Antagonistic functions of Par-1 kinase and protein phosphatase 2A are required for localization of Bazooka and photoreceptor morphogenesis in Drosophila

Sang-Chul Nam a,1, Bibhash Mukhopadhyay b, Kwang-Wook Choi a,b,c,*

a Department of Molecular and Cellular Biology, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA
b Program in Developmental Biology, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA
c Department of Ophthalmology, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA

Received for publication 28 September 2006; revised 27 March 2007; accepted 28 March 2007
Available online 1 April 2007

Abstract

Establishment and maintenance of apical basal cell polarity are essential for epithelial morphogenesis and have been studied extensively using the Drosophila eye as a model system. Bazooka (Baz), a component of the Par-6 complex, plays important roles in cell polarity in diverse cell types including the photoreceptor cells. In ovarian follicle cells, localization of Baz at the apical region is regulated by Par-1 protein kinase. In contrast, Baz in photoreceptor cells is targeted to adherens junctions (AJs). To examine the regulatory pathways responsible for Baz localization in photoreceptor cells, we studied the effects of Par-1 on Baz localization in the pupal retina. Loss of Par-1 impairs the maintenance of AJ markers including Baz and apical polarity proteins of photoreceptor cells but not the establishment of cell polarity. In contrast, overexpression of Par-1 or Baz causes severe mislocalization of junctional and apical markers, resulting in abnormal cell polarity. However, flies with similar overexpression of kinase-inactive mutant Par-1 or unphosphorylatable mutant Baz protein show relatively normal photoreceptor development. These results suggest that dephosphorylation of Baz at the Par-1 phosphorylation sites is essential for proper Baz localization. We also show that the inhibition of protein phosphatase 2A (PP2A) mimics the polarity defects caused by Par-1 overexpression. Furthermore, Par-1 gain-of-function phenotypes are strongly enhanced by reduced PP2A function. Thus, we propose that antagonism between PP2A and Par-1 plays a key role in Baz localization at AJ in photoreceptor morphogenesis.

Keywords: Cell polarity; Photoreceptor; Rhabdomere; Bazooka; Par-1; PP2A

Introduction

Genetic control of apical basal cell polarity is essential for epithelial morphogenesis and asymmetric cell division during cell fate specification. It is also important for development of polarized subcellular structures with specialized functions such as the light sensing organelles of photoreceptor cells. A small number of evolutionarily conserved proteins play important roles in diverse types of apical basal cell polarization. These polarity proteins form two major heterotrimeric cassettes consisting of Crumbs (Crb)-Stardust (Sdt)-Dpatj (Crb complex) and Par-6-aPKC-Baz (Par-6 complex) in the apical cell membrane. Recent studies have shown that these two protein complexes function in a coordinated fashion by direct protein–protein interactions (Hurd et al., 2003; Lemmers et al., 2004; Nam and Choi, 2003; Sotillos et al., 2004). However, the functional significance of these interactions and the mechanisms for precise subcellular targeting of these proteins remain to be elucidated. The Drosophila eye provides an excellent system to study in vivo functions of these interacting polarity proteins in control of cell polarity and organization of the rhabdomere, the light-sensitive apical structure of photoreceptor cells.

In Drosophila, about 800 ommatidial clusters comprising of 8 photoreceptor cells (R1–R8) are generated in the eye disc epithelium during the third instar larval stage, but morphogenesis of
photoreceptor cells takes place mainly during the following pupal stage. By 40% pupal development (pd), the apical region of each photoreceptor cell is involuted by 90°, which reorients the apical side toward the center of the cluster (Longley and Ready, 1995). In photoreceptor cells, Crb complex proteins are localized immediately apical to AJs (Fig. 1A). At 55% pd, when the rhabdomeres begin to develop from the apical surface of photoreceptor cells, Crb complex proteins are positioned to the region called the rhabdomere stalk, which links the rhabdomere with the AJ. During this time, developing rhabdomeres undergo dramatic vertical extension from the distal region of photoreceptor cells to the proximal base of the retina.

