Combover/CG10732, a Novel PCP Effector for *Drosophila* Wing Hair Formation

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Introduction

Planar Cell Polarity (PCP), the polarity within the epithelial plane, is a characteristic of many epithelia in vertebrates and invertebrates and is established under the control of the non-canonical Wnt/Fz-PCP signaling pathway. In vertebrates, PCP signaling is evident in the alignment of hair follicles [1,2] and stereocilia in the inner ear [3], and required for limb growth [4]. Non-canonical Wnt signaling also regulates directional cell migration and intercalation during convergence and extension (C&E) during vertebrate gastrulation and kidney development [5–8] and aberrant PCP signaling thus can lead to severe birth defects (reviewed in [9,10]). In *Drosophila*, PCP signaling controls cell fates and orientation of ommatidia in the facet eye as well as the formation and orientation wing hairs (trichomes; reviewed in [11–14]).

A set of core PCP factors including the transmembrane proteins Fz, Flamingo (Fmi; aka. Stan), Van-Gogh (Vang, aka. Sbm), the adaptor proteins Dishevelled (Dsh), and Prickle are required for PCP establishment in all tissues (reviewed in [11–13]). Their interplay during PCP establishment leads to their asymmetric localization within cells with Fz and Dsh localizing to the distal and Vang and Prk localizing to opposite proximal vertex of hexagonal wing cells. These asymmetries are thought to act as cues interpreted by downstream effector genes for the establishment of polarity dependent structures [11–13]. In particular, each wing cell initiates the growth of a single trichome, an actin and tubulin rich wing hair [15,16] at the distal vertex at around 30 hrs after puparium formation (APF) [15–18]. About 17 hrs later, the trichome has developed into a cuticle ensheathed, rose-thorn shaped spike filled with a highly organized actin and microtubule fibers [19] that points towards the distal wing tip. In core PCP mutants, a wing hair typically forms in the center of a cell and shows aberrant polarity.

Planar cell polarity effector (PPE) genes, such as inturned (in), fuzzi (fy), and fritz (frtz), act downstream of the core PCP genes. In contrast to core PCP mutants, in PPE mutant wings, an average of two independent trichomes are initiated at various positions in the apical periphery of a wing cell (‘multiple hair cell’ (MHC) phenotype) [18]. A distinct phenotype with four hairs per cell is seen in *multiple wing hair* (*mwh*) mutants, some of which appear to be smaller secondary hairs splitting from larger ones [18]. Epistasis analyses and colocalization studies suggest that a complex of Int, Frtz, and Fy localizes to proximal, apical cell vertices in a core PCP gene dependent manner and prevents local hair initiation
and/or promotes distal hair initiation [18,20,21]. Specifically, PCP effector genes recruit and/or activate Mwh via direct interaction with In [22] leading to proximal enrichment of Mwh trailing off towards the distal end of cells [21,23]. mwh encodes a protein that resembles formins in that it contains a Rho family GTPase binding domain followed by a formin homology 3 domain with a potential for dimerization, but lacks a FH2 domain able to catalyze actin polymerization. Mwh may inhibit ectopic actin filament formation either directly, or by interfering with Rho GTPase activation of formins, or formin mediated actin polymerization [21,23]. Consistent with this, growing actin pimpls are initially seen all over the apical surface of a mwh mutant wing cell [21]. At around 34 hrs APF, Mwh relocates to the base of the forming forehair, where it prevents the formation of secondary trichomes [23].

Fz-PCP signaling also leads to the activation of Rho family GTPases such as RhoA, which in turn activates Rho kinase (Rok) to ensure proper cytoskeletal responses required for trichome formation in the wing and ommatidial rotation in the eye in Drosophila or directed cell migration during C&E in vertebrates [24–26]. In particular, loss of rok causes the appearance of multiple hairs per cell, albeit these trichomes still form at distal vertices and their appearance is thus mechanistically distinct from the action of other PPE genes such as fy or in [26]. The best-known substrate of Rok is Myosin II light chain regulatory kinase (MRLC, sgh), phosphorylation of which is required for myosin activity. Indeed, based on genetic interaction assays, it has been postulated that a proper balance between actin/myosin activities is essential for the formation of a single wing hair, as Myosin II can affect actin bundling [26].

To date, it is unknown how the In/Fy and Mwh PCP effectors cooperate with Rok during wing hair formation. We thus performed a genome-wide molecular screen for novel Rok substrates and identified CG10732 (now called Combover; Cmb) as a novel substrate of Rok. Overexpression of Cmb causes the formation of MHCs, a phenotype that was dominantly enhanced by removal of a gene copy of rok. In addition, the MHC phenotype of Cmb overexpression is enhanced by the fy/in group of PPE genes and mwh. We show that Cmb binds to Mwh and that mutation of cmb suppresses mwh in double mutants. We propose that Cmb is a novel PCP effector, the first one known to act downstream of mwh in wing cells during trichome formation.

Results

Combover (Cmb)/CG10732 is a novel direct substrate of Rok

To identify novel effectors of Planar Cell Polarity signaling, we performed a genome-wide, gel-shift based screen for Rho kinase substrates [27,28]. Briefly, pools cDNA clones of the Drosophila Gene Collections 1&2 were in vitro translated and internally labeled with [35]S-Methionine and incubated with the catalytic fragment of Rok (Rokcat) in the presence of unlabeled ATP. Candidate substrates were identified based on a reduced mobility on Anderson gels [28]. Compared to untreated control, incubation with Rokcat induced a gel-shift of in vitro translated clone GH01088 coding for CG10732-PB (Fig. 1A; blue arrows in Fig. 1C indicate the Rok dependent slow-migrating form of CG10732-PB). Importantly, incubation of the kinase reaction with calf intestinal alkaline phosphatase (CIP) strongly reduced the gel-shift (Fig. 1C, lane 3), indicating that the retarded gel migration is dependent on a (direct or indirect) phosphorylation event. CG10732 is thus a novel and uncharacterized potential target of Rho kinase phosphorylation.

