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Rho-kinase planar polarisation at tissue boundaries depends on phospho-regulation of membrane residence time

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Summary

The myosinII activator Rho-kinase (Rok) is planar-polarised at the tissue boundary of the *Drosophila* embryonic salivary gland placode through a negative regulation by the apical polarity protein Crumbs that is anisotropically localised at the boundary. However, in inner cells of the placode both Crumbs and Rok are isotropically enriched at junctions. We propose that modulation of Rok membrane residence time by Crumbs’ downstream effectors can induce both behaviours. Using FRAP combined with *in-silico* simulations, we find that the lower membrane dissociation rate ($k_{off}$) of Rok at the tissue boundary with low Crumbs explains this boundary-specific effect. The S/T-kinase Pak1, recruited by Crumbs and Cdc42, negatively affects Rok membrane association *in vivo*, and *in vitro* can phosphorylate Rok near the PH-domain that mediates membrane association. These data reveal an important mechanism of modulation of Rok membrane residence time via affecting the $k_{off}$ that may be widely employed during tissue morphogenesis.
Introduction

Tissues arise during development through specification of primordia that will then initiate morphogenetic movements (Castelli-Gair Hombria, 2016). Many primordia are epithelial in nature and give rise to tubular epithelial organs, such as lung, kidney, intestine in vertebrates or the equivalent organs in invertebrates. How are epithelial primordia physically set aside from the surrounding tissue, apart from the inductive change in transcription factor expression? We know from studies in the *Drosophila* early embryonic epidermis as well as in larval wing discs that differently fated compartments are physically clearly segregated and cell mixing across compartment boundaries is restricted (Dahmann and Basler, 1999; Tepass et al., 2002). In both tissues, this seems to be in part achieved through an increased tension at the compartment boundary that coincides with junctional accumulation of actomyosin into a seemingly supra-cellular structure, a so-called actomyosin cable (Röper, 2013). In the embryonic epidermis, the actomyosin cables found at parasegmental boundaries physically restrain boundary-challenging divisions within the correct compartment (Monier et al., 2010). The molecular mechanisms that drive actomyosin cable assembly are not well understood, and where aspects have been uncovered, a variety of tissue-specific mechanisms seem to contribute. Parasegmental cables for instance arise over time from dorso-ventrally-polarised junctional myosin accumulations (Tetley et al., 2016) that might themselves depend on a code of Toll receptor expression within the early epidermis (Pare et al., 2014). In the wing disc, the dorso-ventral boundary requires Notch-signalling (Major and Irvine, 2005, 2006). Actomyosin-based compartment boundaries are not restricted to invertebrates, but have in fact also been found in vertebrates, with key examples being the rhombomere boundaries in the mammalian hindbrain (Calzolari et al., 2014) as well as the neural plate-ectoderm boundary during neurulation (Galea et al., 2017).

We have previously identified that an actomyosin cable is positioned at the boundary of the salivary gland placode in the *Drosophila* embryo (Röper, 2012). Two epithelial placodes of about 100 cells on either side of the ventral midline become specified at stage 10 of embryogenesis and will invaginate to form the two salivary glands ((Girdler and Röper, 2014; Sidor and Röper, 2016); Fig. 1A,A'). This invagination process is driven by a combination of isotropic apical constriction, mediated by dynamic pools of apical-medial actomyosin, and directed cell intercalation, mediated by apical junctional pools of actomyosin (Booth et al., 2014; Sanchez-Corrales et al., 2018). The placodes are contained within parasegment 2 of the embryo, and remnants of previous parasegmental actomyosin cables are
specifically retained near the ventral portion of the placode, whilst a new section forms around the dorsal part, so that by mid stage 11 a circumferential actomyosin cable surrounds each placode (Röper, 2012). The formation of this cable is transcriptionally initiated, as it is lacking in mutants for the most upstream specifying transcription factor Sex Combs Reduced (Henderson and Andrew, 2000; Röper, 2012). We have previously identified the apical transmembrane protein Crumbs as a key determinant of actomyosin cable positioning at the boundary of the placode. Crumbs levels are transcriptionally upregulated within the placode, whereas levels are reduced in the epidermal cells surrounding the placode (Fig. 1C” and Fig. 2A) (Kerman et al., 2008; Myat and Andrew, 2002). Crumbs’ ability to form homophilic interactions (Fletcher et al., 2012; Röper, 2012; Zou et al., 2012) leads to a highly anisotropic distribution of Crumbs apically within placodal cells at the placode boundary: this Crumbs anisotropy leads to accumulation of actomyosin and Rho-kinase (Rok) reporters at the placode boundary, away from membranes with high levels of Crumbs. Moreover, ectopic boundaries of high/low Crumbs levels lead to myosin cable formation (Röper, 2012). Thus, Crumbs exerts a negative regulatory effect on actomyosin cable formation, likely through the upstream myosin activator Rho-kinase. Based on studies in mammalian cells that reported that the kinase aPKC is able to phosphorylate and thereby inactivate Rok (Ishiuchi and Takeichi, 2011), we previously suggested that the effect in the placode was mediated by aPKC, which in fly embryos closely follows Crumbs distribution (Röper, 2012). Complementary localisation of Crumbs and actomyosin is not restricted to flies, but has also more recently been reported in mouse (Ramkumar et al., 2016).

The negative influence of Crumbs on Rok localisation and activity is clear at the placode boundary where Crumbs anisotropy triggers the actomyosin cable assembly in boundary junctions with lower levels of Crumbs. However, in the inner placodal cells, where Crumbs is high due to transcriptional control and Rok is required for the actomyosin-dependent invagination process, both Crumbs and Rok membrane levels are high and isotropic. Thus, it appears that it is not the overall levels of Crumbs but rather the difference in levels experienced within a single cell at the boundary that allows negative regulation of Rok. This is also supported by the fact that introduction of a new step change in Crumbs levels within the placode, now adding a stripe of Crumbs expression even higher than the already elevated placodal levels, triggers Crumbs anisotropy and ectopic myosin cable formation (Röper, 2012). Here we propose a molecular mechanism by which these different scenarios can be reconciled, based on a modulation of Rok residence time downstream of Crumbs at the apical junctional plasma membrane, mediated by modulation of the $k_{off}$
through phosphorylation of Rok. We suggest that such mechanism might be widely used in membrane receptors to allow combination of patterning activity with other molecular functions.
Results

*Rok is planar polarised and complementary to Crumbs at the placode and other tissue boundaries*

We previously proposed that the apical transmembrane protein Crumbs actively influences myosin II accumulation through a negative regulatory effect on Rok, the most common activator of myosin II during morphogenetic processes (Röper, 2012). However, current visualisation of Rok localisation and activity depends on over-expression of tagged active or kinase-dead versions of the protein (Simoes Sde et al., 2010). In order to examine endogenous Rok localisation, we used CRISPR/Cas9 to engineer an endogenously N-terminally tagged Rok with the bright mNeon-Green (mNG) fluorescent protein, tagging major isoforms A, B and C ((Shaner et al., 2013); Fig. 1B). Isoform D has an alternative start but it is not expressed during embryogenesis (mod et al., 2010). Importantly, mNG-Rok did not form aggregates like many of the overexpression lines (see Fig.4B) and mNG-Rok flies were homozygous viable, indicating that Rok function was not impaired by the mNG-tag.

In epithelial cells of the epidermis, as well as in larval imaginal discs, endogenous mNG-Rok localised similarly to previously described tagged Rok, being enriched both in the apical-junctional as well as in the apical-medial region of certain cells (Fig. 1C-H). At several boundaries of differently fated epidermal domains, such as in the embryo the salivary gland placode (Fig. 1C), posterior spiracles (Fig. 1 D, E), ventral midline, (Fig. 1F), epidermal leading edge/amnioserosa interface (Fig. 1G), and larval eye discs (Fig. 1H, I), mNG-Rok was strongly enriched at the boundary, as was the downstream morphogenetic effector myosin II (Figure S1; Jacinto et al., 2002; Röper, 2012). These were also all boundaries where Crumbs levels show a clear step change of high versus low expression.