Crb, together with Sdt and Dpatj (Bachmann et al., 2001; Hong et al., 2001; Nam and Choi, 2003, 2006; Richard et al., 2006), is required for extension of rhabdomeres and formation of AJ along the distal–proximal axis of the photoreceptor cell, although it is not essential for establishing apical basal cell polarity (Izaddoost et al., 2002; Pellikkka et al., 2002). The mammalian homolog of Crb, CRB1, is also localized to the inner segment of photoreceptors, the structure analogous to the rhabdomere stalk, between the outer segment and the AJ (Pellikkka et al., 2002). Furthermore, mutations in CRB1 cause retinal diseases including retinitis pigmentosa 12 and Leber Congenital Amaurosis (LCA) in humans (den Hollander et al., 1999, 2001).

In addition to Crb, Par-6 complex is also required for proper organization and maintenance of apical photoreceptor membranes in the eye (Hong et al., 2003; Nam and Choi, 2003). Genetic evidence suggests that Par-6 complex is required for localization of the Crb complex to the apical membrane whereas the Crb complex may be necessary for maintenance of Par-6 complex proteins in the apical region (Hong et al., 2003; Nam and Choi, 2003). However, it is unknown whether any one of the Par-6 complex proteins plays a primary role in the Par-6 complex function during the organization of photoreceptor cells. Interestingly, although Par-6 and aPKC, like the Crb complex proteins, are localized to the apical membrane of photoreceptors, Baz is targeted to the AJ domain where Armadillo (Arm, Drosophila β-catenin) and DE-cadherin are localized (Nam and Choi, 2003). Recent studies have also shown the localization of Baz to AJ in the embryonic epithelia (Harris and Peifer, 2005). The differential localization of Baz and Par-6/aPKC raises the question of how Baz localization is regulated and whether Baz functions independently from Par-6 and aPKC.

In C. elegans, the Baz homolog, Par-3, is asymmetrically localized to the anterior cortex of the embryo whereas the serine/threonine protein kinase Par-3 is asymmetrically localized to the posterior side (Kemphues et al., 1988). In Drosophila follicle cells, Baz localization to the apical membrane is also regulated by basolateral Par-1 (Benton and St Johnston, 2003). Thus, in both C. elegans embryo and Drosophila follicle cells, Par-3 and Par-1 localize in a complementary cellular pattern. Interestingly, Baz is a phosphorylation substrate for Par-1 kinase, and the apical localization of Baz requires Par-1 phosphorylation (Benton and St Johnston, 2003). In follicle cells, Par-1 is required to exclude Baz from the basolateral membrane. Loss of Par-1 results in the basolateral expansion of Baz–Par-6 complex, indicating that Par-1 is essential for the regulation of apical basal cell polarity (Benton and St Johnston, 2003). As opposed to the essential role for Par-1 in follicle cell polarity, no significant function of Par-1 in regulation of cell polarity in the developing eye disc has been identified (Bayraktar et al., 2006). However, it has not been studied whether Par-1 is required at a later time point in photoreceptor morphogenesis for the control of Baz localization during the pupal stage when the photoreceptor cells undergo massive reorganization of their apical basal cell structure.

Since Par-1 is a protein kinase, an interesting question is whether Baz localization might be regulated by an opposing phosphatase activity. Evidence from mammalian studies has suggested that the formation of tight junctions is regulated by the interaction of protein phosphatase 2A (PP2A) and aPKC (Nunbhakdi-Craig et al., 2002). However, understanding the in vivo role of PP2A in cell polarity requires additional study, and the use of an animal model such as Drosophila eye can be a powerful tool to address these questions. PP2A is a heterotrimeric serine/threonine phosphatase composed of invariant catalytic (‘C’) and structural (‘A’) subunits and a variable regulatory subunit (‘B’) that directs the AC core complex to different substrates (Janssens and Goris, 2001). In Drosophila, the catalytic subunit is encoded by the microtubule star (mts) gene. mts mutants die in embryogenesis with defects in mitosis (Snaith et al., 1996), but the function of Mts in apical basal cell polarity has not been examined.

To understand the role of Baz and the regulation of Baz localization during photoreceptor morphogenesis, we investigated the functional relationship among Baz, Par-1 and PP2A in early/mid pupal eye development. First, we show that Baz plays an essential role in the localization of Par-6-aPKC complex to the apical domain. Second, Par-1 is required for the localization and/or maintenance of Baz and proper morphogenesis of photoreceptors. Thirdly, Baz unphosphorylated at the Par-1 sites is preferentially targeted to AJ whereas phosphorylated Baz is ectopically localized. Lastly, we identify an important function of PP2A in photoreceptor cell organization and its role as an antagonizing factor for Par-1. This study establishes the role for Baz as a central player in the localization of cell polarity proteins during photoreceptor morphogenesis and provides new insight into how Baz is regulated by antagonistic roles of Par-1 and PP2A.