Searches in Flybase [29] showed that CG10732, which we named Combover (Cmb), encodes four predicted splice isoforms (RA–RD). The open reading frame of the short isoform (PB) being contained within the longest one (PA; Fig. 1A). Isoforms cmb-RC and cmb-RD lack 96 bp or 99 bp towards the end of Exon 5 of cmb-RA (corresponding to Exon 4 of cmb-RB), due to removal of an extra intron possibly leading to in frame deletions of 32 and 33 aa, respectively. In contrast to the existence of cmb-RA and cmb-RB that is strongly supported by genomic data, the existence of cmb-RC and cmb-RD is only moderately supported [29]. Neither isoform contains known protein domains beyond a potential coiled-coil domain similar to SMC (structural maintenance of chromosomes) proteins (not shown) [30]. To assess when cmb evolved, we performed a database search using the BLAST algorithm against model metazoan sequences (see Fig. S1). cmb is present across a wide range of insect orders, with the only notable absence in the pea aphid Acyrthosiphon pisum (Fig. 1B and S2; alignments in Figs. S3 and S4). We identified a cmb ortholog in the crustacean Daphnia magna as well as in the tick Ixodes scapularis. We did not retrieve orthologs of cmb in any non-arthropod metazoan, thus suggesting that cmb evolved in the common ancestor of the Euarthropoda (i.e. chelicerates, myriapods, crustaceans and hexapods).

Phylogenetic analyses revealed that the cmb gene exists as a single ortholog in many dipterans (i.e. flies and mosquitoes), however many culicinorm (= mosquito) species have duplicated cmb paralogs, including up to three distinct paralogs in C. quinquefasciatus and A. aegypti (Fig. S2A and B). Our analysis revealed that cmb has been maintained in the genomes of a wide range of dipterans, and that cmb evolved in the last common ancestor of all arthropod clades.

We then confirmed that Cmb was a direct substrate of Rok in vitro. As full-length Cmb-PB was not soluble, we expressed overlapping GST-Cmb fragments (Fig. 1A) in E. coli and tested purified fusion proteins in kinase assays in the presence of [γ-32P]ATP. GST alone, GST-BB (aa 384–580 relative to Cmb-PB), and GST-SX (aa 496 to stop) were not phosphorylated by Rokcat (Fig. 1D, lanes 1, 2, 4; note autophosphorylation of Rokcat [31]). However, the N-terminal ES fragment of Cmb-PB (aa 1–495) was directly phosphorylated by Rokcat (Fig. 1D, lane 4), leading to the prediction that the Rok phosphorylation site(s) lie within the first 382 amino acids of Cmb-PB (Fig. 1A). GST-ES thus was thus subject to phosphorylation by cold ATP and the phosphorylation sites mapped by mass spectrometry. Relative to the start codon of Cmb-PB, T46, T026, S300, T368, and T370 were identified with T368 being phosphorylated at very low levels. Except for S300, all sites are followed by Proline, uncommon for Rok phosphorylation sites that are usually preceded by a positively charged amino acids at position [−1] or [−2] (see [32–36]). We mutated these five candidate sites to Ala and introduced an additional mutation (T372A) as it is a third Thr in a [TP]3 repeat together with T368, and T370. The mutated Gst-ES wild type was then tested in direct kinase assays. Fig. 1E shows that compared to Gst-ES (middle set), most phosphorylation of the phosphorylation by Rokcat is lost in the Gst-ESA mutant (right panel; Fig. 1F shows quantification of the triplicates shown in Fig. 1E) indicating that the major phosphorylation sites of Cmb were correctly identified and that Cmb is a novel Rok substrate in vitro.

A role of Combover in actin wing hair formation

In order to assess the physiological role of combover, we mutagenized the gene using homologous recombination techniques [37,38]. Briefly, we replaced 1023 bp of Exon 3 including the start codon and roughly half of Exon 4 of cmb-RB (equivalent to Exon 4 and half of Exon 5 of the other isoforms; Fig. 2A) with a
Figure 1. Cmb is a Rho kinase substrate. (A) Schematic of the PA and PB Cmb isoforms as annotated in Flybase. Clone GH01088 is a full-length clone corresponding to cmb-RB identified in the Rok target screen. Grey bars indicate GST fusions of Cmb-PB used to map phosphorylation sites in direct kinase assays in vitro, and Cmb-NT, Cmb-CT, and Cmb-Int indicate fragments used in two-hybrid and co-immunoprecipitation experiments. Phosphorylation sites identified by mass-spectrometry are numbered relative to Cmb-PB. Direct phosphorylation results of the Cmb Gst fragments are indicated towards the right. (B) Summary of the presence and/or absence of cmb and mwh genes in model arthropod genomes (see also Figure S2; phylogenetic reconstruction shown is based on [62–64]). mwh and cmb likely evolved in the last common ancestor of euarthropods (as represented by an ortholog in the tick Ixodes scapularis). The Diptera clade has been collapsed (see Figures S1 and S2). (C) Treatment of in vitro translated Cmb-PB (clone GH01088) with the catalytic fragment of Rho kinase (Rokcat) causes the formation of a slower migrating form of Cmb on an Anderson gel (compare blue arrow position with black one in lanes 1 and 2). The gel shift is due to phosphorylation, as it is reverted upon treatment with alkaline phosphatase (CIP; lane 3). Note that for unknown reasons (such as translation initiating at an internal methionine), two Cmb bands are seen upon in vitro translation of Cmb in reticulocyte lysates. (D) Phosphorylation of Cmb by Rok is direct. Indicated purified fragments of Cmb fused to Gst were incubated with Rokcat in the presence of [32] P-ATP and separated on a 12% SDS PA gel. Only GST-ES (lane 3) is a substrate of Rok. Top panel is an autoradiograph of the Coomassie stained gel in the lower panel. Arrow show Rokcat autophosphorylation (Rok) and indicated GST-fusion proteins. (E) Kinase assays in triplicate of indicated Gst fusion proteins. The upper panels show autoradiographs of the Coomassie stained gel in the lower panel. Arrow show Rokcat autophosphorylation (Rok) and indicated GST-fusion proteins. (F) Quantification of kinase assay shown in E. Error bars indicate standard deviation; T-test: ***p<0.0001.

