Drosophila Big bang regulates the apical cytocortex and wing growth through junctional tension

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Growth of epithelial tissues is regulated by a plethora of components, including signaling and scaffolding proteins, but also by junctional tension, mediated by the actomyosin cytoskeleton. However, how these players are spatially organized and functionally coordinated is not well understood. Here, we identify the Drosophila melanogaster scaffolding protein Big bang as a novel regulator of growth in epithelial cells of the wing disc by ensuring proper junctional tension. Loss of big bang results in the reduction of the regulatory light chain of nonmuscle myosin, Spaghetti squash. This is associated with an increased apical cell surface, decreased junctional tension, and smaller wings. Strikingly, these phenotypic traits of big bang mutant discs can be rescued by expressing constitutively active Spaghetti squash. Big bang colocalizes with Spaghetti squash in the apical cytocortex and is found in the same protein complex. These results suggest that in epithelial cells of developing wings, the scaffolding protein Big bang controls apical cytocortex organization, which is important for regulating cell shape and tissue growth.

Introduction

Epithelial tissue morphogenesis and growth are regulated by a plethora of mechanisms and components, including the actomyosin cytoskeleton, polarity regulators, various signaling pathways, systemic cues, and cell–cell and cell–matrix contacts (Zhang et al., 2010; Lye and Sanson, 2011; Röper, 2015). Many of the participating components are organized as multiprotein complexes in the apex of the cell, such as adhesion or signaling complexes, and are instrumental in regulating cell and tissue behavior—for example, cell size, cell division and shape, and tissue growth and folding. Signals can modulate actomyosin activity, thereby inducing morphogenetic changes. On the other hand, there is increasing evidence that mechanical forces originating from the actin cytoskeleton are essential regulators of tissue morphogenesis and growth by modulating signaling pathway activities (Lye and Sanson, 2011; Colombelli and Solon, 2013; Clark et al., 2014; Choi et al., 2016; LeGoff and Lecuit, 2016; Vásquez and Martin, 2016). Excess actin polymerization, for example, induced by various actin-binding proteins, can result in excess growth (Fernández et al., 2011; Sansores-Garcia et al., 2011; Yu and Guan, 2013; Gaspar and Tapon, 2014; Rauskolb et al., 2014; Deng et al., 2015; Sun and Irvine, 2016). How tension is sensed and how it is converted into chemical signaling to modify gene expression and ultimately cell behavior is still poorly understood. So far, no general concept has emerged, which may also be a result of a variety of cell- and tissue-specific tension sensors and their cellular effectors. Among the known tension sensors involved in growth control are cytoskeletal components, e.g., Spectrin and actin (Sansores-Garcia et al., 2011; Deng et al., 2015; Fletcher et al., 2015; Gaspar et al., 2015), but also the junctional components α- and β-catenin and p120-catenin, which act either indirectly via other proteins or directly, by translocating into the nucleus (Spadaro et al., 2012; Rauskolb et al., 2014). These few examples underscore the important role of cytoskeleton–junction-mediated tension in growth control, but at the same time they unveil the complexity of growth regulation by tension. Among the effectors are signaling pathways, such as ECM-mediated signaling or the Hippo pathway, which are conserved from flies to mammals (Ingber, 2006; Badouel et al., 2009; Halder et al., 2012; Dupont, 2016; Sun and Irvine, 2016).

These results also indicate that we are far from a complete picture of how tissue tension controls growth. Given that adherens junctions, a major site of tension modulation, reside apically in epithelial cells, and that many of the regulatory and signaling molecules localize apically as well, one important question remains, namely, which components help to organize the apical cytocortex itself. Solving this question is crucial to understand how the different factors involved are coordinated and how they impact junctional tension. To identify these components, we conducted a genetic modifier screen aimed to find novel regulators of wing growth (Nemetschke and Knust, 2016). One of the modifiers turned out to be big bang (bbg). bbg encodes a scaffolding protein with three PSD-95/Discs large/ZO-1 (PDZ) domains, which has previously been shown to regulate...
Results

\textbf{bbg regulates wing growth during \textit{Drosophila} development}

The \textit{Drosophila} wing imaginal discs have turned out as an ideal model in which to study the genetic, molecular, and cell biological basis of various aspects of tissue morphogenesis and growth. To identify novel regulators of wing growth, we performed a genetic screen by scoring for mutations that dominantly modulate changes and has been described to control border cell migration in \textit{Drosophila} (Nemetschke and Knust, 2016). One of the enhancers identified in this screen was \textit{bbg}, \textit{bbg} encodes a scaffolding protein with three PDZ domains and has been described to control border cell migration in the follicle (Kim et al., 2006) and to modulate the gut immune tolerance (Bonmay et al., 2013). To determine whether \textit{bbg} controls wing size on its own, we knocked down \textit{bbg} activity in developing wings. RNAi-mediated knockdown of \textit{bbg} by using two different Ga4 lines resulted in smaller wings (Fig. 1, A–F; quantified in Fig. 1 M). Reduction of \textit{Irhp}, a predicted off-target of \textit{bbg} RNAi (Aranjuez et al., 2012), did not show any growth defect in wings of adult flies (Fig. S1 A–F; quantified in Fig. S1 G), \textit{bbg} homozygous mutant flies, which are viable (Kim et al., 2006), as well as \textit{bbg}\textsuperscript{bbgB211}/\textit{Df(3L)4543} hemizygotes, develop even smaller wings (Fig. 1, G–L; quantified in Fig. 1 M).

