mattie et al., 2010). Therefore, dynamic MTs in DME cells grow along each other or along acetylated stable MTs.

In the cell body of shot mutant DME cells, a slight increase in MT growth rate was detected indicating that Shot inhibits MT polymerization. This function of Shot is reflected by the faster turnover of tubulin measured by FRAP in shot mutant DME cells. A similar role for spectraplakins in MT growth speed regulation has been demonstrated in cultured Drosophila primary neurons and in human U2OS cells (Alves-Silva et al., 2012; Nishimura et al., 2012). Upon reaching the cell cortex, fast growing MTs of shot mutant DME cells continue to grow and either push out the cell cortex or bend along the lattice. The abnormal bending of the MTs could be responsible for the disorganized appearance of the MT network in the cell body of the shot mutant DME cells. Thus, Shot-mediated regulation of plus-end dynamics at the cell cortex contributes to the global organization of the MT network in the cell body.

MTs of the third subset reach the leading edge with their growing plus-ends and grow into protrusions. This fraction of MTs is severely affected by shot mutations: MTs are abnormally long and are frequently bent or curled. Similar MTs have been observed upon depletion of shot or ACF7 at the cell cortex in many other cell types, such as mammalian keratinocytes, endodermal cells, Drosophila S2 cells and neurons (Applewhite et al., 2010; Kodama et al., 2003; Sanchez-Soriano et al., 2009; Wu et al., 2008). Here, we show that, in addition to the Gas2-mediated MT association, Shot has to bind actin filaments simultaneously to regulate MT behavior. This observation is consistent with the hypothesis that bending of MTs occurs when they are not captured in the cortical actin network (Applewhite et al., 2010, 2013; Gierke and Wittmann, 2012). For cross-linking-mediated stabilization of MTs at the DME cell periphery, Shot associates with the MT lattice via its Gas2 domain and with cortical actin through its CH1 domain. Cortical tethering of MTs enables spatial control of MT dynamics by localized modification of plus-tip components, which promotes further cytoskeletal rearrangements required for the dorsal closure process. It remains to be determined what these cytoskeletal mechanisms are, but possible processes are protrusion formation or regulation of cell adhesion dynamics.

**Shot promotes zippering by regulating protrusion formation**

Depending on domain composition and cellular context, spectraplakin isoforms are able to regulate actin and MT networks separately or coordinate their interactions by simultaneously binding to both filament systems. As distinct steps of dorsal closure involve both actin- and MT-based mechanisms, there are many potential points for Shot to regulate the closure process.

Dorsal-ward movement of the epithelial sheets is mainly driven by actomyosin contraction-generated forces in the amnioserosa (Ducuing and Vincent, 2016; Pasakarnis et al., 2016). Dorsal-ward displacement of the epithelial fronts is unaffected in shot mutants, indicating that shot is not required for the regulation of actin-based processes at this stage of the closure.

The last sealing step of the closure, zippering of the hole, however, requires shot function. Two major cellular mechanisms have been shown to facilitate closure at this stage: formation of protrusions that establish initial contacts between the opposing DME cells at the onset of zippering and the resolution of overlapping lamellar regions between them after they have met (Eltsov et al., 2015). Our data suggest that Shot regulates zippering by promoting protrusion formation rather than by supporting rearrangement of the lamellar interaction surfaces between DME cells. Resolution of lamellar overlaps has been suggested to be driven by microtubule-generated forces (Eltsov et al., 2015). However, our rescue experiments demonstrated that in addition to its function in MT regulation, the actin-binding activity of Shot is also required for zippering. While we cannot exclude the direct involvement of the MT regulatory function of Shot in overlap rearrangement, Shot-mediated stabilization of MTs alone is clearly not sufficient to ensure proper zippering.

In DME cells, proper protrusion dynamics requires both actin and MT activities (Eltsov et al., 2015; Hakeda-Suzuki et al., 2002; Harden et al., 1999; Jankovics and Brunner, 2006; Woolner et al., 2005). Here, we show that the coordinated action of the two filament types is required for proper filopodia function, and this coordination is orchestrated by Shot. In mediating zippering, the actin- and MT-binding activities of Shot are simultaneously required in the same molecule, indicating that Shot acts as a MT–actin crosslinker in this process. In DME cells, close alignment of MTs and actin filaments has been found exclusively in filopodia, thus, Shot exerts its crosslinking function in cell protrusions. This conclusion is supported by the reduction of the number of protrusions in shot mutants. We hypothesize that lack of actin-MT crosslinking causes abnormal protrusion formation in shot mutants, which in turn leads to inefficient zippering.