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The allele obtained was homozygous viable (see below). The mutation was verified by inverse PCR using primers of the White cassette and genomic primers outside of the homology arms (not shown) and by directly assessing the deletion with primers specific for cmb (see Fig. 2A). While cmb primers were able to amplify the expected fragment of 945 bp from wild-type (w¹¹¹B) genomic DNA, no product was obtained from DNA of homozygous cmbKO flies (compare lanes 1 and 2 of Fig. 2B), as predicted for successful deletion. Control amplification of an unrelated locus confirmed the integrity of the DNAs (Fig. 2B, lanes 4, 5).

To further characterize the cmbKO mutant, we generated an antiserum against the ES fragment of Cmb-PB. cmb-RA and cmb-RB encode predicted proteins of 1657 amino acids (aa) and 809 aa with calculated molecular masses of 189 kDa and 89 kDa, respectively. Western blot analysis of lysates of w¹¹¹B 3rd instar larvae revealed two predominant proteins recognized by the antiserum of about 190 and 110 kDa (as calculated by their relative migration to markers) that were absent from cmbKO lysates (Fig. 2C; αTubulin was used as loading control, lower panel). Although the smaller isoform migrates at a higher apparent molecular mass than predicted by conceptual translation, the absence of both bands in the mutant show that the cmbKO allele is a protein null allele of cmb.

As cmbKO flies are viable and showed no gross anatomical defects, we assessed if cmb loss caused PCP phenotypes similar to loss of rok. Trichome polarity in the wing was normal and we found no multiple hair cell phenotype (compare wild-type wing area shown in Fig. 2E with cmbKO wing in Fig. 2F). Similarly, sections of adult eyes of homozygous cmbKO flies were normal, with no PCP defects (Fig. S5).

We next created transgenic flies overexpressing each Cmb isoform under UAS control to assess gain of function (GOF) phenotypes. Overexpression of either Cmb isoform under the control of sevenless-Gal4, usually an excellent driver to induce PCP defects in the eye [39–41], had no effect (Fig. S5C, D). However, overexpression of cmb-RA and cmb-RB in several independent transgenic lines under the control of en-Gal4 (compare en>Gal4 control wing in Fig. 3A with en>cmdRb4 and en>cmdRA3 in Fig. 3B and C, respectively) or nubbin-Gal4 (not shown) caused the formation of a multiple hair cell phenotype similar to loss of rok (and other PCP effectors) [18,26]. These data

**Figure 2. cmbKO is a protein null mutant.** (A) Genomic locus of cmb/CG10732 showing the RA and RB isoforms that are well supported by genomic data. 1023 bp of genomic DNA (red) was replaced with a White+ marker by homologous recombination to generate the cmbKO allele. The deleted fragment includes the start codon of the PB isoform. Arrows indicate approximate location of the PCR primers used to verify the deletion. (B) Analytical PCR shows that cmb specific primers amplify a 945 bp fragment from w¹¹¹B control DNA (lane 2), but not from homozygous cmbKO DNA (lane 1). Control primers amplify the expected 532 bp fragment from both DNAs showing their integrity (lanes 4, 5). Lanes 3, 6: No DNA controls. (C) Western blot analysis of 3rd instar larval lysates separated on a 12% SDS-PA gel shows that, in contrast to a w¹¹¹B lysate (left lane), neither Cmb-PA nor Cmb-PB (arrows; predicted MWs 189 kDa and 89 kDa, respectively) are detected in lysates of homozygous cmbKO flies. The minor form running above Cmb-PB may be a modified form and was not detected in all preparations. αTubulin was used as loading control (lower panel). (D–F) Wing hairs and their orientation of cmbKO flies are normal. Compare enlarged wing area of a w¹¹¹B wing (E) with a cmbKO wing in (F; area corresponds to blue box in D). Scale bar is 50 μm.

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thus suggest a function of Cmb in actin hair initiation or modification of actin myosin contractility. The frequency of ectopic hairs varied strongly between UAS-Cmb lines and conditions (not shown), suggesting strong dosage sensitivity.

To assess the specificity of the Cmb GOF phenotype and to assess if there may be a physiological relevance for the in vitro phosphorylation of Cmb by Rok, we tested genetic interactions between rok and Cmb overexpression. As shown for rok1 and rok2 alleles in Fig. 3D, E, removal of one gene dose of rok significantly enhanced the MHC frequency of cmb-RA overexpression (quantified in Fig. 3H). Similarly, cmb-RA overexpression is enhanced by a deficiency uncovering rok (Fig. 3F, quantified in Fig. 3H). Wings of flies heterozygous for rok look wild-type (data not shown). These results suggest that rok exerts a negative effect on overexpressed Cmb in vivo.

The multiple hair cell phenotype of Cmb overexpression is enhanced by PPE genes

To further define a potential connection between Cmb and PCP effectors, Cmb was overexpressed in the Drosophila wing in a heterozygous mutant background for members of the PCP effector family. Intriguingly, the MHC phenotype of cmb-RA overexpression (quantified in Fig. 3H). Similarly, cmb-RA overexpression is enhanced by a deficiency uncovering rok (Fig. 3F, quantified in Fig. 3H). Wings of flies heterozygous for rok look wild-type (data not shown). These results suggest that rok exerts a negative effect on overexpressed Cmb in vivo.

Figure 3. Overexpression of Cmb causes a MHC phenotype which is dominantly enhanced by alleles of rok. (A–C) Compared to control (A), overexpression of either isoform of Cmb (RB and RA in (B), (C), respectively) under the control of en-Gal4 causes the formation of multiple hairs cells specifically in the posterior compartment. (D–F) This MHC phenotype is dominantly enhanced by the removal of one gene dose of rok (rok1 (D), rok2 (E)), or a deficiency uncovering rok (DF(1)FDD-033126; F). (G) Schematic of wing indicating the approximate areas shown in panels A–F (blue box) and the second posterior wing cell scored for quantification (rose). (H) Quantification of MHC phenotype of Cmb overexpression in second posterior wing cell and enhancement by indicated alleles. 29°C. Depicted are mean and SEM; T-tests (*p<0.05; ***p<0.001); n=5. Scale bars are 20 μm. doi:10.1371/journal.pone.0107311.g003

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To assess the mechanistic basis of the genetic interactions, we tested whether Cmb would physically interact with PPE genes in yeast two-hybrid assays. Due to the size of Cmb, we tested interactions of Fy, Frtz, In, and Mwh bait proteins for interaction with the N- and C-terminal halves of Cmb as prey (see schematic in Fig. 1A). Interestingly, the N-terminal half of Cmb-PA interacted with Mwh in this assay under stringent selection conditions on medium simultaneously lacking Ade and His (Fig. 5A). In addition, growth correlated with the activation of the lacZ gene, a third reporter present in the two-hybrid tester strain. Neither Fy, Frtz, or In interacted with Cmb (Fig. 5A), also suggesting the Cmb-Mwh interaction is specific.