The adult fly wing develops from the wing imaginal disc, an epithelial sac built from a single layered epithelium. Specified during embryogenesis, wing discs expand about a 1,000-fold through proliferation during larval stages. The wing blade originates from the central area of the disc, the pouch (Fig. 2 I, green). To analyze the role of \textit{bbg} in wing growth, we studied the proliferation behavior of \textit{bbg}\textsuperscript{bbgB211} homozygous cells by inducing \textit{bbg} mutant clones at two different developmental stages. To exclude any cell competition, GFP-positive \textit{bbg} mutant clones were studied in \textit{bbg} mutant discs. Their behavior was compared with that of GFP-positive WT clones induced in WT discs. The total clone area per wing pouch was determined in third instar larvae (L3) discs. GFP-positive \textit{bbg} mutant clones in \textit{bbg} mutant discs were ∼50% or 70% smaller than GFP-positive WT clones in WT discs when induced at 48 h or 72 h after egg laying (AEL), respectively (Fig. 2, A–F; quantified in Fig. 2 E). In addition, the number of GFP-labeled \textit{bbg} mutant clones was reduced by ∼50% compared with the number of WT clones when induced 48 h AEL, and ∼60% less mutant clones were observed upon induction at 72 h AEL (Fig. 2 F). These results indicate that \textit{bbg} is required for normal wing growth in \textit{Drosophila}.

To further determine whether the smaller wings of flies lacking \textit{bbg} were a result of cell cycle arrest, we identified the cell cycle stages in WT and \textit{bbg} mutant wing disc cells by FACS analysis. Notably, we compared exactly the same number of events both in WT and \textit{bbg} mutants. The two different peaks shown in the histogram (Fig. 2 G) allowed us to distinguish the G0/1 and G2 phases (black arrows in Fig. 2 G). In the absence of \textit{bbg}, the number of cells in G2 are reduced by ∼19% and those in G0/G1 are increased by ∼19% compared with the corresponding numbers of WT cells (Fig. 2 G). From this we conclude that loss of \textit{bbg} perturbs cell cycle progression, because there are fewer cells in G2 and more cells in G1/G0. To better understand the basis of the perturbed cell cycle, we determined cell number, cell division, and apoptosis in wing discs of L3 larvae. The analysis was restricted to the center of the wing pouch (red rectangle in Fig. 2 H). Cell borders were marked by an antibody against Discs large (Dlg). L3 wing pouches of animals expressing \textit{bbg}\textsuperscript{RNAi} or of \textit{bbg} homozygous mutant animals exhibited 20% and 35% fewer cells, respectively, in comparison to control animals (Fig. 2, J–L; quantified in Fig. 2 M).

Cell numbers and hence wing size can be regulated through cell divisions or cell death or a combination of these, and many genes have been identified that regulate this process (Hariharan, 2015). To study the effect of \textit{bbg} on proliferation, the number of mitotic cells in the whole pouch area (Fig. 2 I, green) was counted, using an antibody that detects mitosis-specific phosphorylation of histone H3 (PH3). Compared with WT control animals, the number of mitotic cells was reduced by 22% in wing pouches of L3 larvae expressing \textit{bbg} RNAi and by 29% in \textit{bbg} homozygous mutant animals (Fig. 2, J', K', and L'; quantified in Fig. 2 N). This result, together with a comparable decrease in cell number in \textit{bbg} mutant discs (see Fig. 2 M) and an increase in the number of cells in G1/G0 (Fig. 2 G), pointed to an increase in apoptosis. To corroborate this assumption, apoptosis was analyzed by TUNEL assays in wing discs. In the wing pouch of WT L3 discs, the number of apoptotic cells was very low (Fig. 2 I), as reported previously (Milán et al., 1997). In contrast, the number of TUNEL-positive cells was significantly increased upon knockdown or loss of \textit{bbg} (Fig. 2, K' and L'; quantified in Fig. 2 O). This result is in agreement with the observation that fewer clones were observed in mutant discs when induced at 72 h AEL (Fig. 2 F). To exclude the possibility that \textit{bbg}\textsuperscript{RNAi} cells are development-
tally delayed, we expressed RNAi only in the posterior compartment using en-Gal4 and compared the anterior/posterior ratio of PH3- and TUNEL-positive cells. The results support the previous data, in that the posterior compartment with reduced level of Bbg reveals less proliferating, PH3-positive cells and more apoptotic cells (Fig. S2, A–F). Collectively, the reduced wing-size in bbgB211 mutants can be attributed, at least partially, to increased apoptosis.