How Shot contributes to protrusion dynamics in DME cells remains elusive. In neuronal growth cones, the interaction of EF-hand domain of Shot with Kra (also known as eIF5C) has been
described to be essential for filopodia formation (Sanchez-Soriano et al., 2009). In DME cells, however, the EF-hand domain is dispensable for dorsal closure, but the actin- and MT-binding functions of Shot are required for filopodia-mediated zipper. The differences in domain requirement indicate that Shot can promote protrusion dynamics by distinct mechanisms in various cell types. In cultured mammalian cells, interaction of the Shot homolog ACM with ELMO1 (also known as DOCK180) has been reported to promote protrusion formation by coordination of localized MT stabilization and Rac GTPase activity (Margaron et al., 2013). Possibly, a similar mechanism works in DME cells in which the interaction of Shot with the ELMO complex (ELMO is known as Mbc in Drosophila) stabilizes MTs by crosslinking them with the actin network and targets Rac activation to the leading edge. In support of this hypothesis, Rac activation has been shown to be required for protrusion formation and efficient zipperping (Hakeda-Suzuki et al., 2002; Woolner et al., 2005).

The effect of the complete lack of Shot function during dorsal closure is very subtle, which can be explained by two, not mutually exclusive, hypotheses. First, that many additional crosslinkers or crosslinking mechanisms may act in parallel with Shot. A possible redundant cytoskeletal linker might be Pigs, which has been demonstrated to have CH1, Gas2 and C-tail domains by which it binds both actin and MTs (Girdler et al., 2016). Second, that the function of Shot is restricted to supporting protrusion formation in DME cells. Inefficient protrusion formation in shot mutants does not abolish zipperping but reduces its efficiency (Eltsov et al., 2015; Jankovics and Brunner, 2006).

**MATERIALS AND METHODS**

*Drosophila* stocks

OregonR was used as wild type. The ZASP2**Δ**C**Δ**L**Δ**3 protein trap line was used to visualize the leading edge of the DME cells (Morin et al., 2001; Stringer, 2014). *Pnr-GAL4* and *en-GAL4* were used for inducible expression of the selected genes. For rescue experiments, the following transgenic constructs were used: Shot-L(A)–GFP, ΔEFhand–GFP [Shot-L (A)ΔEFHand–GFP], ΔGas2–GFP [Shot-L(A)ΔGas2–GFP], ΔCH1–GFP [Shot-L(C)–GFP]. These overexpression constructs were generated based on the *shot* mRNA as a transcript *shot-KE* in FlyBase (Lee and Kolodziej, 2002b). The encoded Shot protein contains all protein domains (CH1, CH2, plakin family domain, spectrin repeats, EF, Gas2 and the C-tail) apart from the plakin repeat region that is encoded by a single large exon and not contained within this mRNA (illustrated in Fig. 2C) (Röper and Brown, 2003; Röper et al., 2002). In addition, the following domain-deletion or single-domain transgenes derived from Shot-L(A)–GFP were used: ΔCH1ΔGas2–GFP [Shot-L(C)Gas2–GFP] (Lee and Kolodziej, 2002a), ΔC-tail–GFP [Shot-L(A)–Ctail–GFP], C-tail–GFP, Shot-L(A)–3Mls–GFP (Alves-Silva et al., 2012), EGC–GFP (Shot-EFhand-Gas2-Ctail–GFP) (Subramanian et al., 2003).

For loss-of-function analyses, we used *shot**Δ**D*, *shot**Δ**API* and *shot**Δ**EGC* alleles and the *Df(2R)BSC383* deficiency covering the *shot* locus.

Actin was visualized with UAS-actin–EGFP (Inoue et al., 2004). FRAP experiments were performed with a Leica SP5 confocal microscope. A 405 nm laser was used to photobleach a 2 μm-wide stripe at the leading edge of the DME cells. Recovery after photobleaching was recorded for three minutes at 1 frame every 2 s. DME cells moving out of focus during image acquisition were ignored. Fluorescence recovery curves were analyzed using the easyFRAP software as described in Bancaud et al. (2010). Statistical tests were performed with GraphPad Prism.