*Crumbs induces Rok planar polarisation at the placode boundary*

In the salivary gland placode, mNG-Rok was enriched apically in junctions in comparison to the surrounding epithelium (Fig. 1C and Fig. 2A-A”). However, in placode boundary cells, where Crumbs protein is highly anisotropically localised within the sub-apical domain (Fig. 2A, arrow, and 2C), apical junctional Rok was planar polarised. In these boundary cells, mNG-Rok accumulated in junctions with lower levels of Crumbs, the boundary junctions, and appeared depleted from junctions with high Crumbs levels, the side junctions (Fig. 2A-A” and C), suggesting that Crumbs negatively regulates endogenous Rok membrane accumulation. In order to examine the effect of Crumbs on endogenous Rok membrane localisation, we
induced Crumbs overexpression in a stripe of cells in mNG-Rok embryos using the en-Gal4 driver (UAS/Gal4 system; (Brand and Perrimon, 1993)). This ectopic stripe of higher Crumbs levels created a new high/low boundary of Crumbs protein within the salivary gland placode (Fig. 2B, arrow). At the new boundary Crumbs localisation was highly anisotropic, with Crumbs enriched in junctions with neighbouring cells that also expressed high levels of Crumbs, and lower at junctions with the surrounding placode cells (Fig. 2 B). Again, only within the cells at the stripe boundary, mNG-Rok was planar polarised and enriched at the new ectopic boundary junctions with lower levels of Crumbs, and it was depleted from junctions with higher levels of Crumbs (Fig. 2 B,"B"). This negative regulatory effect was also visible within the apical-basal extent of the epidermal cells, as an expansion of the apico-lateral distribution of Crumbs due to ectopic overexpression lead to a basal shift in mNG-Rok localisation within the lateral membrane (Figure S2). Thus, Crumbs negatively regulates endogenous Rok membrane association.

A model to explain Rok polarisation at the boundary via modulation of Rok dynamics
We were intrigued by the fact that the negative regulatory effect of Crumbs appeared to only affect Rok in cells at the boundary of the salivary gland placode, but not all throughout the tissue. In inner placodal cells, where Crumbs was highly enriched isotropically at all sub-apical junctions, mNG-Rok was able to localise at apical junctions despite the high Crumbs levels (Fig. 2A-A", C). In contrast, in boundary cells where Crumbs localisation was highly anisotropic, mNG-Rok was planar polarised. Furthermore, an ectopic boundary induced by Crb overexpression could also polarise Rok (Fig. 2B-B"). Thus, it was the anisotropic distribution of Crumbs within a cell, rather than its absolute levels in the plasma membrane, that affected Rok membrane localisation (Fig. 2C).

One way to mechanistically reconcile these two situations (boundary versus inner placode) is to consider the dynamics of Rok membrane localisation (Fig. 2D). If we assume that at equilibrium Rok is able to associate and dissociate from the cell membrane at specific rates, Crumbs could influence Rok membrane accumulation by selectively increasing the Rok membrane dissociation rate ($k_{on}$). We propose that within inner placodal cells, high levels of Crumbs in all apical junctions would lead to a higher, though isotropic, Rok turnover. With Rok membrane recruitment, i.e. $k_{on}$, remaining unaffected (in inner cells just as in boundary cells), Rok would still localise at the plasma membrane of these inner cells. Within the boundary cells, Rok would dissociate more often from junctions with high levels of Crumbs (side junctions) and
would therefore accumulate at junctions with lower levels of Crumbs and lower $k_{\text{off}}$ (boundary junction), resulting in the planar polarisation of Rok at this tissue boundary.

**Endogenous Rok $k_{\text{off}}$ is lower at the boundary of the placode and leads to planar polarisation of Rok in simulations**

The above model clearly predicts that Rok’s mobility and in particular Rok’s $k_{\text{off}}$ at the boundary junctions should differ from Rok’s $k_{\text{off}}$ at other junctions within the placode, both the side junctions of a boundary cell as well as the inner junctions of a placode inner cell (with both types of junctions showing the same high levels of Crumbs). To test this prediction, we performed Fluorescence Recovery After Photobleaching (FRAP) experiments in stage 11-12 embryos expressing the endogenously tagged mNG-Rok (Fig. 3A,B). We bleached small circular regions of mNG-Rok at the apico-lateral plasma membrane and imaged at ~500-700ms intervals to capture the mNG-Rok fluorescence intensity pre- and post-bleach. Supporting our predictions, mNG-Rok located at junctions within the placode, both inner junctions within the centre of the placode as well as side junctions of boundary cells, recovered significantly faster than mNG-Rok located at the boundary (Fig. 3B-D). As demonstrated in Supplemental Information, given our experimental settings, the recovery rate can be interpreted as a $k_{\text{off}}$ of a reaction-diffusion equation (Bulinski et al., 2001; Sprague et al., 2004). $k_{\text{off}}$ values were estimated from fluorescence recovery as 0.148 (+/- 0.018) s$^{-1}$ for the inner placodal cells, 0.173 (+/- 0.024) s$^{-1}$ for side membranes of boundary cells and 0.057 (+/- 0.008) s$^{-1}$ for the boundary membrane of boundary cells. Estimated $k_{\text{off}}$ values were found to be significantly lower in boundary junctions compared to junctions within the placode, both side and inner ones (p-values [bootstrap/boxplot]: 0.0002; Fig. 3C,D). Thus, the difference in dynamic behaviour of mNG-Rok at junctions forming the boundary of the placode compared to other junctions within the placode supports the model for Rok planar polarisation presented above.

In order to assess whether the difference in Rok’s $k_{\text{off}}$ measured at the placode boundary and within the side/inner junctions of the placode was in itself sufficient to produce planar polarisation of Rok at the tissue boundary, we developed an *in silico* simulation of the process, using particle-based stochastic reaction-diffusion (Fig. 3E,F; Suppl.Fig. 3; for details of the implementation see STAR Methods). We tested a variety of $k_{\text{off}}$ combinations for the boundary and side/inner placodal cell membranes (keeping the $k_{\text{on}}$ constant; Suppl.Fig. 3). Uniform $k_{\text{off}}$ values in all membranes led, as expected, to isotropic membrane accumulation of Rok (Fig.
3G, simulation 1). Setting a high $k_{off}$ in inner junctions compared to boundary junctions led to a strong Rok polarisation at the boundary, but markedly reduced Rok membrane localisation in inner cells (Fig. 3G, simulation 2). By contrast, $k_{off}$ values deduced from the in vivo FRAP measurements of mNG-Rok were sufficient to trigger Rok planar polarisation in simulated boundary cells, at a level similar to the polarisation measured in vivo (in silico polarisation value of 1.48, Fig. 3G, simulation 3; compare to values for mNG-Rok in vivo in Fig. 6E and Fig. 7G). Under these conditions, Rok accumulation at the membrane of inner cells with higher $k_{off}$ was preserved, leading to a pattern that closely resembled Rok localisation in the salivary gland placode in vivo (Fig. 1C’ and 2 A’).

Thus, modulation of Rok membrane $k_{off}$ within the placode is sufficient to elicit Rok planar polarisation specifically at the tissue boundary whilst preserving Rok membrane accumulation in the rest of the tissue. We next investigated the molecular basis for this mechanism.

**The Rok C-terminal region is required for its planar polarisation**

*Drosophila* Rok is a large protein of 1391 amino acids, containing an N-terminal kinase domain followed by a coiled-coil region, and a C-terminal Shroom-binding domain (SBD), Rho-binding domain (RBD) and PH-domain (PH) (Fig. 4A)(Amano et al., 2010; Simoes Sde et al., 2014). In order to identify which of these domains were required for apical junctional planar polarisation, truncated versions of a Venus-tagged kinase-dead Rok localisation reporter were overexpressed in embryos using the UAS/Gal4 system under the control of Daughterless-Gal4, an early ubiquitous zygotic driver (Simoes Sde et al., 2014)

The Venus-Rok[K116A]-ΔSBD lacking the coiled-coil/Shroom-binding domain was able to accumulate in apical membranes (at lower levels than the control Venus-Rok[K116A]) and could still polarise at the placode boundary (Fig. 4B vs C, and F’). Venus-Rok constructs lacking the Rho-binding domain, Venus-Rok[K116A]ΔRBD, or the PH-domain, Venus-Rok[K116A]ΔPH, although showing a strongly reduced overall membrane accumulation, were able to localise at low levels to the membrane. Both constructs failed to polarise at the tissue boundary (Fig. 4D-E’, and F’). Thus, in addition to promoting Rok apical junctional membrane recruitment, both the Rho-binding domain and the PH domain are important for Rok planar polarisation in boundary cells.