To independently confirm the interaction in a different system, we transfected HEK293 cells with GFP tagged Cmb-PA, Cmb-PB, or Cmb-Int, a fragment consisting of the N-terminal half of Cmb overlapping with Cmb-PB (Fig. 1A) together with Myc-tagged Mwh. Immunoprecipitation with anti-GFP antibodies efficiently co-precipitated Mwh (lanes 1–3 in Fig. 5C), but not a Myc-tagged control protein (Dazap1, lanes 5–7 in Fig. 5C). Importantly, GFP-Tbx1 did not pull-down Myc-Mwh (Fig. 5C Lane 4), indicating that Cmb specifically interacts with Mwh not only in yeast two-hybrid assays, but also in lysates of transfected cells.

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The Fy/In group of PCP effectors form a complex that is required for the proper localization and probably the restriction of Mwh activity to the more proximal side of early developing pupal wing cells [21–23]. Furthermore, the MHC phenotype of Cmb overexpression is enhanced by reduction of fy, frtz, and mwh (Fig. 4). Our physical and genetic interaction data suggested that Cmb contributed to trichome formation, a function that might be antagonized by PPE genes. If so, we hypothesized that loss of cmb might suppress the phenotype of mwh mutants. We thus recombined the cmbKO mutation with an amorphic mwh1 and a hypomorphic, temperature sensitive mwh6 allele [18,23,42] and assessed the phenotypes of homozygous double mutants raised at 25°C. Indeed, loss of cmb suppressed the phenotype of mwh1 and mwh6 alleles, as well as mwh1/mwh6 transheterozygotes (compare Fig. 6A–C with D–F). Quantification of the multiple hair cells in the second posterior wing cell (Schematic in Fig. 3G) showed that the suppression from a mean of 752±40 MHCs to 517±35 for mwh1, of 697±44 MHCs to 413±39 for mwh1/mwh6, and of 488±13 MHCs to 260±12 for mwh6, respectively, is statistically significant (Fig. 6G), thus supporting a model in which Cmb has a role in actin wing hair formation and is directly regulated by the PCP effector Mwh through a protein-protein interaction in vivo.

Localization of Cmb in pupal wing discs

In order to restrict wing hair formation to the distal end of cells, In and Frtz are localized to the apical, proximal side of developing wing cells [21,23]. Similarly, Mwh is enriched at the proximal edge of wing cells [21,23]. We thus stained pupal wing discs with our Cmb antibodies. Unfortunately, our antiserum does not detect endogenous Cmb in tissue samples (Fig. 7A, A'). cmb-RB and cmb-RA driven protein is localized apically in posterior wing cells at 30 hrs APF prior to wing hair formation (Fig. 7A, D). We noticed that the expression level of the transgenes was somewhat variable between different cells. In cells expressing Cmb at lower levels, it appears cortically enriched. At 36 hrs APF, once wing hairs started forming, Cmb protein remains distributed in a grainy pattern in the apical region of wing cells (Fig. 7B, B' for cmb-RB and not shown) and is largely excluded from more basal regions of cells (Fig. 7 C for cmb-RB). Apical restriction of both isoforms is also evident in Z-sections (Fig. 7E, F). We do not find Cmb localized to the actin hair itself (Fig. 7B). Significantly, the ectopic hairs form at the distal vertex (Fig. 7B) and no obvious difference in Cmb localization is seen between cells that show one or multiple hairs due to overexpression of Cmb (Fig. 7B, B' and B''). To express Cmb at lower levels and to address if Cmb can localize asymmetrically in cells, we
induced ‘flip-out’ clones expressing cmb-RA under control of the actin-Gal4. While we find apical cortical enrichment of Cmb-RA, there is no evidence for proximal-distal asymmetry (Fig. 7G, G’; expressing cells are labeled in blue in G). In conclusion, (overexpressed) Cmb is localized apically in pupal wing cells and is cortically enriched in cells that appear to express lower protein levels and thus overlaps with areas where Mwh (and other PCP effectors) localize.

Discussion

Rho kinase, a member of the AGC kinase family which also includes PKC and Akt (for review see [43,44]) was originally identified as a RhoA effector reorganizing the cytoskeleton by promoting the formation of actin stress fibers [45]. In Drosophila, Rok was shown to act downstream of Fz and Dsh in the non-canonical Wnt/Planar Cell Polarity pathway causing ommatidial rotation and structural defects in the eye and multiple hairs cells in the wing [26]. Here, we have identified Combover/CG10732 as a novel substrate of Rok. We created a cmb protein null allele lacking both Cmb protein isoforms (Fig. 2C) that is homozygous viable. Homozygous cmb mutants display no visible phenotype in the wing or in sections of the adult eyes (Figs. 2 and S5). As a reduction or an excess of actin polymerization can cause MHCs, we assessed the overexpression phenotype of Cmb. Indeed, overexpression of either Cmb isoform caused a multiple hair cell phenotype that is strongly dominantly enhanced by rok and the fy/in/mwh PCP effectors, validating our in vitro screening approach to identify PCP effectors. Importantly, the cmb mutation suppresses the MHC phenotype of mwh in double mutants.
data thus indicate that Cmb, while not essential for wing hair formation, nevertheless promotes trichome formation in vivo.