**Protrusion and MT dynamics measurements**

For time-lapse imaging of protrusions, *shot**Δ**D*, *shot**Δ**AP1*, and *Df(2R)BSC383*/*shot**Δ**EGC*, and *shot**Δ**API*/*shot**Δ**EGC* embryos were used. The number of filopodia extended in 15 min was calculated from measurements over more than 30 min. For measuring the MT growth rate and growth direction, *en-Gal4/EB1–EGFP* and *en-Gal4, Df(2R)BSC383/shot**Δ**D*, UAS-EB1–EGFP embryos were imaged with an Olympus CellR microscope. Quantitative parameters were measured using ImageJ and analyzed with GraphPad Prism.

**Immunohistochemistry**

Immunostainings were performed as described earlier (Jankovics and Brunner, 2006). Mouse-anti-Tubulin (1:10, E7, DSHB), rabbit-anti-EGFP (1:500, Life Technologies), mouse-anti-acetylated-tubulin (1:1000, 6-11B-1, Sigma), mouse-anti-FasciclinIII (1:20, 7G10, DSHB) and guinea pig anti-Shot-spectrin-repeats (1:1000) primary antibodies were used. To label actin, manually deventilated embryos were incubated for one hour in Rhodamine-phallolidin (Molecular Probes). Specimens were examined with Leica TCS SP5 confocal microscope.

Acknowledgements

We thank A. Prokop and T. Millard for fly stocks and reagents. We also thank G. Tick, and B. Irvine for critical reading of the manuscript.

Competing interests

The authors declare no competing or financial interests.

**Author contributions**

F.J. and M.E. conceived and designed the experiments. Z.T. and F.J. performed the experiments. Z.T. and F.J. analyzed the data. P.L. and K.R. contributed reagents and analysis tools. F.J., K.R. and M.E. wrote the paper.

**Funding**

This work was supported by the János Bolyai Research Fellowship of the Magyar Tudományos Akadémia (Hungarian Academy of Sciences) and by the National Research, Development and Innovation Office (NKFI-K117010, OTKA-K108538 and GINOP-2.3.2-15-2016-00001). K.R. was funded by the Medical Research Council (MC_UP_120111). Deposited in PMC for immediate release.

Supplementary information

Supplementary information available online at http://jcs.biologists.org/lookup/doi/10.1242/jcs.193003.supplemental
References

Akmanova, A. and Steinmetz, M. O. (2015). Control of microtubule organization and dynamics: two ends in the limelight. *Nat. Rev. Mol. Cell Biol.* 16, 711-726.

Alves-Silva, J., Sánchez-Soriano, N., Beaven, R., Klein, M., Parkin, J., Millard, T. H., Bellen, H. J., Venken, K. J. T., Ballesteros, C., Kammerer, R. A. et al. (2012). Spectrapsin promotes microtubule-mediated axonal growth by functioning as structural actin-microtubule-associated proteins and EB1-dependent +TIPs (tip interacting proteins). *J. Neurosci.* 32, 9143-9158.

Applewhite, D. A., Grode, K. D., Duncan, M. C. and Rogers, S. L. (2015). Coordinating neuronal actin organization and dynamics: two ends in the limelight. *Nat. Rev. Mol. Cell Biol.* 21, 1714-1724.

Bottenberg, W., Sanchez-Soriano, N., Alves-Silva, J., Hahn, I., Mende, M. and Ducuing, A. and Vincent, S. (2014). An EB1-kinesin complex is sufficient to steer microtubule growth in vitro. *Curr. Biol.* 24, 316-321.

Bottenberg, W., Sanchez-Soriano, N., Alves-Silva, J., Hahn, I., Mende, M. and Prokop, A. (2009). Context-specific requirements of functional domains of the spectraplakin short stop in vivo. *Mech. Dev.* 126, 499-505.

Bottenberg, W., Sanchez-Soriano, N., Alves-Silva, J., Hahn, I., Mende, M. and Prokop, A. (2009). Context-specific requirements of functional domains of the spectraplakin short stop in vivo. *Mech. Dev.* 126, 499-505.

Bottenberg, W., Sanchez-Soriano, N., Alves-Silva, J., Hahn, I., Mende, M. and Prokop, A. (2009). Context-specific requirements of functional domains of the spectraplakin short stop in vivo. *Mech. Dev.* 126, 499-505.