Rok membrane recruitment has been shown to be dependent on the small GTPase Rho and its exchange factor RhoGEF2 (Amano et al., 2010; Mason et al., 2016; Nakamura et al., 2017). In order to assess whether Rok planar polarisation is
caused by planar polarisation of these upstream activators, we examined the localisation of an sfGFP tagged version of RhoGEF2 (Sarov et al., 2016) and the GFP tagged Rho-binding domain of Anillin (AniRBD-GFP), a reporter for activated GTP-bound Rho (Munjal et al., 2015). Neither RhoGEF2-sfGFP nor AniRBD-GFP showed any apical junctional planar polarisation within the boundary cells of the salivary gland placode (Fig. 4 G-I). Thus, Rok planar polarisation is not driven by an upstream polarisation of active Rho.

These data suggest that, although Rho-binding is important for Rok membrane recruitment, once at the membrane Rok planar polarisation is regulated by a Rho-independent mechanism involving the Rok C-terminal membrane association region.

**Phosphorylation of the Rok C-terminal region by Pak1 regulates Rok membrane association**

Alternatively, regulation of Rok membrane association could be achieved via modulation of Rok’s binding to Rho or to phospholipids, for instance through phosphorylation in the vicinity of Rok’s RBD or PH domain. Phosphorylation near phospholipid-interacting sequences in several Par-complex substrates has recently been suggested to inhibit their membrane binding (Bailey and Prehoda, 2015). We had previously suggested that Crumbs could negatively regulate Rok through one of its downstream interactors, aPKC (Fig. 5A; (Röper, 2012)), which binds to the Crumbs intracellular domain through its binding partner Par6 (Bulgakova and Knust, 2009). Previous data from mammalian tissue culture cells revealed that aPKC could phosphorylate human ROCK1, one of the two mammalian Rho-kinases, near the RBD and PH-domains, and a phospho-mimetic version of ROCK1 showed loss of plasma membrane association (Ishiuchi and Takeichi, 2011). Furthermore, the p21-activated kinase 1 (Pak1), which is activated by the Par6 binding protein Cdc42, has recently been shown to act semi-redundantly with aPKC to phosphorylate shared target proteins (Fig. 5A; (Aguilar-Aragon et al., 2018; Bokoch, 2003)). Importantly, several putative phosphorylation sites for both Pak1 and aPKC are highly conserved between *Drosophila* and mammalian Rho kinases (Blom et al., 1999; Blom et al., 2004; Rennefahrt et al., 2007; Fig. 5B, Suppl. Fig. 4). We therefore set out to examine whether aPKC and/or Pak1 contributed to a regulation of Rok membrane association.

We overexpressed membrane targeted versions of aPKC or Pak1 in *enGal4* stripes in embryos with endogenously tagged mNG-Rok. While overexpression of aPKC using *UAS-aPKC[CAAX]* did not significantly affect mNG-Rok in the embryonic
epithelium in comparison to control cells (Fig. 5C-"C"), overexpression of Pak1 using UAS-Pak1\textsuperscript{mmy} strongly decreased mNG-Rok membrane localisation (Fig. 5D-D").

Mass-spectrometric analysis of mammalian EpH4 tissue culture cell lysates performed by (Ishiuchi and Takeichi, 2011) identified nine phosphorylated sites in human ROCK1, four of which are conserved in Drosophila Rok. Strikingly, all four sites are located in the Rok C-terminal region, close to the RBD and PH domains (Fig. 5B and Fig.S4). Furthermore, all are recognised as putative phosphorylation substrates for Pak1 and aPKC (Fig. 5B; Fig. S4). We designed three short peptides covering these putative sites and performed \textit{in vitro} kinase assays with the purified kinase domains of human Pak1 and human PKCi, using a small Bazooka peptide (BazS980) as a positive control (Fig. 5E). Both Pak1 and aPKC kinase domains strongly phosphorylated the control Bazooka peptide, but no aPKC phosphorylation of any of the Rok peptides was detected. By contrast, Pak1 strongly phosphorylated Rok peptide 2 (Rok Pep2), containing serine S1365 and threonines T1366 and T1369, which are all located close to the C-terminal end of the PH domain (Fig. 5B, E). This phosphorylation was completely abolished in a peptide that had these three residues, S1365, T1366 and T1369, mutated to alanine (Fig. 5E; Rok pep2AAA).

Thus, Pak1 negatively regulates Rok membrane accumulation by phosphorylating its C-terminal region.

\textbf{Phosphorylation of Rok contributes to its planar polarisation in vivo}

To confirm that phosphorylation of Rok by Pak1 played a role in its planar polarisation \textit{in vivo}, we used the previously generated mNG-Rok strain to mutate the four conserved putative phosphorylation sites described above, as well as eight serines and threonines in close proximity (Fig. 5B, asterisks) using CRISPR/Cas9 and homologous recombination repair (Fig. 6A).

Although flies carrying this mNG-tagged phospho-site-mutant Rok (mNG-Rok[Pmut]; Fig. 6 B vs C) were homozygous viable, a two-fold increase in embryonic lethality (12.1% of fertilised embryos) was observed in comparison to the parental mNG-Rok strain (5.9% of fertilised embryos) (Fig. 6D). There was no increased lethality at later developmental stages. We analysed planar polarisation of mNG-Rok[Pmut] in comparison to mNG-Rok by quantifying fluorescence intensity ratio between boundary junctions and side junctions in boundary cells that showed a clear Crumbs anisotropy (Fig. 6 E-G). Whereas, in agreement with quantifications shown in Fig. 7E, mNG-Rok showed a clear polarisation at the boundary, with a mean ratio of 1.36, mNG-Rok[Pmut] was on average not polarised, with a mean ratio of 0.94.
(Fig. 6G). This suggests that phosphorylation of these consensus aPKC/Pak1 sites contributes to Rok planar polarisation.

We could not detect any phenotype in tissue bending and salivary gland placode invagination at early stages, i.e. stages 11-12, in mNG-Rok[Pmut] embryos, but could detect a fraction of embryos beyond stage 13 (23.3% in mNG-Rok[Pmut] compared to 0.0% in the mNG-Rok control) that showed a striking phenotype: we observed a disruption and altered cell shapes within the epidermis at the positions where the salivary gland placodes were located (Fig. 6 H-J, arrows). The rest of the epidermis appeared largely unaffected. At this stage in control embryos, the circumferential actomyosin cable at the placode boundary was strongly enriched in F-actin (Fig. 6 H,H'', arrows; (Röper, 2012)). By contrast, in mNG-Rok[Pmut] embryos we did not detect any accumulation of F-actin at comparable junctions where the cable would be positioned in the wild-type (Fig. 6 I,I''). mNG-Rok[Pmut] was still localised to junctions in the placode area but there was no accumulation suggesting cable localisation at this stage (Fig. 6 I').

Thus, phosphorylation of the Rok C-terminal domain, which appears to be mediated by Pak1, is required in vivo for Rok planar polarisation.

**Pak1 regulates Rok planar polarisation downstream of Crb/Cdc42**

Although zygotic loss of Pak1 is lethal, maternal contribution allows embryos to develop normally as far as early stage 11, making it possible to investigate Pak1’s role during salivary gland placode morphogenesis. Interestingly, Pak1 zygotic loss of function has been shown to induce defects in embryonic dorsal closure as well as in the late embryonic salivary glands (Bahri et al., 2010; Conder et al., 2007; Pirraglia et al., 2010). In early stage 11 pak1'14 zygotic mutant embryos, Pak1 was still detectable in the epithelium and early salivary placodes were not affected. However, in later placodes from late stage 11/stage 12 onwards mNG-Rok planar polarisation at the salivary gland boundary was strongly reduced (Fig. 7A-B'). Moreover, UAS-Pak1myr overexpression in the salivary placode using fkh-Gal4 (Henderson and Andrew, 2000) completely abolished mNG-Rok planar polarisation at the boundary most obvious at later stages, when the placode seemed to pull away from the surrounding epithelium (Fig. 7D). These results show that Pak1 plays a crucial role in salivary gland morphogenesis by regulating Rok membrane localisation and planar polarisation.