Bbg localizes in the apical cytocortex and regulates growth through Sqh

To further elucidate the mechanism by which bbg exerts its function, we raised antibodies that should detect all predicted Bbg isoforms. The specificity of the antibody was confirmed by Western blot (WB) analysis (Fig. S3 A) and immunohistochemical staining of wing discs, in which bbg was reduced in the posterior compartment by RNAi-mediated knockdown (Fig. S3, B–B”). In wing disc cells of third instar WT larvae, Bbg localized in the apical cytocortex (Fig. 3, A–A”), apical to Dlg (Fig. 3, B–B”). Here, Bbg colocalized with Sqh-GFP (Fig. 3, C–C”), the Drosophila regulatory light chain of nonmuscle myosin II (Karess et al., 1991). Sqh has been reported to be enriched in the apical cytocortex (Landsberg et al., 2009) and is required, together with the myosin II heavy chain, encoded by zipper (zip), for many morphogenetic processes, including imaginal discs morphogenesis (Edwards and Kiehart, 1996; Aldaz et al., 2013). In the wing disc, Bbg was enriched in the cell cortex along the anterior-posterior (AP) compartment boundary (Fig. S3, C–C”, white arrows), thus reproducing the pattern of actin

Figure 1. Loss of bbg results in smaller wings. (A–C) Control (69B-Gal4) wing (A), wing expressing UAS-bbgRNAi with 69B-Gal4 (B), and overlay (C). (D–F) Control (C765-Gal4) wing (D), wing expressing UAS-bbgRNAi with C765-Gal4 (E), and overlay (F). (G–I) bbgB211/– heterozygous wing (G), bbgB211/Df(3L)ED4543 wing (K), and overlay (L). (M) Wing size measurement of the surface area of 50 (A–C and G–I) and 15 (D–F and J–L) independent females per genotype. The statistical analysis (M) used t test and ANOVA. *, P ≤ 0.1; ***, P ≤ 0.001. G and J show the same wing, because all figures depicted here were obtained in the same experiment. Error bars show SD. Bar, 500 µm.
Figure 2. **Loss of bbg results in fewer cells and increased apoptosis in L3 wing discs.**

(A–D) GFP expressing clones in WT (A and C) and bbgB211 mutant (B and D) induced at different time points (heat-shock [hs] 48 h and 72 h AEL) and stained with an anti-GFP. 

(E and F) Ratio of GFP-positive clones to the whole pouch (E), and number of GFP-positive clones in the pouch of WT and bbgB211 mutant L3 wing discs (F), induced by 48 and 72 h AEL, using 10 independent discs per genotype.

(G) FACS analysis from cells of ~20 L3 wing discs (10,000 events/cells per condition) of WT and bbgB211 mutant. Histograms display DNA content/fluorescent intensity (x-axis) and cell numbers (y-axis). Diagram: Mean of WT and bbgB211 mutant cells in every cell cycle stage (three biological replicates per condition).

(H and I) Cartoons representing the wing pouch (green), the area measured in J, K, and L (red box, H) and the outline of the pouch measured in J', K', L', and L'' (gray outline, I). Control (69B-Gal4; J–J’), 69B-bbgRNAi (K–K’), and bbgB211 mutant (L–L’). L3 wing discs stained with anti-Dig, anti-PH3, and TUNEL, respectively. Measurement of cell numbers (M), PH3-positive cells (N), and TUNEL-positive cells (O) in all three genotypes, respectively, using eight independent L3 wing discs per genotype. The statistical analysis (E–G and M–O) used t test and ANOVA.

* P ≤ 0.1; ** P ≤ 0.01; *** P ≤ 0.001. Error bars show SD. Bars, 25 µm.
and Sqh localization (Landsberg et al., 2009), and in the cortex of cells entering mitosis (Fig. S3 C′, magenta arrowheads).

To analyze a possible link between bbg and sqh, we looked at the role of sqh in WT and bbg mutant discs. Therefore, we asked whether Sqh itself regulates wing size. Knocking down sqh reduced wing size to a similar extent as knocking down bbg (Fig. 4, compare A–C with D–F; quantified in Fig. 4 M). Strikingly, concomitant knockdown of sqh and bbg in the whole wing resulted in nearly 100% lethality. To further unveil the relationship between bbg and sqh, we overexpressed Sqh E20E21, a variant in which the two regulatory phosphorylation sites, Thr-20 and Ser-21, are mutated to phosphomimetic Glu residues. This variant has been shown to act as a constitutive active form of Sqh (Winter et al., 2001). As previously shown (Rauskolb et al., 2014), lowering the activity of Rok also gives rise to smaller wings (Fig. 5, A–C and G). Therefore, we also tested the genetic interaction between bbg and rok, and found that concomitant knockdown of rok and bbg also resulted in lethality (Fig. 5 G), supporting the link between bbg and sqh. These results suggest that bbg acts upstream of sqh to control wing size.

**Bbg stabilizes Sqh in the apical cytocortex of wing disc cells**

Next, we set out to study the molecular relationship between Bbg and Sqh. To analyze the localization of Sqh, we used animals expressing Sqh-GFP under the endogenous promoter in a sqh mutant background. This transgene completely rescues all sqh mutant phenotypes (Royou et al., 2004). Upon reduction of Bbg, Sqh-GFP was more diffuse (Fig. 6, A–B′). In addition, the amount of Sqh-GFP, as measured by fluorescence intensity, was reduced by 30% upon reduction of bbg in the posterior compartment of the wing disc in comparison to the control, anterior compartment (Fig. 6 C). Furthermore, the phospho-tylated form of Sqh was reduced in the absence of Bbg as well.