Rok phosphorluates unconventional sites on Cmb

It has been noted that known phosphorylation sites of Rok targets such as ERM proteins, Vimentin, Myosin regulatory light chain, or Adducin, often follow the consensus site [R/K]XX[S/T] or [R/K][X][S/T] [32–36]. Of the five Rok sites we have identified in our in vitro kinase assays followed by MS analysis, only S300 is preceded by a basic residue at position [-2] (RT[S]). In all other cases, no basic amino acid is found at position [-1] or [-2]. However, T46, T206, T368, and T370 are all followed by a Proline, more typical of MAP kinase phosphorylation sites [46]. Nevertheless, mutation of these sites strongly reduced Cmb phosphorylation in vitro (Fig. 1E).

In rok mutants, multiple hairs form at the distal end of wing cells [26]. Similarly, overexpression of either Cmb isoform causes MHCs that originate at the distal end of cells (Fig. 7B), distinct from the in/fy group of PCP effectors and mwh, which form MHCs around the periphery of the cells (note that in mwh mutants, actin patches are initially even formed all over the apical cell surface) [18,21]. Importantly, reduction of rok activity by the removal of one gene dosage (by two different alleles or a deficiency) increases the number of MHCs (Figure 3), suggesting an inhibitory effect of Rok on Cmb. It was suggested that Myosin II, which is concentrated at the site of prehair initiation and whose activity is regulated by Rok via phosphorylation of its regulatory light chain (MRLC), must be within an optimal range to properly bundle actin and to ensure the formation of a single hair [26]. Consistent with the genetic interaction between cmb and rok, it is possible that in addition to regulating MRLC, Rok might also inhibit a potential hair promoting activity of Cmb (see model in Figure 8), although we cannot exclude that Rok/MRLC activity acts in parallel to the effect Cmb exerts on wing hair formation.

Cmb as a PCP effector during wing hair formation

mwh and the in/fy group of PCP effectors all have been implicated in restricting actin hair initiation to the distal vertex of the cells by inhibiting proximal hair assembly [18,20,21,23,47]. Core PCP signaling ensures proper proximal localization of In, Frtz, and Fy proteins (and thus inhibition of prehair formation) to the proximal end of wing cells [18,21,23] leading to the formation of a single wing hair on the distal side.

cmb-RA or cmb-RB cause the formation of MHC phenotypes upon overexpression with several wing drivers. Importantly, this overexpression phenotype is enhanced by the removal of one gene dosage of the PCP effectors fy, frtz, in, and mwh as well as deficiencies uncovering those loci. These genetic interactions suggest cmb could exert a positive effect on hair initiation, although such a function would play a supportive or redundant role as neither a lack nor ectopic trichomes are found in cmb mutants.

Significantly, we showed that Cmb physically interacts with Mwh in yeast two-hybrid and communoprecipitation assays (Fig. 5). Interestingly, while we were unable to identify vertebrate homologs of cmb, we found orthologs of both cmb and mwh outside of the insects in the genomes of the crustacean Daphnia magna and of the tick Ixodes scapularis (Figs. 1B, S1, S2; see also [21,23]). Ixodes is a member of the Chelicerata, the most basally-branching
Figure 7. Cmb localization. Pupal wing discs expressing en>cmb-RB or en>cmb-RA (green) in the posterior compartment (posterior is down, distal to the right in all panels). Actin is shown in red; greyscale pictures show indicated single channels. (A) Endogenous Cmb (anterior, above compartment boundary indicated by a yellow dotted line) is not detected by our antibody. At 30 APF prior to hair initiation, cmb-RB overexpressed in the posterior compartment localizes apically in a punctate pattern. (B–C) At 36 APF, after hair initiation, en>cmb-RB localizes apically (B, B') but is largely absent from more basal confocal sections (C, C'). Note the multiple wing hairs due to Cmb overexpression emerge on the distal side (C ''). Cmb is not present in the wing hairs. (D) en>cmb-RA localizes apically in 30 hr APF wing discs and appears cortically enriched in cells that express at a lower level. (E, F) Optical Z-sections of 36 hrs pupal wing discs shows strong apical enrichment of cmb-RB (E, E') and cmb-RA (F, F'; note that the dorsal and ventral wing layers are seen, with their basal sides touching and their apical sides facing away from each other, respectively). (G) Flip-out clones (marked by UAS-GFP in blue in G) expressing cmb-RA under the control of actin-Gal4 show that while cmb-RA appears cortically enriched, it is not asymmetric with respect to the P/D axis. Scale bars are 2 µm.

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The presence of mwh and cmb in Isodes may be indicative of an ancient protein-protein interaction that has been retained throughout arthropod evolution. Because both Isodes and Daphnia lack wings, the Mwh/Cmb interaction likely performed different, possibly additional function in the ancestral arthropod. Consistent with this, mwh mutants cause other cuticular hair defects in other regions of the Drosophila body [49]. Alternatively, the Mwh/Cmb interaction evolved much later than the appearance of both of these genes in the genome of the ancestral arthropod. The roles of and interactions between Cmb and Mwh proteins in more non-insect arthropods needs to be further explored.

The presence of both mwh and cmb orthologs in the genomes of members of all holometabolous insect orders may indicate that the Mwh and Cmb interaction is also conserved in this insect clade. The retention of these two genes in members of the more basally-branching hemipteran orders, however, is less conserved. The presence of both mwh and cmb in Holometabola may be due to the shared mode of wing development, i.e. via internal wing imaginal discs. This is in contrast to the mode of wing development in hemimetabolous insects by which the wings develop as buds outside of the body. Further study into the association of wing development and Mwh/Cmb interactions in other insect orders is needed to elucidate these findings.

Interestingly, PCP effector mutations generally enhance each other. For example, the hypomorphic frtz allele is enhanced by weak alleles of in or fy in double mutants [47]. Analogously, removal of a gene dosage of mwh in a fy or in background, enhances their MHC phenotype [50]. In contrast, the MHC phenotype of mwh null allele by a cmb null mutation further suggests that the role of Cmb on hair formation is antagonized by Mwh.