To understand how Crumbs might regulate Pak1 to control Rok planar polarisation, we examined Pak1 protein localisation in the embryonic epidermis. Like Crumbs protein, Pak1 was localised apically at junctions and was enriched in the
salivary placode (Fig. 7F-F’). Pak1 function depends on its activation by the small GTPase Cdc42 (Bokoch, 2003). An mCherry-Cdc42 reporter expressed under the control of the sqh promoter localised apically in junctions in the embryonic epithelium and showed anisotropic distribution at the salivary gland placode boundary similar to Crumbs (Fig. 7G-G” and quantified in 7J). Ectopic Crumbs expressed in enGal4 stripes also recruited this Cdc42-reporter to ectopic locations (Fig. 7H-H’’). In order to confirm that Cdc42 activated Pak1 in the embryonic epidermis we examined the localisation of Pak1 in cells expressing constitutively active Cdc42\textsuperscript{V12} (Welch et al., 1998). In stripes of cells expressing Cdc42\textsuperscript{V12} Pak1 localisation to junctions was strongly enhanced (Fig. 7I-I’’, K). Thus, the negative regulatory effect of Crumbs on Rok is likely mediated by the recruitment and activation of Pak1 by Crumbs-bound Cdc42.

We then wanted to directly test the effects of zygotic loss or overexpression of Pak1 on Rok dynamics at the membrane. To do so, we measured mNG-Rok recovery upon photo-bleaching (as above) in the salivary gland placode of pak\textsuperscript{14} mutant embryos as well as in embryos overexpressing membrane-targeted Pak1 (using UAS-Pak1\textsuperscript{myr} x fkhGal4). Zygotic loss of Pak1 slowed the recovery of mNG-Rok in both boundary and inner junctions, whereas placodal expression of myristoylated Pak1 increased it (Fig. 7 L, M). mNG-Rok $k_{\text{off}}$ values estimated from fluorescence recovery in pak\textsuperscript{14} placodes (boundary junctions: 0.032 (+/- 0.013) s\textsuperscript{-1}, inner junctions: 0.044 (+/- 0.008) s\textsuperscript{-1}) were significantly lower than in control placodes (p-value [bootstrap/boxplot]: 0.0002 and 0.008, respectively), supporting our hypothesis that Pak1 promotes Rok membrane dissociation. Corroborating this data, mNG-Rok $k_{\text{off}}$ values in placodes overexpressing Pak1\textsuperscript{myr} were significantly increased compared to the original control, with values of 0.170 (+/- 0.022) s\textsuperscript{-1} at the boundary and 0.266 (+/- 0.025) s\textsuperscript{-1} within the placode (p-value [bootstrap/boxplot]: 0.0002 for both; Fig. 7M).

Thus, Pak1 is able to modulate Rok membrane dynamics. Altogether, our data demonstrate that Pak1 is a direct modulator of Rok dynamics responsible for Rok planar polarisation downstream of Crumbs at the boundary of the salivary gland placode.
Discussion

In epithelial cells, junctional proteins as well as morphogenetically active pools of actomyosin are concentrated within the apical and apico-lateral region of the cells. This placement is controlled by the epithelial polarity network (Tepass, 2012). Patterning of cytoskeletal activity and junctional changes, key ingredients of morphogenetic changes in epithelial tissues, then takes place within this apical domain. This can lead to apical junctional planar polarisation of whole tissues or smaller domains or rows of cells that are morphogenetically active, as for instance is the case at the boundary of the salivary gland placode. In this case, the apical polarity determinant Crumbs plays a dual role, firstly in maintaining apical-basal polarity of epidermal cells including the salivary gland placode, and secondly in patterning cytoskeletal behaviour within the apical domain. Crumbs levels, in fact, show dynamic variations across much of the Drosophila embryonic epidermis until stage 14, and as described here, step changes in Crumbs levels tend to be accompanied by actomyosin accumulation at these boundaries (Röper, 2013).

Rok as the key morphogenetic activator of non-muscle myosin II is crucial to development in many animals. Thus, regulation and activity of Rok in cells is closely controlled. Historic views of Rok activation assumed a potential fold-back mechanism whereby the known regulatory activity of Rok’s C-terminus would contact the N-terminal kinase domain and block its activity (Amano et al., 1999; Julian and Olson, 2014). An alternative view is supported by recent evidence and suggests that the C-terminal domain is crucial for membrane interaction and that Rok is in fact always found as a homodimer in an extended conformation (Truebestein et al., 2015; Truebestein et al., 2016). Such an extended conformation and role of the C-terminus in membrane binding is in agreement with our in vivo findings that phospho-regulation of this domain is critical for membrane localisation.

aPKC and Pak1 are both important kinases with a multitude of roles in development and tissue homeostasis (Hong, 2018; Rane and Minden, 2014). Their overlapping function though has only been appreciated recently (Aguilar-Aragon et al., 2018). Due to the recently published data on Pak1 and aPKC overlapping function, we were prompted to identify Pak1 as the main effector downstream of Crumbs responsible for Rok planar polarisation at the placode boundary. But why ‘charge’ the Crumbs intracellular domain with two kinases with highly overlapping targets? The fact that Pak1 depends on Cdc42 for its activation might add another layer of control and allow differential kinase usage or amplification of kinase activity depending on the tissue context.
The differential expression level of Crumbs between salivary gland placode and surrounding epidermis, or epidermis and amnioserosa, demarcates clear boundaries. These boundaries at the plasma membrane level can then be turned into cytoskeletal planar polarisation, leading to physical boundaries due to for instance increased tension. Crumbs is not the only homophilic interactor that can exert such effects. Recent examples include E- and N-Cadherin patterning during eye morphogenesis in the fly (Chan et al., 2017), the fly nectin Echinoid at clonal boundaries (Chang et al., 2011), as well as tissue-specific expression of a Cadherin, Cad2, selectively in the neural cells in Ciona robusta, thereby patterning myosin activity at the neural/epidermal boundary where there is a step change in Cad2 expression (Hashimoto and Munro, 2018). Interestingly, in Ciona Cad2 is also titrated away from the tissue boundary due to homophilic interactions and also exerts a negative regulatory effect on myosin II accumulation. In this context though, and in contrast to the salivary gland placode boundary, it is RhoA activity that is polarised and increased at the tissue boundary, due to a selective recruitment of a RhoGAP by Cad2 (Hashimoto and Munro, 2018). Thus, although common principles in patterning tissue boundaries and patterning of cytoskeletal activity are repeatedly used in development, the fine molecular details vary depending on the tissue context.

We identified a mechanism that allows for a boundary-specific regulation of Rok and myosin accumulation at membranes between differently fated groups of cells. This mechanism only occurs when a pre-existing molecular anisotropy of an upstream factor is detected, thereby circumventing an on/off negative regulatory interaction. In the Ciona example above, Cad2 levels are high in constricting neural cells that internalise into the embryo, very reminiscent of the situation in the salivary gland placode. Thus, the effect of Cad2 on myosin is not absolute here either. We suspect that such a mechanism of modulation of membrane residence time via affecting the $k_{off}$ might be widely employed during tissue morphogenesis, as it allows for fine-tuning of cytoskeletal activity being integrated with other essential cell biological functions of the upstream regulators. In the case of Crumbs, its crucial role in maintaining apical-basal polarity and thereby epithelial integrity can be combined with its planar tissue patterning role.

In summary, this example of planar patterning in a morphogenetic process illustrates that the study of such processes needs to take the dynamic behaviour of components into account. The current exciting advances in light microscopy, live imaging and image quantification will be of crucial help to facilitate and transform such analyses in live tissues during development.
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Author contribution

Declaration of Interests
The authors declare no competing interests.
Figure titles and Figure legends

Figure 1. Wide-spread apical planar polarisation of Crumbs and Rho-kinase during morphogenesis.