Figure 3. Bbg localizes in the apical cytocortex of L3 wing disc epithelial cells. (A–B′) WT L3 wing discs stained with anti-Bbg (A and B), anti-DE-Cadherin (DE-Cad; A′), anti-Dlg (B′), and the respective overlays (A″ and B″). (C–C′) sqh-GFP L3 wing disc stained with anti-Bbg (C), Sqh-GFP [endogenous signal, C′] and the respective overlay [C″]. The projection in B was taken from a more lateral view compared with that of A and C. Insets, top right: Respective pouch areas. xz projection shows the central area of the same L3 wing discs. Bars: [A″, B″, and C″] 25 µm; [xz projections and small boxes] 5 µm.
The colocalization of the two proteins in the apical cortex let us to speculate that the two proteins may be part of the same complex. To address this question, we immunoprecipitated Sqh-GFP from protein extracts of wing disc lysates, using an anti-GFP antibody. Bbg was pulled down from discs expressing Sqh-GFP (Fig. 6 E, right two lanes), but not from control (WT) discs (Fig. 6 E, left two lanes). To conclude, Bbg can be found in a protein complex with Sqh in wing imaginal discs, where it stabilizes Sqh in the apical cytocortex.

Bbg is required to stabilize junctional tension in wing imaginal discs

We noticed that the apical surface of cells in bbg<sup>B211</sup> mutant L3 wing discs appeared larger in size in comparison to WT cells (Fig. 2, J and L). We confirmed this observation by knocking down bbg in the posterior compartment of L3 wing discs (Fig. 7, A–B'). Quantification of apical cell surface area in wing discs stained for DE-cadherin to outline the cell apex (Fig. 7, C and D) revealed a ∼23% increase of apical cell surface area

![Image](image-url)
in bbgΔ211 mutant cells compared with corresponding WT cells (Fig. 7 E). Similar to loss of bbg, RNAi-mediated reduction of sqh in the posterior compartment had no effect on DE-cadherin localization and tissue integrity (Fig. 7 F), and resulted in enlarged cell surface areas (Fig. 7, F–F′), supporting the previous conclusion that bbg and sqh act on a common pathway.

Because Sqh is a major regulator of actin, we asked whether absence of bbg affects actin localization. We knocked down bbg by expressing RNAi in the posterior compartment of wing discs and stained with phalloidin. Under these conditions, F-actin was reduced by ∼25% both apically and laterally in comparison to the anterior, bbg-positive tissue (Fig. 8, A–F′; quantified in Fig. 8 H). Reduction of F-actin upon knockdown of bbg could be prevented by simultaneous overexpression of SqhE20E21 (Fig. 8, G–G′′, quantified in Fig. 8 H).

Actin is a major regulator of tension, and tension has been shown to regulate growth (Mao et al., 2013; Schluck et al., 2013; LeGoff and Lecuit, 2016). To determine whether bbg controls tension in wing imaginal discs, we ablated single cell junctions by laser and quantified the initial velocity of the movement of vertices, which is a suitable readout for mechanical tension (Fig. 8 I; Landsberg et al., 2009). The velocity was reduced by 47% upon RNAi-mediated knockdown of bbg in the posterior compartment in comparison to the velocity in the anterior, bbg-positive control compartments (Fig. 8 J). A similar reduction in the initial velocity (52%) was observed upon knocking down sqh (Fig. 8 J). Strikingly, simultaneous expression of SqhE20E21 and bbg RNAi rescued the reduced velocity observed in bbg RNAi discs and brought it back to WT levels. To conclude, bbg controls junctional tension in wing imaginal discs by promoting the activity of Sqh.

Collectively, our data show that bbg is a key organizer of the apical cytocortex by regulating the localization and hence activity of Sqh. This conclusion is based on the observation that all defects observed upon loss of bbg, namely increased apical surface, reduced junctional tension, and reduced wing growth, were all rescued by the expression of a constitutively active form of Sqh.

**Discussion**

Wing growth is controlled by various signaling pathways, cell–cell and cell–matrix adhesion, cell shape, and cytoskeletal activity (Hariharan, 2015). Here we show that the scaffolding protein Bbg is an organizer of the apical cytocortex, and is required for cell shape and junctional tension, thereby regulating growth of the *Drosophila* wing imaginal discs.