Materials and methods

Genetics

Mitotic recombinations were induced by using FLP/FRT method for clonal analysis (Xu and Rubin, 1993). apks1803 and baz1625 mutants were produced in the eye by eyless (eyless)-FLP in y w par-61625 FRT226 ey FRT226 w Ubi-GFP FRT226, ey-Flip+ w FRT42D apks1803 FRT226 FRT42D Ubi-GFP. and w baz1625 FRT226 w Ubi-GFP FRT226 ey-Flip+ respectively (Petronczki and Knoblich, 2001; Wodarz et al., 1999, 2000). Overexpression of Mts, dmMts, Par-1 or Baz was induced by crossing UAS-Mts, UAS-dmMts (Hannus et al., 2002), UAS-Par-1 (full length of Par-1 a form, Sun et al., 2001), UAS-GFP:Baz; UAS-GFP; Bag1;4;ts18034 (Benton and St Johnston, 2003) with GMR-Gal4 (Freeman, 1996). mtsX2258 is a null allele of mts (Shioni et al., 1994; Wasserman et al., 1996) and par-1146 or par-1163, null alleles of par-1 (Cox et al., 2001; Huyh et al., 2001) were
used. par-1Δ16 is a deletion disrupting both par-1 and mei-W68 (Shulman et al., 2000), whereas par-1Δ16 is a par-1 specific deletion of complete kinase domain and most of the linker region of par-1 gene (Cox et al., 2001).

Antibody staining and histology

The following primary antibodies were used: rabbit anti-PKCζ (Santa Cruz), 1:500; mouse anti-Arm, 1:100; anti-α-Spectrin, 1:10; anti-Na+/K+ ATPase (Developmental Studies Hybridoma Bank), 1:50; rabbit anti-Baz (Wodarz et al., 1999), 1:500; rat anti-Crb (Bhat et al., 1999), 1:400; sheep anti-GFP (Biogenesis), 1:100; rabbit anti-Mts (Shiomi et al., 1994), 1:100; rabbit anti-Par-6 (Pinheiro and Montell, 2004), 1:500; rabbit anti-Par-1 (Vaccari and Ephrussi, 2002), 1:1000; rabbit anti-Dlg (Lee et al., 2003), 1:200; mouse and rabbit anti-Dpatj (Bhat et al., 1999), 1:500. Secondary antibodies conjugated with Cy3, Cy5, or FITC were from Jackson Laboratories. Fluorescent immunostaining of pupal eyes and confocal analysis were performed as reported (Izaddoost et al., 2002; Nam and Choi, 2003).

Results

Baz is essential for apical targeting of Par-6 and aPKC in photoreceptors

During early-mid pupal stage of eye development prior to the formation of the rhabdomere, Baz is localized at AJ of photoreceptor cells. Baz is known to form a protein complex
with Par-6 and aPKC, but it is not colocalized with them in photoreceptor cells, as Par-6/aPKC are targeted apical to the AJ (Fig. 1A) (Nam and Choi, 2003). Because the rhabdomere is not clearly formed at this stage, apical markers like Dpatj and Par-6 are localized in the most central region of each photoreceptor cluster surrounded by AJ markers, Arm and Baz, resulting in a characteristic star shape pattern in the tangential section (Fig. 1A).

Previous studies have shown that Baz is essential for localization of Crb complex and AJ formation during photoreceptor morphogenesis in the pupal eye (Hong et al., 2003; Nam and Choi, 2003). However, it is unknown whether Baz is required for the apical localization of its interacting partners, Par-6 and aPKC. To test this possibility, we analyzed the localization pattern of Par-6 and aPKC in loss-of-function (LOF) clones of baz null mutant cells. As the rhabdомерes develop, Baz is relocated from AJ to the rhabdомер in 65–70% pupal development (pd) (Hong et al., 2003). Therefore, we examined mutant clones in earlier pupal eyes at approximately 40% pd when Baz is restricted to AJ. As shown in Fig. 1, loss of Baz caused a strong reduction of Par-6 (Fig. 1B) as well as aPKC (Fig. 1C) to lower or nearly undetectable levels, whereas basolateral membranes of mutant photoreceptors were affected less severely or relatively intact (Supplementary Fig. 1). In contrast to the strong reduction of Par-6 and aPKC protein levels in baz mutant clones, significant levels of Baz protein expression were present in par-6 or apkc null clones, although Baz protein was abnormally distributed to the basolateral region, and Dpatj was displaced basally (Figs. 1D and E, see also Supplementary Fig. 2). These data suggest that Baz is a nodal component for apical targeting of Par-6 and aPKC whereas Par-6 and aPKC might be required for the maintenance of Baz at AJ.