Chen, Y., Rolls, M. M. and Hancock, W. O. (2014). An EB1-kinesin complex is sufficient to steer microtubule growth in vitro. *Curr. Biol.* 24, 316-321.

Coles, C. H. and Bradke, F. (2015). Coordinating neuronal actin–microtubule dynamics. *Curr. Biol.* 25, R577-R91.

Doddhi, H., Katrukha, A. A., Kaptein, L. C. and Akhamanova, A. (2014). Mechanical and geometrical constraints control kinesin-based microtubule guidance. *Curr. Biol.* 24, 322-328.

Ducluing, A. and Vincent, S. (2016). The actin cable is dispensable in directing dorsal closure dynamics but neutralizes mechanical stress to prevent scarring in the Drosophila embryo. *Nat. Cell Biol.* 18, 1149-1160.

Eltsos, M., Dubé, N. Y., Zu, Y., Pasakarnis, L., Haselmann-Weiss, U., Brunner, D. and Franagakis, A. S. (2015). Quantitative analysis of cytoskeletal reorganization during epithelial tissue sealing by large-volume electron tomography. *Nat. Cell Biol.* 17, 605-614.

Felgner, T. A. and Rørup, K. (2002). Invasive cell migration is initiated by guided growth of long cellular extensions. *Nat. Cell Biol.* 4, 715-719.

Gierke, S. and Wittmann, T. (2012). EB1-recruited microtubule +TIP complexes coordinate protrusion dynamics during 3D epithelial remodeling. *Curr. Biol.* 22, 753-762.

Girdler, G. C., Applewhite, D. A., Perry, W. M. G., Rogers, S. L. and Rørup, K. (2016). The Gas2 family protein Pgs is a microtubule +TIP that affects cytoskeletal organization. *J. Cell Sci.* 129, 121-134.

Gratz, S. J., Cummings, A. M., Nguyen, J. N., Hamm, D. C., Donohue, L. K., Harrison, M. W., Wildonger, J. and O’Connor-Giles, K. M. (2013). Genome engineering of Drosophila with the CRISPR RNA-guided Cas9 nuclease. *gene. *in vivo destabilization of dynamic microtubules by HDAC6-mediated deacetylation. *EMBO J.* 21, 6820-6831.

Matsuyama, A., Shimazu, T., Sumida, Y., Saito, A., Yoshimatsu, Y., Segeier-Beaver, R., Nakada, Y., Komatsu, Y., Nishino, N., Khochbin, S. et al. (2002). Functional genomic screen combined with time-lapse microscopy uncovers a novel set of genes involved in dorsal closure of Drosophila embryos. *PLoS ONE* 1, e22229.

Matov, A., Applegate, K., Kumar, P., Thoma, C., Krek, W., Danuser, G. and Wittmann, T. (2010). Analysis of microtubule dynamic instability using a plus-end growing marker. *Nat. Methods* 7, 761-768.

Matsuura, Y., Shimazu, T., Sumida, Y., Saito, A., Yoshimatsu, Y., Segeier-Beaver, R., Nakada, Y., Komatsu, Y., Nishino, N., Khochbin, S. et al. (2002). Functional genomic screen combined with time-lapse microscopy uncovers a novel set of genes involved in dorsal closure of Drosophila embryos. *PLoS ONE* 1, e22229.

Matov, A., Applegate, K., Kumar, P., Thoma, C., Krek, W., Danuser, G. and Wittmann, T. (2010). Analysis of microtubule dynamic instability using a plus-end growing marker. *Nat. Methods* 7, 761-768.

Millard, T. H. and Martin, P. (2008). Dynamic analysis of filopodial interactions during the zipper phase of Drosophila dorsal closure. *Development* 135, 621-626.

Morin, X., Daneman, R., Zavortink, M. and Chia, W. (2001). A protein trap strategy to detect GFP-tagged proteins expressed from their endogenous loci in Drosophila. *Proc. Natl. Acad. Sci. USA* 98, 15050-15055.

Morphism, Y., Applegate, K., Davidson, M. W., Danuser, G. and Waterman, C. M. (2012). Automated screening of microtubule growth dynamics identifies MARK2 as a regulator of leading edge microtubules downstream of Rac1 in migrating cells. *PLoS ONE* 7, e41413.