bbg expression is highly dynamic in a variety of epithelia throughout development. Previous data (Kim et al., 2006) and results presented here suggest that the subcellular localization of Bbg proteins is cell type–specific, which could explain the distinct functions observed in different tissues. In the adult midgut, where bbg mediates the gut immune response, Bbg colocalizes with Coracle at septate junctions (Bonnay et al., 2013). In wing imaginal discs, however, Bbg could
not be detected at the septate junctions but rather in the apical cytocortex (this study). According to Flybase (http://flybase.org/), bbg encodes eight isoforms. Therefore, it is possible that the difference in Bbg localization is a result of the expression of alternative isoforms in different tissues, which may organize distinct protein complexes with cell-type specific localization and function. In fact, unlike in wing discs, knocking down bbg in migrating border cells had no obvious effect on actomyosin organization (Aranjuez et al., 2012). mRNAs of all predicted Bbg isoforms are expressed in wing discs (Tsoumpekos, 2016); therefore, the function described here cannot be allocated to any specific isoform. All Bbg proteins are scaffolding proteins with two or three PDZ domains. PDZ domains are protein–protein interaction modules, often found together with other protein–protein interaction domains in molecules that organize supramolecular protein complexes, which are involved in diverse biological processes, such as signaling, trafficking, adhesion, or growth (Subbaiah et al., 2011). Many of these processes depend on the ability to cluster functionally related components at defined cellular compartments at or close to the plasma membrane. Here we show that Bbg resides in a protein complex together with Sqh in the apical cytocortex of wing imaginal disc cells, but whether the two proteins interact directly remains to be elucidated. The organization of Bbg with three PDZ domains makes it an ideal candidate to coordinate components of the actomyosin network, including Sqh, and regulators of growth control.

Various results point to a close functional link between bbg and sqh in wing imaginal discs: both are required for proper wing size and show genetic interactions, their proteins colocalize in a common complex in the apical cytocortex of wing disc epithelial cells, both proteins are needed for maintaining proper junctional tension in the wing imaginal disc, and Bbg stabilizes Sqh in the apical cytocortex. Increasing evidence points to an important role of tension in growth regulation (Mao et al., 2011; Rauskolb et al., 2011, 2014; Sansores-Garcia et al., 2011; Le-
Wing pouch of L3 discs expressing UAS-sqhRNAi′ per genotype). (F and F′) area of each cell per image (three independent samples were quantified (E)). Measurement of the mean pixel

D E-cadherin staining from fixed tissues. (D) L3 wing discs tracked using the “Tissue Analyzer” plug-in from Fiji to quantify the surface area of the cells. Cell outlines were tracked from anti–

E-cadherin (F) and anti-Patched (Ptc; F′). en-Gal4, stained with anti–Dg and anti-Dlg, respectively. The dotted lines in A and A′ highlight the AP boundary. Images of bbg211 mutant (C) and WT (D) L3 wing discs tracked using the “Tissue Analyzer” plug-in from Fiji to quantify the surface area of the cells. Cell outlines were tracked from anti–DE-cadherin staining from fixed tissues. (E) Measurement of the mean pixel area of each cell per image (three independent samples were quantified per genotype). (F and F′) Wing pouch of L3 discs expressing UAS-sqhRNAi′ with en-Gal4, stained with anti–DE-cadherin (F) and anti-Patched (Ptc; F′). The dotted line in H highlights the AP boundary. The statistical analysis (F) used t test and ANOVA. *** P ≤ 0.001. Error bar shows SD. Bars: (A–B and F and F′) 25 µm.

Figure 7. Loss of bbg or sqh results in cells with larger surface areas. en>bbg′ [A and A′] and control [en-Gal4; B and B′] L3 wing discs, stained with anti-Bbg and anti-Dlg, respectively. The dotted lines in A′ and B′ highlight the AP boundary. Images of bbg211 mutant [C] and WT [D] L3 wing discs tracked using the “Tissue Analyzer” plug-in from Fiji to quantify the surface area of the cells. Cell outlines were tracked from anti–DE-cadherin staining from fixed tissues. (E) Measurement of the mean pixel area of each cell per image (three independent samples were quantified per genotype). (F and F′) Wing pouch of L3 discs expressing UAS-sqhRNAi′ with en-Gal4, stained with anti–DE-cadherin (F) and anti-Patched (Ptc; F′). The dotted line in H highlights the AP boundary. The statistical analysis (F) used t test and ANOVA. *** P ≤ 0.001. Error bar shows SD. Bars: (A–B and F and F′) 25 µm.

Goff and Lecuit, 2016; Sun and Irvine, 2016). Therefore, we hypothesize that Bbg controls growth in the imaginal discs by regulating tension through the organization of the actomyosin network. This assumption is supported by various observations. First, the protein complex pulled down with an antibody against Bbg contained, besides Sqh, other regulators of the actomyosin network, including the nonmuscle myosin heavy chain (called Zipper in flies), actin 57B, and α- and β-spectrin (Tsoumepkos, 2016). Spectrins are cytoskeletal scaffolding proteins, which are important for plasma membrane integrity and cytocortex organization (Machnicka et al., 2014), and modulate the activity of the apical actomyosin, thereby controlling the Hippo signaling pathway (Deng et al., 2015). In addition, mutations in bbg were found to modify the synapse growth phenotype induced by a dominant-negative mutation in Glued (Chang et al., 2013).