**Par-1 is required for maintenance of apical and AJ proteins in pupal photoreceptor cells**

Based on the critical role of Baz for the localization of apical polarity proteins, it is important to determine how localization of Baz protein is regulated. In ovarian follicle cells, Par-1 is required to exclude Baz from the basolateral membrane, thereby restricting Baz localization to the apical membrane (Benton and St Johnston, 2003). Because Baz localizes to AJ rather than the apical membrane domain in photoreceptors, we examined whether the localization and function of Par-1 in photoreceptors might be different from that in ovarian follicle cells. At 40% pd, Par-1 is broadly distributed in the cytoplasm of photoreceptor cells. Double-labeling for the AJ marker Arm or a basolateral marker α-Spec showed weak and diffused Par-1 staining near basolateral membrane, but its level was considerably elevated in the apical region near the center of each ommatidial cluster (Supplementary Fig. 3).

The enhanced distribution of Par-1 in the apical region of photoreceptor cells suggests that Par-1 might play a role in photoreceptor morphogenesis. Thus, we examined whether Par-1 is required for correct localization of apical/AJ markers and photoreceptor development. Using par-1Δ16 null mutation, we generated par-1 LOF clones in pupal eyes and examined mutant phenotypes at about 40% pd. Par-1 protein expression was not detected in par-1Δ16 null mutant clones (Fig. 2A), confirming that this allele is protein-null (Cox et al., 2001). par-1 mutant clones showed a range of abnormalities, depending on the position along the proximal-distal axis of the photoreceptor cells. In the distal region of photoreceptor cells, toward the periphery of the retina, Arm was mildly affected in approximately 80% of mutant ommatidia (N>100, Fig. 2B). About 20% (N>100) of ommatidia showed relatively normal pattern of Arm and Dpatj staining (Fig. 2B, arrowheads). However, serial confocal sections of such par-1 mutant ommatidia showed mislocalization of Arm staining to basolateral position and occasional loss of AJ marker staining in the deeper proximal region of the retina (Fig. 2B, Supplementary Fig. 4). As shown in Fig. 2C, Baz is prominent in the distal section but greatly reduced at more proximal sections. An apical polarity protein, Crb, was also reduced or lost at proximal region of par-1 mutant ommatidia (Fig. 2C). These phenotypes suggest that Par-1 is necessary for the proper proximal–distal extension or maintenance of AJ and apical domains during the distal–proximal elongation of photoreceptors in pupal stages. However, despite such distal–proximal defects, Dpatj and Crb remained apical to Arm, indicating that apical basal cell polarity is not noticeably affected in par-1 mutant photoreceptor cells. Basolateral marker Dlg appears to be intact despite the ectopic presence of Arm, suggesting that a low level of ectopic Arm is not sufficient to disrupt the basolateral membrane property (Figs. 2D and E). Clonal analysis of par-1Δ16, a deletion mutant of par-1 and adjacent mei-W68 genes (Cox et al., 2001), showed the same phenotype of par-1Δ16 (not shown), further supporting that the par-1Δ16 phenotypes are due to loss of Par-1 function.