Unfortunately, our Cmb antibodies do not detect endogenous Cmb protein in the developing pupal wing. Nevertheless, Cmb expressed in the posterior compartment of the wing under the control of en-Gal4 localizes apically in a punctate pattern. In cells that appear to express at a lower level (seen particularly for Cmb-RA in Figure 7D, G) Cmb is enriched at the circumference of the cells, but shows no proximo-distal enrichment. Although we cannot exclude that Cmb localization is an overexpression artifact, this appears unlikely, because we would expect Cmb to fill the cells rather than to localize specifically apically (Fig. 7G). Importantly, Cmb likely localizes to the area of wing cells where Mwh is present, as Mwh known to be initially enriched apically towards the proximal side [21,23], further supporting our model that a positive effect of Cmb as a novel PCP effector on wing hair formation may be restricted by Mwh.

Materials and Methods

Ethics Statement

This study was carried out in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The antibody generation protocol was approved by the Institutional Animal Care and Use Committee of the Albert Einstein College of Medicine (Protocol number 20130514).

Fly strains

mwh1, mwh6, in1, fy2, fy3, frtz3 are described in Flybase [23,47,51,52]. Df(3L)Fpo2 (mwh), Df(2L)ED611 (fy), Df(2L)ED7762 (frtz), Df(3L)BSC667 (in) were obtained from the Bloomington stock center. rok1, Frt19A and rok2, Frt19a were a kind gift of Dr. L. Luo (Stanford University). Constructs for transgenic flies were injected by Rainbow Transgenic Flies and Genetic Services.

For adult wing analysis, wings were incubated in 0.1% Triton-X100 in PBS for at least one hour and subsequently mounted in 80% glycerol in 1xPBS. Multiple hair cells (MHCs) were counted on a minimum of 5 wings was utilized. Statistical analyses were performed using R version 3.0.2 (2013-09-25).

Plasmids

All PCR products used for cloning were sequenced. pCS3_Myc_Dazap1 was a kind gift of Dr. F. Marlow, Einstein. pCS_Tbx1_GFP was a kind gift of Dr. B. Morrow (Einstein).

pFastBacHisC was constructed by inserting Fast_bac_his_upper and Fast_bac_his_lower into the BamHI/HindIII sites of pFastBac1 (Life Technologies). pCRIITopo_rok was made by cloning the Rok open reading frame as a PCR product amplified with drok_Upper_BglII and drok_lower_BamHI into pCRIITopo (Invitrogen). pFastBacRGSHis_rokcat was made by cloning a BglII/EcoRI fragment of pCRIITopo_rok into the BamHI/EcoRI sites of pFastBacRGSHis. pFastBacRGSHis_rokcat was transformed into DH10BAC cells and Baculovirus was produced according to the instructions of the manufacturer (Invitrogen). 11 of SF9 cells (10⁶ cells/ml) was inoculated at a MOI of 1 and Rokcat was expressed for 72 hrs at room temperature in a spinner flask. Cells were incubated in 40 ml lysis buffer (50 mM Tris, 150 mM NaCl, 1% glycerol, 3 mM β-mercaptoethanol, 10 mM imidazole, pH 8.0 supplemented with 1x Complete protease inhibitors (Roche) for 10 minutes on ice and sonicated. The lysate was centrifuged for 20 min at 10,000 g and the supernatant incubated with 2 ml 50% NiNTA agarose (Qiagen) for 1 hour. After 3 washes with 10 ml lysis buffer, the protein was eluted with 3 ml lysis buffer containing 300 mM imidazole and the eluate dialyzed.

Figure 8. Model of Cmb function. Based on the genetic interaction data we suggest that Rok may antagonize a positive role of Cmb on wing hair formation. Mwh is enriched on the proximal side of wing cells by the Fy/In group of PCP effectors where it prevents hair initiation. The suppression of the MHC phenotype of mwh null allele by a cmb null mutation further suggests that the role of Cmb on hair formation is antagonized by Mwh.

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in

mwh1

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Interestingly, PCP effector mutations generally enhance each other. For example, the hypomorphic frtz allele is enhanced by weak alleles of in or fy in double mutants [47]. Analogously, removal of a gene dosage of mwh in a fy or in background, enhances their MHC phenotype [50]. In contrast, the MHC phenotype of mwh mutants was (partially) suppressed in mwh cmb double mutants, as significantly fewer cells formed additional hairs. This interaction is likely specific, because we find it with the spontaneous mwh6 allele [18] and with the spontaneous mwh allele, two alleles of independent origin unlikely to carry a similar second site mutation and thus further supporting the physiological function of Cmb as a PCP effector. To our knowledge, cmb is the only gene reported so far to suppress mwh. Importantly, as mwh1 and cmb both are null alleles (cmbknockout lacks expression of both protein isoforms, Fig. 2; [23]), this result suggests that Mwh acts upstream of and normally antagonizes Cmb and that the derepression of Cmb thus may contribute to the MHC phenotype of mwh mutants (Fig. 8).

Unfortunately, our Cmb antibodies do not detect endogenous Cmb protein in the developing pupal wing. Nevertheless, Cmb expressed in the posterior compartment of the wing under the control of en-Gal4 localizes apically in a punctate pattern. In cells that appear to express at a lower level (seen particularly for Cmb-RA in Figure 7D, G) Cmb is enriched at the circumference of the cells, but shows no proximo-distal enrichment. Although we cannot exclude that Cmb localization is an overexpression artifact, this appears unlikely, because we would expect Cmb to fill the cells rather than to localize specifically apically (Fig. 7G). Importantly, Cmb likely localizes to the area of wing cells where Mwh is present, as Mwh known to be initially enriched apically towards the proximal side [21,23], further supporting our model that a positive effect of Cmb as a novel PCP effector on wing hair formation may be restricted by Mwh.

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against lysis buffer (without protease inhibitors andimidazole). Protein concentration was estimated to be 300 ng/μl using BSA standards on a Coomassie stained gel.