Glued encodes Dynactin 1, a subunit of the dynactin complex, which associates with cytoplasmic dynein, a motor protein involved in microtubule-based transport processes. Second, the apical surface was enlarged in bbg mutant wing disc epithelial cells, which is likely to be caused by a decrease in F-actin. In the absence of dachs, for example, which encodes an unconventional myosin, the apical surface of wing disc cells is larger, and wing size is reduced (Mao et al., 2011).

Third, bbg mutant wing discs showed increased apoptosis. This could also be a consequence of F-actin destabilization, because increased F-actin levels induced by overexpressing of the capping proteins α and β can decrease apoptosis (Amândio et al., 2014). Finally, in WT wing discs, Bbg, together with actin and Sqh, is enriched at the AP compartment boundary, an area of increased tension required to prevent cell mixing along the compartment boundary (Landsberg et al., 2009; Umetsu and Dahmann, 2015). In addition, Bbg is enriched in dividing cells, which require increased tension during rounding up (Rosa et al., 2015).

How Bbg, by maintaining proper junctional tension, regulates tissue growth remains to be elucidated. Tension has been reported to be a regulator of the transcriptional coactivator Yorkie (Yki), the Drosophila orthologue of mammalian Yes-associated protein/transcriptional coactivator with PDZ-binding motif (YAP/TAZ; Halder et al., 2012; Piccolo et al., 2014). This regulation can occur via the Drosophila kinase Warts (Wts; large tumor suppressor [LATS] in vertebrates; Wada et al., 2011; Rauskolb et al., 2014), a component of the Hippo pathway. Other studies suggest a more direct influence of the actomyosin on Yki activity (Dupont et al., 2011; Aragona et al., 2013). Reduced Yki phosphorylation (e.g., in the absence of the Warts or Hippo kinase) induces Yki translocation into the nucleus, where it up-regulates expression of antiapoptotic and proproliferation genes (Halder et al., 2012; Finch-Edmondson and Sudol, 2016; Sun and Irvine, 2016). Our preliminary results show that reduced growth in the absence of bbg is associated with reduced expression of the Hippo target gene Diap1, suggesting that bbg may regulate growth via the Hippo signaling pathway. This conclusion is in line with recent results showing that overexpression of Sqh in wing discs increases the expression of the Hippo target genes expanded and Diap1 (Rauskolb et al., 2014).

However, the canonical kinase cascade of the Hippo pathway is only one of several pathways that can regulate Yki activation and hence growth. For example, a recent study performed in Madine-Darbine canine kidney cells showed that tension mediated by the apical, circumferential actin belt represses translocation of Yki into the nucleus and hence tar-
get gene expression, a process that involves the interaction between E-cadherin and Merlin (Furukawa et al., 2017). Furthermore, vertebrate Yki has been documented to act as an apical sensor in some epithelial cells (Elbediwy et al., 2016). Although loss of bbg in wing discs does not affect overall epithelial polarity, E-cadherin localization, and tissue integrity, preliminary data indicate that some apical proteins (e.g., Crb) are affected in mutant cells. Therefore, it is tempting to speculate that Bbg, as a multi-PDZ scaffolding protein, is a versatile organizer of the apical cytocortex, thereby regulating the overall apical organization. This, in turn, is important for proper control of Yki or other regulators of growth. Identifying additional components recruited by Bbg and unraveling their function and possible interactions will further our understanding of the protein network localized in the apical cytocortex that fine-tunes tissue growth.

Materials and methods

Genetics

bbg^{B211} has previously been described a null allele (Kim et al., 2006), and was provided by G.L. Boulianne (University of Toronto, Toronto, Canada). The following RNAi lines were used: UAS-bbg^{RNAi} (III), UAS-bbg^{RNAi} (II), UAS-sqh^{RNAi} (III), and UAS-Irhp^{RNAi} (II; Vienna Drosophila Resource Center 15974, 15975, 7917, and 16758). UAS-Sqh^{E20E21} (III) and en-Gal4; DE-Cad::GFP/DE-Cad::GFP; UAS-GFP (Alice et al., 2012) and engrailed (en)-Gal4; UAS-RFP, sqh-GFP (II) were gifts from...
C. Dahmann (Technical University Dresden, Dresden, Germany). 69B-Gal4 and C765-Gal4, expressed in the whole wing disc, and engrailed-Gal4(hen), provided by S. Eaton (Max Planck Institute of Molecular Cell Biology and Genetics, Dresden, Germany), expressed in the posterior compartment of the wing, were used. Df(3R)ED4543/TM6C (8073) and yw, sqh^{WT}; sqh-GFP (Royou et al., 2004; 57144) were obtained from the Bloomington Stock Center (http://flystocks.bio.indiana.edu/). Unless otherwise stated, flies were raised at 25°C on standard food. Flip-out clones were generated using yw;hsflp;actinpromoter.FRT;STOP.FRT. clones were generated using otherwise stated, flies were raised at 25°C on standard food. Flip-out clones were generated using yw;hsflp;actinpromoter.FRT;STOP.FRT. clones were generated using