**Par-1 overexpression causes mislocalization of Baz and disruption of cell polarity**

Our data suggest that Par-1 is required for maintenance of AJ and apical markers during distal–proximal extension of the photoreceptor apical domain but not for regulation of apical basal polarity per se. The possibility remains that Par-1 may be involved in controlling cell polarity but its function in cell polarity may be compensated by unknown redundant factors. To test whether Par-1 can induce changes in cell polarity, we overexpressed Par-1 in differentiating retinal cells by GMR-Gal4 and examined whether it alters the localization pattern of Baz and other polarity proteins. Overexpression of wild-type Par-1 at 25 °C resulted in dramatic disruption of the apical basal pattern of photoreceptor cells (Figs. 3A and B). The AJ region marked by Arm was displaced irregularly and was diffused more basally or apically and into the cytoplasm from the normal AJ position. The Dpatj apical marker was also mislocalized and diffused, often more basally to AJ (Fig. 3B, arrows). Baz, which normally colocalizes at the AJ with Arm, was mislocalized in a domain overlapping the Arm mislocalization (Fig. 3C). Misplaced positioning of the AJ and apical markers suggests that overexpression of Par-1 leads to a disruption of apical basal photoreceptor cell polarity. This phenotype was highly consistent with 100% (n>100) of ommatidia showing...
similar defects in cell polarity. In contrast to the severe defects caused by overexpressing wild-type Par-1, no significant effects were observed when a kinase-null Par-1 mutant, Par-1(KN) (Sun et al., 2001), was expressed with the same GMR-GAL4 control under similar conditions (Fig. 3D). As in the wild-type eye, Arm was tightly localized to AJ, and Dpatj was restricted to...
the apical region. Therefore, the mislocalization of AJ and apical markers induced by Par-1 appears to be due to the Ser/Thr kinase activity of Par-1.

**Unphosphorylatable Baz protein is localized to the normal position at AJ**

As Par-1 kinase activity is important for inducing Baz mislocalization, and Baz is a biochemical substrate for Par-1 protein kinase (Benton and St Johnston, 2003), it is possible that Par-1 phosphorylation of Baz may be responsible for the mislocalization of AJ and apical markers (Fig. 4). To test this possibility, we expressed GFP-tagged wild-type Baz (BazWT) in differentiating retinal cells and examined whether the GFP-Baz proteins are normally localized to AJ or are recruited to ectopic positions in the photoreceptor cells. As shown in Fig. 4A, most GFP-BazWT was severely displaced to apical or basolateral regions. The apical marker Dpatj was also diffused and mislocalized basolaterally from the apical domain, implying the disruption of apical basal cell polarity (Figs. 4A–A″). This phenotype of BazWT overexpression was similar to the defects caused by overexpression of Par-1 (Figs. 3A and B), suggesting that the mislocalization effects of Baz overexpression might be correlated with the Par-1 kinase activity to Baz.

To test this idea further, we examined the distribution of a unphosphorylatable BazS151A,S1085A (BazSA) that has serine-to-alanine substitutions at the Par-1 phosphorylation sites, S151 and S1085 (Benton and St Johnston, 2003). As hypothesized, most BazSA protein was predominantly localized to the normal AJ positions basal to the apical Dpatj domain (100% ommatidia, N>100) (Fig. 4B). Consistent with the relatively normal localization of BazSA, adult flies with BazSA overexpression showed near normal eyes (not shown), although the expression levels of BazWT and BazSA were similar based on the level of GFP tag (Figs. 4A″ and B″).

**Inhibition of PP2A phenocopies Par-1 overexpression**

Our data shown above suggest that the absence of phosphorylation at the Par-1 phosphorylation sites of Baz is required...
to restrict Baz localization to AJ and to maintain photoreceptor cell polarity. A potential mechanism for inhibiting the Par-1 kinase function may involve the activity of a protein phosphatase(s). One candidate phosphatase, PP2A, has been implicated in a variety of functions, including establishment or maintenance of apical basal cell polarity in mammalian epithelial cells (Nunbhakdi-Craig et al., 2002). Therefore, we examined the possibility of whether Mts, the catalytic subunit of Drosophila PP2A, might be required for proper localization of Baz and the organization of cell polarity by antagonizing the Par-1 function. Because mts null mutant cells in the eye fail to survive even to a mid-pupal stage (Hannus et al., 2002; Sathyanarayanan et al., 2004; Wassarman et al., 1996), we took an alternative approach to inactivate wild-type Mts function by expressing a dominant-negative mutant form of Mts (dnMts). dnMts was generated by a truncation in the N-terminal region of Mts and shown to interfere with the PP2A activity specifically (Hannus et al., 2002; Sathyanarayanan et al., 2004).