To generate the cmbKO knock-out allele, left and right homology arms were amplified by PCR from Bac clone RP80-17E13 (DGRC, CHORI, CA) using primers CG10732_left_for_KpnI, CG10732_left_rev_SacI and CG10732_right_for_BglIII, and CG10732_right_rev_AvrII, respectively and cloned into pSCA_KanAmp (Agilent). After sequence verification, the left arm was cloned as KpnI (blunt)/SacI fragment into the NotI (blunt)/SacI II fragment prPK2 [37], followed by insertion of the right arm as BglII/AvrII fragment into the corresponding sites to give prPK2 CG10732_fnal. The knock-out mutant of cmb was made according to [37]. The integrity of the cmbKO allele was verified by inverse PCR (not shown) and by PCR analysis using primer Cmb KO Verification F and Cmb KO Verification R (predicted product size 945 bp; Fig. 2A). To control for DNA integrity, primers EY10165_1 and EY10165_2 were used to amplify a fragment of CG7177 of 532 bp.

pGEX4T3_ES (Gst-ES) was generated by isolating the EcoRV/Sal fragment from pOT2_CmbRB (DGC clone GH01088) and cloning it into the Smal/Sall sites of pGEXT3. pGEX4T2_SX (Gst-SX) was made by isolating the SalI/XhoI fragment of pOT2_CmbRB and cloning it into the Sall site of pGEX4T2. pGEX4T2_BB (Gst-BB) was generated by isolating BamHI fragment (blunt) from pOT2_CmbRB and cloning it into pGEX4T_Smal vector.

pSCA-K/A-CmbRA_Nterm and pSCA-K/A-CmbRA_Cterm were made by cloning PCR products amplified from oligo-dT primed total ovarian cDNA using CG10732_RA_For and CG10732_RA_Nrev, and CG10732_RA_Cfor and CG10732_RA_rev, respectively, into pSCA-K/A-CmbRA (Agilent). pSCA-K/A-CmbRA was then assembled in a triple ligation from pSCA-K/A-CmbRA_Nterm (DraIII/EcoRV), pOT2_CmbRB (EcoRV/DraIII), and pSCA-K/A-CmbRA_Cterm (DraIII fragment; note that the different DraIII sites have different spacer sequences and could thus only ligate in one way). A gateway entry clone for CmbRB was made by inserting a XbaI (blunt)/BglII fragment of pSCA-K/A-CmbRA into the EcoRV/BamHI sites of pENTR_3C. A CmbRB entry clone was generated by cloning a PCR product amplified with CG10732_rarmLong_rev_AvrII, respectively and cloned into pENTR_3C. A CmbRB entry clone was then assembled in a triple ligation from pSCA-K/A-CmbRA_Nterm and pSCA-K/A-CmbRA_Cterm.

Site directed mutagenesis of the Rok phosphorylation sites of Cmb-RA with CG10732-NdeI-F1 and CG10732-XmaI-R1, and CG10732-NdeI-F2 and CG10732-XmaI-end-R2, respectively, as NdeI/XmaI fragments into pGADT7. pGBK7 constructs of mwh andfy were made by cloning PCR products amplified with DBD-mwh3 and DBD-mwh5, and pGBK7T7-3_Fy and pGBK7T7-3_Fy, respectively, as NdeI/EcoRI fragments into pGBK7T7. Analogously, fritz was cloned into the EcoRI site of pGBK7T7 after amplification using primers 5’ fritz-hybrid and 3’ fritz-hybrid, and in was cloned into NheI/BamHI of pGBK7T7 after amplification with Intron-th5 and Intron-th3.

**Antibody production and Western analysis**

Gst-ES was expressed and purified by the Macromolecular Therapeutics Development Facility of Einstein. Briefly, after purification over a Gsth-Sepharose column, Gst was cleaved off with His8-tagged Thrombin and further purified over a Ni-column and by size-exclusion chromatography. 100 μg ES fragment was injected per boost into Guinea pigs by Covance. For Western analysis, lysates of 7th instar w1118 and cmbKO larvae were prepared as the HEK293 cells for Co-IP were and separated on a 12% SDS PAGE gel, transferred to PVDF membrane and probed with #1051 anti Cmb at a dilution of 1:5,000. Signals were detected with a rabbit anti-Guniea pig antibody (Invitrogen) at a dilution of 1:10,000 and imaged after ECL treatment (Promega).

**Kinase assays**

For gel shift assays, 0.5 μl miniprep DNA of pOT2_CmbRB (DGC clone GH01088) were in vitro translated in the presence of [35] S-methionine in a 10 μl reaction using the TNT coupled transcription-translation system (Promega) according to the instructions of the manufacturer. The translation was then diluted with 40 μl 1.25x kinase buffer (5x kinase buffer: 125 mM HEPES pH 7.2, 15 mM MgCl2, 5 mM EDTA; unless phosphate assays were performed, 5x kinase buffer also contained 25 mM β-glycerophosphate and 5 mM Na3VO4). 5 μl of this dilution was incubated for 1 h with 5 μl 2x kinase mix (1 μl 5x kinase buffer, 1 μl 5 mM ATP, 0.1 μl 100x Cycloheximide (Sigma), 0.5 μl Rokcat and, if required, 10 u calf intestinal alkaline phosphatase (Roche). The reaction was stopped by the addition of 5 μl 5x SDS loading dye and boiling for 5 min at 95°C. Kinase reactions were separated on a 12% Anderson gel [28].

Gst proteins were expressed as described [40,41]. Radioactive kinase assays consisted of 250 ng of purified GST-tagged protein, 4 μl 5x Kinase buffer, 1.5 ul of 1 mM ATP, 0.5 ul of 32P γATP (3000 ci/mmol), 0.5 ul of Rokcat and, if required, 10 u calf intestinal alkaline phosphatase (Roche). Protein concentration was estimated to be 300 ng/μl using BSA standards on a Coomassie stained gel.

**Phylogenetic analyses**

Putative orthologs of Cmb were identified by BlastP searches of GenBank as well as other arthropod genome repositories (see Fig. S1) using the sequence of the Cmb-RA isoform. To deduce the phylogenetic position of cmb among putative heretofore unidentifed dipiteran cmb orthologs (see Fig. S2A and B), a multiple sequence alignment of Cmb-RA against the deduced amino acid sequences of Dipteran orthologs was performed using T-Coffee.
(v6.85) [56,57] using the following pair-wise alignment methods: the 10 best local alignments (Lalign_pair), an accurate global alignment (slow_pair) [56,57]. This alignment was used to construct a maximum likelihood tree using the PhyML program (v3.0 aLRT) [50] using the WAG substitution model assuming an estimated proportion of invariant sites (of 0.018) and 4 gamma-distributed rate categories to account for rate heterogeneity across sites, with the gamma shape parameter being estimated directly from the data (gamma = 1.306) [59]. Internal branch support was assessed using the aLRT test (SH-Like).