Nishimura, Y., Applegate, K., Davidson, M. W., Danuser, G. and Waterman, C. M. (2012). Automated screening of microtubule growth dynamics identifies MARK2 as a regulator of leading edge microtubules downstream of Rac1 in migrating cells. *PLoS ONE* 7, e41413.

Pasakarnis, L., Frei, E., Caussinus, E., Affolter, M. and Brunner, D. (2016). Aminoferre cell constriction but not epidural actin cable tension autonomously drives dorsal closure. *Nat. Cell Biol.* 18, 1161-1172.

Pecloado Lópe, M., Huber, F., Gregor, I., Simonet, M. O., Akhmanova, A., Wolf, D., Golembo, L. and Donohue, L. K. (2013). Actin–microtubule coordination at growing microtubule ends. *Nat. Commun.* 4, 7578.

Røper, K. and Brown, N. H. (2003). Maintaining epithelial integrity: a function for gigantic spectraplakin isoforms in adherens junctions. *J. Cell Biol.* 162, 1305-1315.

Røper, K., Gregory, S. L. and Brown, N. H. (2002). The “spectraplakins”: cytoskeletal giants with characteristics of both spectrin and plakin families. *J. Cell Sci.* 115, 4215-4225.

Sanchez-Soriano, N., Travis, M., Dajas-Bailador, F., Gonçalves-Pimentel, C., Whitmarsh, A. J. and Prokop, A. (2009). Mouse ACF7 and drosophila short stop modulate filopodia formation and microtubule organisation during neuronal growth. *J. Cell Sci.* 122, 2534-2542.

Sanchez-Soriano, N., Gonçalves-Pimentel, C., Beaven, R., Haessler, U., Ofner-Ziegenfuss, L., Ballesteros, C. and Prokop, A. (2010). Drosophila growth cones: a genetically tractable model for the analysis of axonal growth dynamics. *Dev. Neurobiol.* 70, 58-71.

Sebo, Z. L., Lee, H. B., Peng, Y. and Guo, Y. (2014). A simplified and efficient germline-specific CRISPR/Cas9 system for Drosophila genome engineering. *Fly* 8, 52-57.

Stronach, B. (2014). Extensive nonmuscle expression and epithelial apicalbosal localization of the Drosophila ALP/Enigma family protein, Zasp52. *Gene Expr. Patterns* 15, 67-79.

Subramanian, A., Prokop, A., Yamamoto, M., Sugimura, K., Uemura, T., Betschinger, J., Knoblich, J. A. and Volk, T. (2003). Shortstop recruits EB1/ APC1 and promotes microtubule assembly at the muscle-tendon junction. *Curr. Biol.* 13, 3021-3034.

Suozzi, K. C., Wu, X. and Fuchs, E. (2012). Spectraplakins: master orchestrators of cytoskeletal reorganization. *J. Cell Biol.* 197, 465-475.
Tran, A. D.-A., Marmo, T. P., Salam, A. A., Che, S., Finkelstein, E., Kabarriti, R., Xenias, H. S., Mazitschek, R., Hubbert, C., Kawaguchi, Y. et al. (2007). HDAC6 deacetylation of tubulin modulates dynamics of cellular adhesions. *J. Cell Sci.* 120, 1469-1479.

Webster, D. R. and Borisy, G. G. (1989). Microtubules are acetylated in domains that turn over slowly. *J. Cell Sci.* 92, 57-65.

Wolf, N., Regan, C. L. and Fuller, M. T. (1988). Temporal and spatial pattern of differences in microtubule behaviour during Drosophila embryogenesis revealed by distribution of a tubulin isoform. *Development* 102, 311-324.

Woolner, S., Jacinto, A. and Martin, P. (2005). The small GTPase Rac plays multiple roles in epithelial sheet fusion—dynamic studies of Drosophila dorsal closure. *Dev. Biol.* 282, 163-173.

Wu, X., Kodama, A. and Fuchs, E. (2008). ACF7 regulates cytoskeletal-focal adhesion dynamics and migration and has ATPase activity. *Cell* 135, 137-148.

Zilberman, Y., Ballestrem, C., Carramusa, L., Mazitschek, R., Khochbin, S. and Bershadsky, A. (2009). Regulation of microtubule dynamics by inhibition of the tubulin deacetylase HDAC6. *J. Cell Sci.* 122, 3531-3541.