Overexpression of dnMts by GMR-Gal4 in developing eyes resulted in severe defects in the apical basal pattern of most photoreceptor clusters (N > 100), as indicated by abnormal positioning of Dpatj and Arm markers (Figs. 5A and B). In most cases, the apical and AJ domains were mislocalized. Arm staining was mislocalized to basolateral or more apical positions as compared to Dpatj (Fig. 5B, arrows), suggesting that the apical basal polarity is altered. Baz was also mislocalized together with Arm when dnMts was overexpressed (Fig. 5C). In contrast to severe mislocalization of apical and AJ markers in the photoreceptor cells of pupal eyes, overexpression of dnMts in larval eye discs caused no significant change in relative localization of polarity markers Dpatj and Arm. Eye discs from GMR>dnMts larvae cultured at 18 °C are normal, but transient overexpression of dnMts during pupal stage by a temperature shift to 29 °C caused similar defects in mid-pupal eye as shown in Figs. 5A–C. These observations indicate that the dnMts phenotype is likely to be due to an inhibition of PP2A function specifically during a later time window of pupal development.

Remarkably, the pattern of apical and AJ marker mislocalization induced by Par-1 overexpression was almost indistinguishable from that of dnMts (Figs. 3B and 5B). This result is consistent with the idea that Par-1 and PP2A have opposing functions in the localization of polarity proteins.

**PP2A function is antagonistic to Par-1 and Baz**

To further probe the suggestion that Par-1 and Mts have opposing functions, we examined whether these two genes show an antagonistic genetic interaction. Since the eye phenotype of Par-1 overexpression is similar to that of PP2A inhibition, we expected that coexpression of Par-1 and dnMts would enhance the phenotype of either Par-1 overexpression or dnMts. Indeed, concomitant expression of Par-1 and dnMts resulted in 100% lethality, an outcome not seen by overexpressing either protein alone. To confirm such genetic interaction in the eye, we overexpressed Par-1 in the mts/+ heterozygous background to reduce the Mts level. Par-1 overexpression led to a reduction of the eye size and roughening of external morphology (Fig. 6B). This phenotype was strongly enhanced by the reduced mts gene dosage (+/mts), resulting, with 100% penetrance, in flies with smaller and rougher eyes (Fig. 6C).

We also examined whether the enhanced pupal eye phenotype by Par-1 overexpression in +/mts background correlated with the phenotypes in the pupal photoreceptor cells. As shown earlier, Par-1 overexpression causes mispositioning of the AJ and apical markers in pupal eyes (Fig. 6E). The reduced mts gene dosage under this condition (GMR>Par-1; mts/) resulted in a strong
reduction or loss of AJ and apical markers in photoreceptor cells at 40% pd (Fig. 6F), suggesting that these markers are not targeted to the membrane or may not be stably maintained. This result further supports the idea of an antagonistic relationship between Par-1 and PP2A.

Discussion

Par-6, aPKC and Baz play important roles in asymmetric polarization of cells. These conserved proteins act in a complex in Drosophila and other animal species, but it is largely unknown how this complex is formed and targeted to specific membrane domains. In this study, we have analyzed the function of Baz, Par-1 and PP2A in apical targeting of polarity proteins during photoreceptor morphogenesis in Drosophila and found antagonistic function of Par-1 and PP2A in controlling the localization of Baz.

Requirement of Baz for Par-6 and aPKC localization in the photoreceptor cells

Our clonal analysis suggests that Baz is crucial for targeting or maintenance of Par-6 and aPKC. In contrast, Baz protein expression was not significantly reduced in par-6 or apkc null clones, although Baz protein distribution was mislocalized to the basolateral region. This implies that Baz plays a nodal role among the Par-6 complex proteins (Figs. 1C and D). A caveat in this analysis is that par-6 and apkc mutant clones are very small (1–2 ommatidia) compared to the relatively large baz mutant clones, raising the possibility that par-6 and apkc mutant clones analyzed may represent rare escaper cells that survive with weaker phenotypes. However, this may not be the case because nearly identical phenotypes were seen in more than 50 par-6 or apkc mutant clones (Supplementary Fig. 2). Our data suggesting the central role for Baz in Par-6 and PKC localization are also consistent with studies on embryonic epithelia, in which Baz localization to the membrane precedes localization of Par-6, implying Par-6-independent membrane localization of Baz (Harris and Peifer, 2005). The requirement of Baz for the localization of Par-6/aPKC but not vice versa suggests that initial Baz localization may be independent of Par-6 and aPKC, as reported in embryonic epithelia (Harris and Peifer, 2005).