A The salivary gland develops from an epithelial placode on the ventral side of the embryo (green area). Cells become specified at late stage 10/early stage 11 and by stage 12 most secretory cells have invaginated to form a tube inside the embryo. A’ The salivary gland placode boundary shows a strong enrichment of myosin II (Zipper-YFP, green) into a supracellular cable, complementary to anisotropic enrichment of Crumbs (Crumbs-GFP, magenta) in the placodal boundary cells (Röper, 2012); stills of a time lapse movie of matching stages to schematics in A (see also Figure S1). B Genomic locus of Drosophila Rok, indicating exon usage in splice variants and the position of the mNeonGreen (mNG) tag inserted at the N-terminus. C-H mNG-Rok enrichment (green) complementary to Crumbs anisotropic localisation (magenta) at many epithelial boundaries: in the embryo, the salivary gland placode (C-C’’), the boundary of the posterior spiracle placode (D-D’’), the spiracular hair precursors (E-E’), the ventral midline (F-F’’), the leading edge during dorsal closure (G-G’’), as well as the boundaries of specified photoreceptor clusters during larval eye disc morphogenesis (H-I’’). Boxes in D, H indicate magnifications in E-E’’ and I-I’’, respectively. Arrowheads point to the boundaries in Crumbs levels that show strong mNG-Rok accumulation.
Figure 2. Rho-kinase planar polarisation at the boundary of the salivary gland placode.

A-A‴ Crumbs is highly enriched in the salivary gland placode and reduced in levels in the surrounding epidermis (A), leading to its strong anisotropic localisation in boundary cells due to homophilic interactions (white arrows in A, A‴). mNG-Rok is enriched complementary to Crumbs in boundary cells, but also strongly enriched isotropically in junctions in the inner cells of the placode (A′, green). B-B‴ Introduction of a new boundary of Crumbs protein levels within the placode, using en-Gal4 x UAS-Crumbs (B, magenta) leads to accumulation of mNG-Rok at the new boundary (B′, green), complementary to anisotropic Crumbs at the new boundary. See also Figure S2. C The anisotropic and complementary localisation of Crumbs and mNG-Rok shows a ‘boundary effect’, the negative regulatory effect that Crumbs exerts on Rok localisation is only apparent in cells that show Crumbs anisotropy (boundary cell), whereas isotropic Crumbs accumulation in the centre of the placode (inner cell) does not prevent high levels of junctional Rok accumulation. D A dynamic model of modulation of Rok residence time at the membrane, depending on local levels of Crumbs, could explain planar polarisation of Rok in cells showing Crumbs anisotropy whilst preserving isotropic Rok localisation in cells with isotropic Crumbs. Asterisks in A‴, B‴ indicate the invagination point.
Figure 3. FRAP analysis and in silico simulation of Rok dynamics at the boundary versus inside the placode.

A Schematic of the placode indicating the boundary with low levels of Crumbs (blue), the side membranes of boundary cells with high levels of Crumbs (magenta) and the inner placodal cells with isotropically high levels of Crumbs (orange) where FRAP analysis was performed. The invagination point (asterisk) and ventral midline position are indicated. B, B’ Examples of boundary (B) and inner (B’) cell junctions of mNG-Rok embryos during FRAP analysis, arrows indicate the positions of the bleached regions (bleach at t=0). C, D Recovery curves fitted to data of FRAP experiments for boundary junctions (blue; n=27), side junctions (magenta; n=7) and inner cell junctions (orange; n=13)(C). k_{off} values were estimated from the fluorescence recovery for the boundary as 0.057 (+/- 0.008), for the inner placodal membranes as 0.149 (+/- 0.018) and for side membranes as 0.173 (+/- 0.024) and inner and side were found to be significantly different from the boundary using a bootstrap procedure, with the p-values determined as 0.0002 for both (***) (D). Data represented are bootstrap sample, median and quartiles. E We modelled the Rok planar polarisation in silico using particle based stochastic reaction-diffusion. F A group of cells representing boundary and inner placodal cells are modelled, imposing different k_{off} values for the boundary with low Crumbs levels (region 2, blue) and a higher k_{off} for membranes with high levels of Crumbs, such as side membrane and inner placodal membranes (region 1, orange). G Examples of steady state outputs of the simulation, with simulation 1 depicting no difference in k_{off}, simulation 2 assuming a 5-fold difference in k_{off}, and simulation 3 using the k_{off} values determined by FRAP as input; numbers on representative junctions are the particle numbers derived from simulations. See also Figure S3.
**Figure 4. Rok’s RBD and PH domain are required for its planar polarisation and placode enrichment.**

A Schematic of Rok protein domains: N-terminal kinase domain followed by a coiled-coil domain, a Shroom-binding domain (SBD), a Rho-binding domain (RBD) and a C-terminal PH-domain. B-F' Expression of variants of a Venus-tagged and kinase-dead localisation reporter of Rok (Venus-Rok[K116A]) throughout the embryo using Da-Gal4: B, B’ full-length Venus-Rok[K116A] similar to mNG-Rok shows anisotropic enrichment at the boundary (F’), though its accumulation at certain sites such as in the apical-medial region is enhanced due to the overexpression. C, C’ Rok lacking the coiled coil and SBD (Rok ∆CC-SBD) is less polarised at the boundary but retains some anisotropy (F’) and also shows apical-medial aggregates. D,D’ Rok lacking the RBD (Rok ∆RBD) is still localised to cell junctions but not enriched in the placode and shows no polarisation at the boundary (F’). E,E’ Rok lacking the PH domain (Rok ∆PH) is still localised to cell junctions but not enriched in the placode and shows no polarisation at the boundary (F’). F,F’ Polarisation quantification expressed as the intensity ratio of boundary junction versus side junctions (F). F’ Mean values are: Rok[K116A]=1.86; Rok ∆CC-SBD=1.433; Rok ∆RBD=1.005; Rok ∆PH=1.701. Data are represented as data points, mean and SEM. Statistical tests used were unpaired t-test, p values are indicated with ** being <0.005, *** being <0.0005, ****<0.0001, ns being not significant. G-I Neither RhoGEF2 (G-G’; sfGFP-RhoGEF2) nor active Rho (H-H’; EGFP-ANI RBD) are polarised at the placode boundary. I Polarisation quantification expressed as the intensity ratio of boundary junction versus side junction. Data are represented as data points, mean and SEM. Mean values are: sfGFP-RhoGEF2=0.9267; EGFP-ANI RBD=0.8862.
**Figure 5. Pak1 can phosphorylate Rok and induce its dissociation from the membrane.**

A Crumbs’ intracellular tail can interact with two kinases with highly overlapping phosphorylation targets: aPKC and Pak1. B Both aPKC and Pak1 have many overlapping predicted phosphorylation sites in Rok, in particular near the C-terminal RBD and PH domains. Residues marked in red and named above the sequence are conserved between *Dm* Rok and *Hs* ROCK1 and are phosphorylated in human EpH4 cells (Ishiuchi and Takeichi, 2011). Residues marked by asterisks were mutated in mNG-Rok[Pmut], see Figure 6. Phosphorylation of Rok by either or both kinases could promote its dissociation from the plasma membrane (A). See also Figure S4. C-C’ Overexpression in stripes of a membrane-targeted version of aPKC (magenta in C; *en-Gal4 UAS-aPKC[CAAX]*) does not significantly affect membrane localisation of mNG-Rok (green) in the overexpressing stripes, quantified in C”. Data are represented as data points, mean and SEM. Unpaired t-test was used to determine statistical significance, p value is 0.278 (ns). D-D” Overexpression in stripes of a membrane-targeted version of Pak1 (red marks overexpression stripe in D; *en-Gal4 x UAS-mCherry, UAS-Pak1myr*) leads to loss of mNG-Rok (green) from the membrane in the overexpressing stripes, as quantified in D”. Data are represented as data points, mean and SEM. Unpaired t-test was used to determine statistical significance, p value is <0.0001 (****). E Using purified kinase domains of either aPKC or Pak1 and short peptide substrates of Rok (indicated in B) in an *in vitro* kinase assay, we can detect phosphorylation of Rok peptide 2 (Rok pep2), that is located near the PH domain, by Pak1, but no phosphorylation of any Rok peptide tested by aPKC. Both kinase domains phosphorylate a known substrate peptide from Bazooka (BazS980). Data show fold enrichment of radioactive P[33]-phosphate counts over ‘no substrate’ control, data points, mean and SEM are shown. Rok pep2AAA is identical to the Rok pep2 peptide, with three potential phosphorylation sites mutated to alanine, S1365A, T1366A, T1369A. See also Figure S4.
Figure 6. A phospho-mutant Rok shows reduced planar polarisation at the boundary.