**Generation of anti-Bbg antibody**

Using a bbg cDNA as template, 0.67 kb, encoding 224 amino acids, was amplified by PCR using the following primers: forward, 5'-ATG TCGACGCAGTGGCAAGAGTCGAGGTCA-3'; reverse, 5'-GTG TCGACTCTTTTAGGGAGATCAGCCTC-3'. The amplicon was subcloned into the plasmid pCR2.1-TOPO (Invitrogen), digested with SalI (restriction sites included in the primers), and subcloned into pGEX-4T-1 (His)_6 C (based on the Amersham vector pGEX-4T-1 with a GST-tag, modified to also include a 6xHis tag; Kim et al., 2006), provided by G.L. Boulinaie. Expression of recombinant protein was induced (500 μM IPTG, 37°C, 4 h) using the BL21DE3 expression strain (Novagen). Tagged protein was purified using Ni-NTA Agarose beads (Invitrogen) according to the manufacturer's instructions. Tag-stagged recombinant protein was injected into female New Zealand White rabbits (2.5–3.0 kg). The serum was purified on HiTrap NHS-activated HP columns and concentrated in an Amicon Ultra 30K. The final concentration of the antibody was 5.64 mg/ml.

**Flow cytometry**

Approximately 20 L3 wing discs were dissociated into single cells using a solution containing trypsin and Hoechst 33342 (1:1000; diluted in PBS) for 1.5 h at RT. The samples were directly sorted using FACS. The flow cytometry was performed on a 5-laser BD FACS-Aria IIIu sorter (BD Bioscience) and analyzed using the FACS Diva software (v8.0; BD Bioscience) and the flow cytometry modeling software ModFit LT. Gates were applied as follows: a P1 gate was set on a side scatter/forward scatter (SSC/FSC) dot plot to identify live cells based on size and shape. The P1 fraction was restricted by setting a P2 gate on a SSC/GFP (exponential, blue laser, 488 nm). The P3 gate was generated on a BV2421-W/BV421-H (linear, UV laser, 375 nm) dot plot to discriminate singlets and to visualize the DNA content using the Hoechst 33342 dye. Out of the P3 population, a histogram for counts/ channel (linear, UV laser, 375 nm) was generated to analyze the cell cycle. Each 10,000 events from P2 were acquired to analyze the cell cycle of the different samples and genotypes. Voltage parameters were set based on WT controls.

**Antibodies**

The following antibodies were used at the indicated concentrations for immunofluorescence (IF) or WB/rabbit anti-Bbg (1:1,000; IF and WB; this work), mouse anti-Dlg (1:1,000; IF; DSHB 4F3), rat anti-DE-cadherin (1:1,000; IF; DSHB DCD2), rabbit anti-PH3 (1:1,000; IF; Millipore 06–570), rabbit anti-IgG (1:1,000; WB; Santa Cruz Biotechnology, Inc.), rat antitubulin (1:3,000; WB; AbD Serotec), rabbit anti-GFP (1:1,000; IF; and WB; Invitrogen), mouse anti-GFP (1:1,000; WB; Sigma-Aldrich), mouse anti-Ptc (1:100; IF DSHB apal), secondary antibodies conjugated with Alexa Fluor 488, 568, and 647 (1:1,000; IF; Invitrogen), or HRP (1:5,000; WB; Santa Cruz Biotechnology, Inc.). Phallolidin-488 (1:1,000; IF; Invitrogen) was used to label F-actin.

**IF and imaging**

Wing discs were dissected on ice, transferred to a sticky glue area on a common slide (e.g., WT with bbg^{251} mutant, together), and processed together. Therefore, IF and imaging conditions for different samples of the same experiment were exactly identical. Discs were fixed in 4% PFA in PBS for 20 min, washed in PBT (PBS and 0.1% Triton X-100) and incubated with the primary antibodies overnight at 4°C in blocking solution (PBT/5% BSA). Tissues were washed with PBT, incubated with secondary antibodies in blocking solution for 2 h at RT, washed with PBT, and mounted in Vectashield medium (Vector Laboratories). Images were acquired using a Zeiss LSM 700 inverted confocal microscope using Zeiss Plan-Neofluar 25x 0.8 Oil/Glycer/Water and Zeiss LCI Plan-Neofluar 63x 1.3 Glycer/Water DIC lenses at 23°C and processed using ZEN2010 and Fiji. For Fig. 6 and for the processing of stained images, the “Tissue Analyzer” plug-in from Fiji was used, which automatically measures different parameters, such as cell surface area. All images shown are projections of 5 µm (except those shown in Fig. 8, A–G’; sections were 1 µm each) and were representatives of the results obtained from several independent experiments (between 5 and 10 individual L3 wing discs and staining per genotype; more details in the legends to Figs. 2, 3, 6, S2, and S3). Fiji was used for quantification of cell numbers, PH3-positive, and TUNEL-positive cells. For this, a square of similar size was placed in the center of the pouch when comparing staining in whole discs, or in the center of the anterior and posterior compartment when comparing expression in these two compartments. For counting cell numbers, the Fiji plug-in “Cell Counter” was used. For measuring fluorescence intensity of Sqh or phallolidin, the same square selection was applied, and pixel intensity was measured using Fiji.