Similar relationships among Par-6 complex proteins have also been observed during asymmetric cell division in C. elegans. PAR-3, the homolog of Baz, is properly localized in the absence of either PKC-3 (aPKC) or PAR-6, whereas it is...
indispensable for localization of both PKC-3 and PAR-6 (Hung and Kemphues, 1999). These studies suggest that Baz/Par-3 is a major component in the control of cell polarity in diverse systems, including the photoreceptors in Drosophila. However, the Par-6 complex may exist in different compositions with unique functions, depending on various developmental contexts. For instance, Baz/Par-3 and Par-6 colocalize to the apical cortex of dividing neuroblasts in the Drosophila CNS (Bilder, 2001) and in dividing cells in C. elegans embryos (Watts et al., 1996), whereas Baz is distinctly localized to the AJ basal to the Par-6 domain in the apical membrane of photoreceptors (Nam and Choi, 2003). It is also striking that whereas Baz is not critically required for photoreceptor differentiation in the larval eye imaginal disc, it becomes crucial during pupal eye development. These data suggest that Baz is required for specific developmental events such as junctional reorganization and rhabdomere formation, although it is expressed in photoreceptors from the time of neuronal fate specification as well as in undifferentiated cells prior to retinal development.

Role of Par-1 in the localization of Baz at AJ

Par-1 is a key regulator of Baz localization in ovarian follicle cells. In this system, Par-1 is localized to the basolateral membrane and is essential for exclusion of Baz expression from the basolateral membrane (Benton and St Johnston, 2003). During early embryogenesis, Par-1 is transiently restricted to the lateral membrane, but at mid-gastrulation it is localized near the apical domain immediately below the region of spot adherens junction (SAJ). Par-1 is required for the restriction of SAJ, preventing the expansion of the E-Cad SAJ marker into the lateral membrane, but not affecting Crb-Dpatj apical markers (Bayraktar et al., 2006).
In contrast to follicle and embryonic epithelia, in eye imaginal discs, par-1 LOF clones show no significant apical basal polarity defects, suggesting that Par-1 is not required for cell polarity in eye imaginal disc epithelia (Bayraktar et al., 2006). This raises a question of whether Par-1 function is dispensable for photoreceptor morphogenesis. In this study, we focused our analysis on pupal eye development, since some cell polarity genes such as crb are not required in larval imaginal discs (Izaddoost et al., 2002), although they become essential later during the pupal stage when the retina undergoes dramatic reorganization of cell junctions and the apical basal pattern in the photoreceptor cells. Our analysis of pupal eyes suggests that Par-1 is required for the distal–proximal growth or maintenance of apical and AJ domains of photoreceptor cells, as Baz and apical markers often fail to form continuous AJ and rhodopsin along the distal–proximal axis of the retina in par-1 null mutant clones. These phenotypes are similar to the defects shown previously in the eyes of Crb complex mutants (Hong et al., 2003; Izaddoost et al., 2002; Nam and Choi, 2003, 2006; Pellikka et al., 2002). Like par-1 mutations, loss of these genes functions also affects the extension/maintenance of AJ and rhodopsin but not apical basal cell polarity. Since Baz is essential for proper targeting of Par-6 and Crb complex proteins, loss of Par-1 function may result in mislocalization of Crb complex through affecting Baz localization, although it is possible that Par-1 may also be directly involved in localization of Crb complex proteins independent of Baz. In the eye imaginal disc, it has been reported that Par-1 is localized to the apical-marginal zone and AJ (Bayraktar et al., 2006). In the pupal eye, we also found that Par-1 is enriched in the apical region of photoreceptor clusters, although a low level of Par-1 is also detected broadly along the basolateral membrane (Fig. 2A). Thus, Par-1 localization is not restricted to the basolateral membrane but appears to be regulated in a complex pattern in different cell types.

**Antagonistic interaction of Par-1 and PP2A**

In ovarian follicle cells, phosphorylation of Baz by Par-1 is required for proper localization of Baz to the apical region of the cells, and BazSA mutated proteins are abnormally localized to the basolateral membrane (Benton and St Johnston, 2003). In contrast, our data show that under conditions of overexpression, Baz protein mutated at Par-1 phosphorylation sites is targeted to the AJ whereas wild-type Baz is ectopically localized. Overexpressed wild-type Baz may be abnormally targeted to non-AJ sites, but it is also possible that ectopic Baz may recruit AJ proteins to form ectopic AJs. Nonetheless, our data suggest that, in photoreceptor cells, Par-1-dependent phosphorylation is not essential for initial localization of Baz to AJ. Instead, dephosphorylation of Baz may be a key for the localization of Baz to AJ. This explanation is consistent with the data that the pattern of AJ and apical markers is severely disrupted by overexpression of Par-1 but not by loss of Par-1, suggesting that Baz phosphorylation by Par-1 must be suppressed to maintain photoreceptor cell polarity.