A Schematic of the potential target sites for aPKC or Pak1 phosphorylation that are mutated in the phospho-mutant Rok (mNG-Rok[Pmut]). B,C Localisation comparison between mNG-Rok (B) and mNG-Rok[Pmut] (C). Arrows mark the boundary, the asterisks mark the invagination point. D Flies homozygous for mNG-Rok[Pmut] are semi-viable, with 12.1% of embryos not hatching compared to 5.9% in the mNG-Rok control. E-G Compared to the mNG-Rok control (E, green) that shows polarisation at the boundary marked by Crumbs (magenta), mNG-Rok[Pmut] (F, green) shows a loss of polarisation at the boundary marked by Crumbs (magenta).

G Polarisation quantification expressed as the intensity ratio of boundary junction versus side junction. Data are represented as data points, mean and SEM. Mean values are: mNG-Rok ctrl: 1.36; mNG-Rok[Pmut]: 0.935. Unpaired t-test was used to compare samples, p value is indicated. H-J Stage 13 mNG-Rok[Pmut] embryos show a strong disruption of the placodal boundary in 23.3% of embryos (compared to 0.0% in the control), Rok is in green in H, I, with phalloidin to label cell outlines in magenta in H, I.

J Quantification of stage 13 gland phenotype prevalence. Arrowheads in H, I point to the pharyngeal ridges and identify embryos as stage 13, arrows in H point to the planar polarised mNG-Rok control at the boundary, magenta arrows in I' point to the disorganised and disrupted boundary in mNG-Rok[Pmut].
**Figure 7. Pak1 acts downstream of Crumbs/Cdc42 and controls Rok planar polarisation and cable formation at the placode boundary.**

A-B’’ In contrast to the wild-type (A-A’’)) where Pak1 is enriched in the placode (A’) and mNG-Rok strongly accumulates at the placode boundary (A), in pak1^{14} zygotic mutants Pak1 antibody labelling is strongly reduced (B’) and mNG-Rok is not polarised to the placode boundary (B, quantification in E). C-D’’ Overexpression of the membrane-targeted form of Pak1 in the placode (UAS-Pak1^{myr} x fkh-Gal4) leads to a disorganised placode at stage 11 (C’’) and loss of mNG-Rok polarisation (C, and quantification in E). At late stage12, many UAS-Pak1^{myr} x fkhGal4 placodes show a disruption that suggests a ripping of cells at the placode boundary (D’’). mNG-Rok remains at remnant cell boundaries (D). C’ and D’ show the increased levels of overexpressed membrane-localised Pak1 by antibody staining. E Polarisation quantification expressed as the intensity ratio of boundary junction versus side junction. Data are represented as data points, mean and SEM. Mean values are: mNG-Rok control=1.304; mNG-Rok in pak1^{14}=0.99; mNG-Rok in fkh-Gal4 UAS-Pak1^{myr} =1.006. Statistical tests used were unpaired t-test, p values are indicated with **** being <0.0001. Dotted lines indicate the placode boundary and asterisks the invagination point.

F-F’’ Pak1 (F’ and magenta; revealed by anti-Pak1 antibody) is enriched in the salivary gland placode compared to the surrounding epidermis. Crumbs is shown in F and green. G-G’’ Cdc42-mCherry localisation (G’ and magenta) follows endogenous Crumbs anisotropic localisation (G and green) at the placode boundary; insets show magnification of boundary cells (arrows point to boundary membrane). Polarisation quantification of mCherry-Cdc42 and Crumbs in corresponding cells expressed as the intensity ratio of boundary junction versus side junction (J). Mean values are: mCherry-Cdc42: 0.906; Crumbs: 0.714. Data are represented as data points, mean and SEM. H-H’’ Overexpression of Crumbs (H and green) in enGal4 stripes leads to ectopic recruitment of mCherry-Cdc42 (H’ and magenta) to sites of ectopic Crumbs, again following the anisotropy (arrows). I-I’’ Overexpression of a constitutively active form of Cdc42, Cdc42^{V12}, in enGal4 stripes leads to a strongly increased membrane association of Pak1 (magenta in I) in the overexpressing stripes. Membranes are labelled with phalloidin to reveal F-actin (green in I). I’ and I’’ are magnifications of the box indicated in I; data are quantified in K. K Mean values are UAS-Cdc42^{V12} x enGal4 stripe: 2.61; control stripe: 1.50. Data are represented as data points, mean and SEM. Statistical tests used were unpaired t-test, p value is indicated with **** being <0.0001.
Dotted lines indicate the boundary of the placode and asterisks mark the invagination point.

**L-M** Modulation of Pak1 levels affects mNG-Rok $k_{\text{off}}$. **L** Recovery curves fitted to data of FRAP experiments for boundary and inner junctions respectively for Pak1 overexpression (using $UAS\text{-}Pak1^{myr} \times fkhGal4$; n[boundary]: 7, n[inner]: 18) and for the $pak1^{14}$ mutant (n[boundary]: 5, n[inner]: 14). **M** $k_{\text{off}}$ values were estimated from the fluorescence recovery for the boundary [$UAS\text{-}Pak1^{myr}$] as 0.17 (+/- 0.022), for inner membranes [$UAS\text{-}Pak1^{myr}$] as 0.266 (+/- 0.025), for the boundary [$pak1^{14}$] as 0.032 (+/- 0.013) and for inner membranes [$pak1^{14}$] as 0.044 (+/- 0.008). Both Pak1 modulations were found to be significantly different from the control using a bootstrap procedure, with the p-values determined as ***= 0.0002, **= 0.008, *= 0.013. See also Figure S5.
LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Dr Katja Röper (kroeper@mrc-lmb.cam.ac.uk).

EXPERIMENTAL MODEL

*Drosophila melanogaster* was cultured using standard techniques at 25°C (unless otherwise noted); both male and female animals were used.

METHOD DETAILS

*Drosophila stocks and genetics*

The following transgenic fly lines were used and are also listed in the Key Resources Table: *sqhAX3; sqh::sqhGFP42* (Royou et al., 2004), *Daughterless-Gal4* and *enGal4* (Bloomington Stock Centre); *fkhGal4* [(Henderson and Andrew, 2000; Zhou et al., 2001) kind gift of Debbie Andrew]; *Crb-GFP* (Huang et al., 2008); *Zip-YFP* (Lowe et al., 2014); *US-Crb* (Wodarz et al., 1995); *UAS-Venus-Rok[K116A], UAS-Venus-Rok[K116A]ΔRBD, UAS-Venus-Rok[K116A]ΔCC-SBD; UAS-Venus-Rok[K116A]ΔPH* (Simoes Sde et al., 2014); *Ubi-EGFP-AnillinRBD* (Munjal et al., 2015); *sfGFP-RhoGEF2* (Sarov et al., 2016); *UAS-aPKC[CAAX]* (Sotillos et al., 2004); *UAS-Pak1myr; UAS-Cdc42[V12]; sqh::Cdc42-mCherry* (Bloomington Stock Centre); *pak1[14]* (gift from B. Thompson); *y[1] sc[1] v[1]; y[+t7.7] v[+t1.8]=nanos-Cas9attp2* (gift from N. Perrimon).

Genotypes analysed are indicated in the figure panels and legends.

**Generation of transgenic fly lines**

To generate *Drosophila rok* transgenic lines, donor (150 ng/µl) and guide RNA (100 ng/µl) plasmids were injected in pools (Bischof et al., 2013) into *nanos::Cas9* (chromosome 3) embryos for the endogenous tagging, or into *mNG-Rok* ; *nanos::Cas9* embryos for the generation of the phospho-site mutant.