**TUNEL assay**

TUNEL assays were performed using the Roche in situ cell death detection kit (fluorescein, 1 684 795). In brief, L3 wing discs were fixed with 4% PFA in blocking solution for 20 min and washed with PBT. Discs were transferred to the reaction/ enzyme buffer containing terminal deoxynucleotidyl transferase (provided by Roche). After enzyme incubation at 37°C for 1 h, discs were rinsed three times in PBT and, after incubation in secondary antibodies in blocking solution for 2 h at RT, discs were washed with PBT and mounted in Vectashield medium (Vector Laboratories). Images were acquired using an LSM 700 inverted confocal microscope (Zeiss) using a 25x lens and processed using Fiji.

**Mounting of adult wings**

The left wings from female flies were dissected in PBS and mounted in Euparal MTNG medium (6372B). Wing size determination is the result of averaging 15 independent wing measurements (except from Fig. 1, which is derived from 50 wings), performed under strict temperature control (25°C), because the Gal4 system is temperature-dependent. By doing so, we tried to compensate for any phenotypic variability. Images were obtained with an Axionplan2 imaging microscope (Zeiss) using a 5x lens and processed using Fiji.

**WBs and immunoprecipitation (IPs)**

20L3 wing discs were dissected in PBS on ice and homogenized in 50 µl 2 × SDS loading buffer (100 mM Tris, pH 6.8, 4% SDS, 0.2% Bromophenol blue, 20% glycercol, and 2% β-mercaptoethanol) using a 1.5-ml Eppendorf tube and pestle. Samples were boiled, clarified by centrifugation, and run on Phos-Tag gels (12.5%; WAKO) for separation of phosphorylated and nonphosphorylated proteins. Proteins were transferred to nitrocellulose membrane 0.45 µm (GE Healthcare) and probed using the antibodies described in the Antibodies section. For IPs, 200 L3 wing discs were collected on dry ice and homogenized before addition of lysis buffer (50 mM Tris, pH 8, 0.5% Triton X-100, 150 mM NaCl, 1 µg/ml leupeptin, 250 µg/
ml Pefabloc, 2 µg/ml aprotinin, and 1 µg/ml pepstatin). The lysate was left on ice for 30 min and clarified by centrifugation. 1 mg total protein was used per IP. After adding the antibody, the lysate was incubated at 4°C for 2 h. 50 µl protein G agarose (GE Healthcare) per IP was added and left to rotate at 4°C overnight. Beads were washed six times with lysis buffer and boiled with loading buffer for 5 min at 100°C, and proteins were analyzed by conventional SDS-PAGE.

Laser ablation

Experiments were performed essentially as described (Farhadifar et al., 2007). Wing imaginal discs (genotypes: en-Gal4;DE-Cad::GFP/DE-Cad::GFP;UAS-GFP/UAS-bbgRNAi, UAS-bbgRNAi;UAS-Sqh20021, or UAS-sqhΔ40) were briefly rinsed twice in 70% ethanol, dissected in M3 medium (S3652; Sigma-Aldrich) and transferred to a 35-mm culture dish with glass bottom (P35G; MakTek). For the analysis of the vertex displacements, an inverted microscope with a 63x/1.2 numerical aperture water immersion objective was used. The laser beam was focused to a fixed spot of ~1.2 µm in the focal plane. The laser focus is targeted at adherens junctions. Images were recorded every 0.25 s over a period of ~30 s.

Online supplemental material

Fig. S1 shows that reduction of Irbp does not affect wing growth. In Fig. S2, RNAi-mediated reduction of bbg results in reduced cell numbers and increased apoptosis in L3 wing discs. Fig. S3 shows that the anti-Bbg antibody specifically detects Bbg molecules.

Acknowledgments

The authors thank G. L. Boulianne, S. Eaton, C. Dahmann, the Vienna Drosophila Resource Center and the Bloomington Stock Centers for flies and antibodies. Special thanks goes to Cagas Dolkás and Marcus Michel, laboratory members of C. Dahmann’s laboratory, for advice on laser ablation/cell tension experiments. We thank the following Max Planck Institute of Molecular Cell Biology and Genetics facilities: Patrick Keller, antibody facility, for antibody production, Ina Nüsslein for FACS analysis, and the Light Microscopy Facility, in particular J. Peychl, for microscopy guidance. The authors are grateful to Marino Zerial for critical and constructive comments on the manuscript, to members of the Knust laboratory for fruitful and constant advice, and to Francois Schweisguth for allowing the authors to conduct a few experiments in his laboratory.

This work was supported by the Max Planck Society. G. Tsoumpekos was a member of the International Max Planck Research School for Cell, Developmental, and Systems Biology and a doctoral student at Technische Universität Dresden.

The authors declare no competing financial interests.

Author contributions: G. Tsoumpekos designed and performed the experiments, interpreted the results, and was involved in writing the manuscript. L. Nemetschke performed the genetic screen and the initial interaction studies and was involved in editing the manuscript. C. Dahmann conceptualized the goal, supported data interpretation, wrote and edited the manuscript, and acquired funding.

Submitted: 16 May 2017
Revised: 22 September 2017
Accepted: 28 November 2017

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