We also tested whether the phenotypes induced by the overexpression of Par-1 might be competitively suppressed by co-expression of the non-phosphorylatable BazSA, but could not detect obvious suppression. In this experiment, both Par-1 and BazSA were overexpressed together using the same Gal4 driver. Therefore, a significant level of Par-1 might be available to phosphorylate endogenous Baz already present at the AJ even in the presence of overexpressed BazSA, thereby causing mislocalization or expansion of endogenous Baz. Suppression of Par-1 overexpression phenotype may be possible if BazSA can be expressed at a much higher level than Par-1, although it may not be straightforward to achieve such condition. Nevertheless, differential localizations of BazSA and wild-type Baz, together with the strong polarity defects by Par-1 overexpression, suggest that Par-1 negatively regulates the Baz localization to AJ of photoreceptors. Our data provide genetic evidence to support that Mts is a major enzyme responsible for antagonizing the effects of Par-1-dependent Baz phosphorylation. PP2A has been implicated in the regulation of tight junction formation in MDCK epithelial cells by interacting with aPKC (Nunbhakdi-Craig et al., 2002). However, it is unlikely that the Mts role in Baz localization is mediated through aPKC. First, wild-type and mutant BazSA are localized to completely different sites even though both have an intact phosphorylation site for aPKC. Second, the phenotype of Par-1 overexpression is mimicked by inhibition of Mts (Figs. 3 and 5) but not by loss of aPKC (Fig. 1D). Based on these observations and available information, we propose a model in which the localization of Baz to AJ in
photoreceptors cells during early pupal eye development depends on the reversal of Par-1-mediated phosphorylation by Mts PP2A activity (Fig. 7). In this model, Mts plays a pivotal role in regulation of Baz localization to AJ by antagonizing the Par-1 function. The idea of specific antagonism between Par-1 and Mts is supported by further enhancement of the Par-1 overexpression phenotype upon reduction of mts gene dosage (Fig. 6) but not apkc (data not shown).

Another approach to support the antagonistic interaction between Par-1 and PP2A is to test whether the Par-1 overexpression phenotypes can be suppressed by coexpression of Mts. However, for unknown reasons, overexpression of wild-type Mts by GMR-Gal4 resulted in early pupal lethality even at low temperatures. An alternative test is to see whether par-1 mutant clones can be rescued by inhibition of PP2A. Overexpression of dnMts in par-1 clones generated by the MARCM (Mosaic Analysis with a Repressible Cell Marker) method (Lee and Luo, 1999) showed no obvious rescue effects (data not shown). This was expected because in the absence of Par-1, there would be no phosphorylated Baz substrate for PP2A to act on. It should be noted that dnMts overexpression in par-1 MARCM clones generated only a few very small par-1 clones, suggesting that further analysis of larger clones under an optimal mutant condition would be desirable. Nevertheless, our data in this study are consistent with the proposed model on the antagonistic roles of Par-1 and PP2A in Baz localization during mid-pupal photoreceptor morphogenesis. It will be interesting to see whether the antagonistic interaction of Par-1 and PP2A plays an important role in regulation of Baz localization and function in various developmental contexts in Drosophila and other animal species.

Acknowledgments

We gratefully acknowledge H. J. Bellen, M. A. Bhat, K.-O. Cho, Y.-N. Jan, S. Eaton, D. St Johnston, J. A. Knoblich, E. Knust, B. Lu, D. J. Montell, T. Uemura, A. Wodarz, the Bloomington Drosophila Stock Center, and the Developmental Studies Hybridoma Bank for providing flies and antibodies. We thank Z. Chen and Y. Chao for technical assistance, and K.-O. Cho, Y.-C. Hsu, J. Lim, B.-J. Park, A. Singh, D. Wycuff and L. Yang for critical reading and comments on the manuscript. Confocal microscopy was supported by a grant from the NIH to D. B. Jones. This work was supported by grants from the NIHROI1 EY012210 and the Retina Research Foundation to K.-W. Choi.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.ydbio.2007.03.522.

References