**mNG-Rok generation by CRISPR/Cas9**

Two gRNAs targeting loci near the start codon of the Rho kinase gene were cloned into pCFD3 vector (Addgene 49410) following the protocol from (Port et al., 2014). A step by step protocol is available at (www.crisprflydesign.org).
Sequences of the guide RNAs were as follows:
gRNA 1: GACCAACAGGAAAGCAGCAGCTGG
gRNA2: GCGCCGGTGAGTGACGAGATGG
PCR primers for cloning into pCFD3:
P56F: 5’-GTCGACCAACAGGAAAGCAGCAGC-3’
P56R: 5’-AAACGCTGCTGCTTCTGTTGTG-3’
P57F: 5’-GTCGCGCCGGTGAGTGACGAGA-3’
P57R: 5’-AAACTCTCGTGCACTCACCAGCG-3’

A donor construct containing the mNeonGreen sequence in fusion with the rok gene in its genomic region was cloned into pBluescript SK(+) using Gibson assembly.
mNeonGreen was cloned between two 1kb-long homology sequences corresponding to the genomic sequence on either side of the insertion site, to create homology arms for directed repair. The 1 kb regions were amplified by PCR on each side of the desired insertion site from genomic DNA. The mNeonGreen gene (Shaner et al., 2013), was amplified by PCR from the mNeonGreen vector (Allele Biotechnology), with the exclusion of the stop codon and the addition of a C-terminal linker. Primers were designed with additional 5’ sequences (underlined below) to allow triple ligation of the three PCR products into a pBluescript SK(+) vector using the Gibson Assembly Master Mix (NEB). PAM sites were mutagenized (indicated in bold below) to prevent re-cutting by Cas9 after transgenesis.

Primer sequences:
Left homology arm PCR primers:
P51F: 5’GACGGTATCGATAAGCTTGATATCGCGCAGCGTCTAATTGAAAC-3’
P51R: 5’GCTGATACTGCTGCTAAGCTGCTGCTGC-3’

Right homology arm PCR primers:
5’TGCCAGCTGGACGAGAAACTGTGAAGACCAGCGCAGCATGGATGTGGAACG
AAGGCCGCGGtgaagtgcagagactgtcggccccaaagc

mNeonGreen PCR primers:
P54F:
5’GGAAGCAGCAGCTGTAGCAGCAGCTATCGAGCTTGTGTTATCTTGCGATTTGCTG
GAGCAAGGGCGAGGAG-3’
P54R:
5’GCTTGGTACAGATTTCTCGTCCAGCTGGCAGATCCGCCGCCCGGATCC
GCCGCCGGATCCGCCCTTGTAAAGTTCATCCATCCCATCCC-3’

Modifications were verified by sequencing of genomic DNA.
**Mutagenesis of putative P-sites in mNG-Rok**

Guide RNAs:
Four gRNAs targeting loci on both sides of the C-terminal region of Rok containing the putative phosphorylation sites to be mutated were cloned into pCFD3 vector (Addgene 49410) following the protocol from (Port et al., 2014).
Sequences: LJ20, LJ21, LJ22, LJ23
Donor plasmid:
A donor construct containing, between two 1kb-long homology arms, a Rok C-term region mutagenized on 13 selected putative phosphorylation sites, was cloned into pBluescript SK(+). The Rok C-term region was amplified by PCR and cloned into pBluescript SK(+)with primers designed to mutagenise the selected 13 putative phosphorylation sites (Fig. 5B) and the 4 PAM sites.
Modifications were verified by sequencing of genomic DNA.

**Embryo Immunofluorescence**
Embryos were collected on apple juice-agar plates and processed for immunofluorescence using standard procedures. Briefly, embryos were dechorionated in 50% bleach, fixed in 10% EM-grade formaldehyde, and stained with primary and secondary antibodies in PBT (PBS plus 0.5% bovine serum albumin and 0.3% Triton X-100). anti-Crumbs and anti-E-Cadherin antibodies were obtained from the Developmental Studies Hybridoma Bank at the University of Iowa (DSHB); anti-Pak1 (Harden et al., 1996); anti-aPKC (Santa Cruz); rhodamine-coupled phalloidin (Molecular Probes). Secondary antibodies used were Alexa Fluor 488/Fluor 549/Fluor 649 coupled (Molecular Probes) and Cy3 and Cy5 coupled (Jackson ImmunoResearch Laboratories). Samples were embedded in Vectashield (Vectorlabs).

**Confocal, and Time-lapse imaging**
Images of fixed samples were acquired on an Olympus FluoView 1200 or a Zeiss 780 Confocal Laser scanning system as z-stacks to cover the whole apical surface of cells in the placode. Z-stack projections were assembled in ImageJ or Imaris (Bitplane), 3D rendering was performed in Imaris.

For live time-lapse experiments embryos of the genotype Crumbs-GFP Zipper-YFP or mNG-Rok were dechorionated in 50% bleach and extensively rinsed in water. Embryos were manually aligned and attached to heptane-glue coated coverslips and mounted on custom-made metal slides; embryos were covered using halocarbon oil 27 (Sigma) and viability after imaging after 24h was controlled prior to
further data analysis. Time-lapse sequences were imaged under a 40x/1.3NA oil objective on an inverted Zeiss 780 Laser scanning system. Z-stack projections to generate movies in Supplementary Material were assembled in ImageJ.

**Embryo Viability Assay**

Embryos of the genotype mNG-Rok control or mNG-Rok[Pmut] were treated as for live imaging and mounted in separate sets of 100 embryos per experiment and let to develop at 18°C. After 48 hours hatched larvae, unfertilised embryos and developed but dead embryos were counted.

**Rho-kinase sequence analysis**

We used published predictive algorithms to identify potential aPKC and Pak1 phosphorylation sites in Dm Rok, DISPHOS (http://www.dabi.temple.edu/disphos/) and NetPhos3.1 (http://www.cbs.dtu.dk/services/NetPhos/). Rok sequences from different species were compared and aligned in EMBOSS Matcher (https://www.ebi.ac.uk/Tools/psa/emboss_matcher/).

**In vitro kinase assay**

The following High-pressure liquid chromatography (HPLC)-purified peptides were ordered from Biomatik:

- BazpepS980: EHFSRDALGRRSISEKHHAA
- Rokpep1T: LPKQKnKARSTAEFLKKEKEM
- Rokpep2STS: SKIPSQSTRSSYKPYAVNV
- Rokpep2AAA: SKIPSQAARSAYKPYAVNV
- Rokpep3S: KPYAVVVQRSATLPANSSLK

For in vitro kinase assays, 10µg of peptide substrate were incubated with either 150pg recombinant human Pak1 kinase domain (AbNOVA) or 0.1 µM recombinant human PKCδ kinase domain (a gift from N. McDonald via B. Thompson) for 30 min at 30°C in kinase reaction buffer (50 mM HEPES [(4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid] pH 7.5, 10 mM MgCl2, 1 mM EGTA, 0.01% Brij35) containing 10 µM cold ATP and 3 µCi γ-P33 ATP (Hartmann Analytic GmbH). Samples were blotted on 2cm x 2cm squares of P81 phosphocellulose paper (Millipore) and washed 3 x 10 min in 1% phosphoric acid, then 5 min in acetone. Dried papers were then transferred to scintillation vials and immersed in liquid scintillation cocktail (Ultima Gold XR, Perkin Elmer). Incorporation of γ-P33 was quantified in counts per minute by scintillation counting (Beckman LS 6500).
QUANTIFICATION AND STATISTICAL ANALYSIS

Fluorescence Intensity Quantifications

Fluorescence intensity was determined in ImageJ using projections covering the apical junctional region (as determined by Crumbs or E-Cadherin staining). Using Crumbs labelling of placodes, boundary cells showing clear Crumbs anisotropy were identified and quantified. 3- to 5-pixel-wide lines to cover the width of junctions (depending on the resolution of the image) were drawn at the boundary and at the sides of the boundary cells (see Fig. 4F). The intensity was divided by the area covered for each junction to determine the intensity/pixel. Values for the boundary junction were divided by the values of the side junctions to determine two polarisation values per boundary cell.