A similar strategy was used to deduce the phylogenetic position of cmb among other identified putative arthropod orthologs (see Figs. S1 and S2C) using the WAG substitution model assuming an estimated proportion of invariant sites (of 0.007) and 4 gamma-distributed rate categories to account for rate heterogeneity across sites [59]. The gamma shape parameter was estimated directly from the data (gamma = 2.195). Reliability for internal branch was also assessed using the aLRT test (SH-Like). All phylogenetic analyses were performed using Phylogeny.fr [60]. All trees were edited and visualized using Geneious [61]. All alignments are available by request.

Protein interactions

Two-hybrid assays were performed using the Clontech system according to the instructions of the manufacturer (see also [40]).

Co-immunoprecipitations were done with a modified version of the technique used in [41]. Briefly, HEK293 cells were transfected with 5 μg of pCS3_GFP_CmbRA, pCS3_GFP_CmbRB, pCS3_GFP_CmbInt or pCS3_GFP_TBX1 together with 5 μg pCSMyc6_Mwh or pCS3_Myc_Dazap1 using Polyethylenimine (PEI). The cells were collected after 48 hours and washed with ice cold 1xPBS and lysed in 500 ul of Buffer A (20 mM Tris, (PEI). The cells were collected after 48 hours and washed with ice cold 1xPBS and lysed in 500 ul of Buffer A (20 mM Tris, 1% Nonidet-P 40, 1 mM EDTA, 1 mM EGTA and 1:200 concentration of 1 mM Benzamidine, 10 mM Leupeptin, and 1 μM Pepstatin) and rotated at 4°C for 15 minutes. Lysates were centrifuged at 1000 g for 15 minutes two times with the supernatant being retained after each step. The supernatants were incubated with 1 ug of mouse anti-GFP antibody (Roche) for four hours at 4°C. 30 ul of 50% Protein G beads were added and followed by overnight incubation at 4°C. Immunoprecipitates were washed 3x with cold Buffer A and suspended in 15 ul of 2x Laemmli buffer. Samples were run on a 12% SDS gel and western blot analysis was performed using standard protocols.

Immunohistochemistry

Flip-out clones were generated using actin>stop>Gal4 with UAS-GFP as lineage tracer. 30 and 36 hpf Pupal wing discs were dissected and stained by following a standard procedure [20]. The monolayers were fixed in 4% Paraformaldehyde, washed with 1xPBS, incubated with the primary antibody overnight at 4°C, rinsed with 1xPBS/0.3% TritonX100, and incubated with a fluorescently labeled secondary antibody, washed and mounted in Prolong gold antifade reagent. For the analysis of the 10 best local alignments (Lalign_pair), an accurate global alignment (slow_pair) [56,57]. This alignment was used to construct a maximum likelihood tree using the PhyML program (v3.0 aLRT) [50] using the WAG substitution model assuming an estimated proportion of invariant sites (of 0.018) and 4 gamma-distributed rate categories to account for rate heterogeneity across sites, with the gamma shape parameter being estimated directly from the data (gamma = 1.306) [59]. Internal branch support was assessed using the aLRT test (SH-Like).

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Supporting Information

Figure S1  Summary and accession numbers of sequences used in alignments and phylogenetic analyses. (TIF)

Figure S2  Cmb phylogenetic analyses. (A) Maximum likelihood tree of the deduced amino acid sequence of D. melanogaster Cmb to other putative Dipteran Cmb proteins showing that Cmb is highly conserved within the Diptera (including flies and mosquitoes). Multiple, lineage-specific cmb duplications may have occurred in the mosquitoes, in that these mosquito cmb orthologs did not resolve into paralog-specific clades. However, the results of this analysis do suggest that the common ancestor of the Anopheles lineage had duplicate cmb paralogs with the most parsimonious explanation being that A. darlingi lost the paralog of A. gambiae AFAP012311. The relationships among other mosquito cmb genes is less clear and may be due the branch length (65). (B) Drosophila portion of the tree in (A) showing the relationship of the deduced amino acid sequences of putative Drosophilid cmb orthologs. (C) Maximum likelihood tree showing the relationships between the deduced amino acid sequences of arthropod Cmb orthologs. A DELTA-BLAST of Cmb indicated that it has partial sequence similarity to SMC (Stability of Mitotic Chromosomes) proteins. The Cmb group clusters as a sister-group to the SMC2 protein group. aLRT test (SH-Like) support at each node is indicated as a percent. The scale bars represent the number of amino acid substitutions per site. All gene numbers were retrieved from the repositories listed in Figure S1. (TIF)

Figure S3  Multiple sequence alignment of the putative amino acid sequences of arthropod Cmb orthologs. All aligned sequences are listed in Figure S1. Sequence logos are shown above each aligned site. Sequences were aligned using the T-COFFEE algorithm (see Materials and Methods). (PNG)

Figure S4  FASTA file of alignment shown in Fig. S3. (FASTA)

Figure S5  Loss or gain of cmb does not cause a PCP phenotype in the eye. Tangential sections of adult eyes with corresponding schematic representation of ommatidial orientations underneath. Black and red arrows represent dorsal and ventral chiral forms of ommatidia. (A) Wild-type. (B) A homozygous cmbKO mutant eye shows no PCP phenotype. (C, D) Eyes overexpressing cmb-RA (C) or cmb-RRB (D) under the control of the ski-Gal4 at 29°C are wild-type. (TIF)

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Author Contributions

Conceived and designed the experiments: JKF GD QL AB JPJ AS CP PA AJ. Performed the experiments: JKF GD QL AB JPJ AS CP PA AJ. Analyzed the data: JKF GD QL AB JPJ AS PA Aj. Contributed reagents/materials/analysis tools: JPJ AS. Contributed to the writing of the manuscript: JKF PA AJ.

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