Membrane versus cytoplasm enrichment was determined in ImageJ using projections covering the apical junctional region (as determined by Crumbs or E-Cadherin staining). 3- to 5-pixel-wide lines to cover the width of junctions (depending on the resolution of the image) were drawn at a cell junction and a comparable line was drawn across the apical cytoplasm. The intensity of each line was divided by the area covered to determine the intensity/pixel. Values for the cell junction were divided by the values of the cytoplasm to determine the membrane versus cytoplasm enrichment.

N values for quantifications are as follows: Figure 4F': **UAS-Rok[K116A]**: 4 placodes (3 embryos), 15 cells (29 polarisation values); **UAS-Rok[K116A]∆CC-SBD**: 6 placodes (3 embryos), 18 cells (36 pol. values); **UAS-Rok[K116A]∆RBD**: 5 placodes (3 embryos), 16 cells (32 pol. values); **UAS-Rok[K116A]∆PH**: 6 placodes (3 embryos), 16 cells (32 pol. values)/ Figure 4I: **Ubi-EGFP-AniRBD**: 4 placodes (3 embryos), 15 cells (30 pol. values); **RhoGEF2 sfGFP**: 3 placodes (3 embryos), 17 cells (34 pol. values)/ Figure 5C'': **UAS-aPKC[CAAX]**: 3 embryos (6 overexpression stripes), 30 cells; **Control for aPKC stripe**: 3 embryos (6 control stripes), 30 cells/ Figure 5D'': **UAS-Pak1[myr]**: 4 embryos (7 overexpression stripes), 35 cells; **Control for Pak1 stripe**: 4 embryos (7 control stripes), 35 cells/ Figure 6E: **mNG-Rok ctrl**: 4 placodes (2 embryos), 23 cells (46 pol. values); **mNG-Rok in pak1[14]**: 3 placodes (2 embryos), 15 cells (30 pol. values); **mNG-Rok fkhGal4 UAS-Pak1[myr]**: 4 placodes (2 embryos), 26 cells (32 pol. values)/ Figure 6J: **mCherry-Cdc42**: 4 placodes (3 embryos), 24 cells (48 pol. values); **Crumbs**: 3 placodes (2 embryos), 14 cells (28 pol. values)/ Figure 7G: **mNG-Rok ctrl**: 4 placodes (4 embryos), 17 cells (34 pol. values); **mNG-Rok[Pmut]**: 4 placodes (4 embryos), 18 cells (34 pol. measurements).
Statistical significance in comparisons was determined using unpaired t-test. Plots show data points, mean and SEM. p-values are indicated as ** being <0.005, *** being <0.0005, ****<0.0001, ns being not significant.

**FRAP imaging and analysis**

Focussing on the apical region of the salivary placode epithelium in stage 11/12 mNG-Rok embryos (or mNG-Rok in the indicated genetic backgrounds), spinning disk confocal/Fluorescence Recovery After Fluorescence (FRAP) was performed on a custom-built set-up based on a Nikon Ti stand equipped with perfect focus system, a fast Z piezo stage (ASI), a PLAN NA 1.4 60x objective and a spinning disk head (Yokogawa CSUX1) followed by 1.2x relay optics. Images were recorded with a Photometrics Prime 95B back-illuminated sCMOS camera run in pseudo global shutter mode and synchronized with the spinning disk wheel. FRAP was performed using a iLAS2 galvanometer module (Roper France) mounted on the back port of the stand and combined with the side spinning disk illumination path using a broadband polarizing beamsplitter mounted in a custom 3D-printed fluorescence filter cube. GFP was excited/bleached by a 150mW 488nm laser (Coherent OBIS mounted in a Cairn laser launch) and GFP fluorescence was imaged using a Chroma 525/50 bandpass filter. System was operated by Metamorph. About 2 μm thick z-sections (either 3 x 0.65μm or 5 x 0.5μm) were acquired to compensate for movement in z during acquisition. For the FRAP, bleach dwell time was 19ms with 28% 488nm laser power. Images were acquired at ~500ms (for 3 z steps) or ~750ms (for 5 z steps) intervals, 6 time points pre-bleach and 60 (or 45) time points post bleach.

We analysed kymographs of recovery regions and surrounding membrane regions to exclude that recovery was due to lateral movement of mNG-Rok in the membrane rather than recovery from the cytoplasmic pool.

Movies were analysed in ImageJ/Fiji (NIH). Fluorescence intensity was measured in a 6-pixel circular ROI at the site of bleach.

Measures were then normalised to account for the general photobleaching caused by image acquisition. All values were multiplied by a photobleaching correction factor determined from a 100μm diameter circle surrounding the bleach site.

\[
C_{\text{photobleaching}}(t) = \frac{\text{background circle intensity at } t_1}{\text{background circle intensity } (t)}
\]

Normalised fluorescence intensity measurements were used to plot the percentage of fluorescence recovery after photobleaching as follows:

\[
F_{\text{prebleach}} = \text{avg } F_{t_1 \text{ to } t_6}
\]
\[
F(t) = \frac{(F_t - F_{\text{postbleach}})}{(F_{\text{prebleach}} - F_{\text{postbleach}})}
\]

**FRAP curve fitting and statistical analysis**

The \( k_{\text{off}} \) was estimated from the whole set of normalized fluorescence recovery curves. We modeled the recovery using a single exponential function in the form of:

\[
a (1 - e^{-k_{\text{off}}t})
\]

A detailed description of the deduction of the \( k_{\text{off}} \) can be found in the Supplemental Information. A non-linear regression algorithm was used to estimate parameters from the data without prior averaging. As individual fluorescence recovery curves were noisy, we used a bootstrap procedure generating 200 estimates to estimate the statistical confidence of the estimated \( k_{\text{off}} \). This allowed us to compute a p-value using a t-test based on bootstrap variance. Note that this p-value does not depend on the number of bootstrap samples.

**DATA AND CODE AVAILABILITY**

**In silico Rok particle simulation**

As illustrated in Figure 3F, we simulated the diffusion of membrane bound and unbound particles (corresponding to Rok) within a pixel representation of a placodal cell layer containing inner and boundary cells. Our intention was to create a deliberately simple two-dimensional model that might nonetheless recapitulate the salient features of the observed planar polarity. Different membrane association and dissociation constants were modelled in different regions of the simulation according to the zones illustrated in Figure 3G, i.e. distinguishing the placode boundary from the inner placode membranes including the side membranes. Here the representation of the cell membranes with superimposed particle positions automatically provided a visualization of the simulation progress. By counting bound and unbound particles in different regions, polarisation at the boundary was quantified and compared with microscopic measurements. An exhaustive grid-search for parameters, which was initially coarse and then more fine-grained, was performed to determine the combinations of diffusion, binding, unbinding and simulation time-step values that matched observations.

In detail, point particles were modelled as moving with random sequential displacements according to 2D Gaussian diffusion within a grid-based (i.e. pixel) representation of a cell layer with membrane boundaries. Particles were set to freely diffuse (off-grid) in the cell interior or diffuse laterally along the interior edge of the cell membrane; i.e. as unbound and bound states. The positional variance for free
diffusion (sigma) was set to correspond approximately to a diffusion coefficient of 25 μm²/s within a 25 μm wide cell area, which is the diffusion coefficient measured in vivo for mammalian GFP-ROCK2 (Truebestein et al., 2015). Diffusion of bound particles within the membrane was restricted to adjacent sites and was much slower than for free particles, with diffusion coefficients tested in the range 0.1 - 0.001 μm²/s (which made no practical difference) and final simulations set at 0.01 μm²/s. Using a probabilistic model, that can be related to $k_{on}$ (association) and $k_{off}$ (dissociation) constants, free particles that collided with the cell membrane were able to bind and bound particles were able to spontaneously unbind/dissociate. Here simulation steps typically corresponded to time segments of 50 ms, though a range of values was tested. $k_{off}$ (in units of per second) was used to set the long-term probability of each bound particle spontaneously unbinding within the simulation time-step. $k_{on}$ was less straightforward to model as, for particles which collide with the membrane, it depends on the concentration of receptive membrane sites and this varies throughout the simulation. Accordingly, at-membrane binding probabilities of initially free particles were set dynamically so that the derived, average $k_{on}$ measured in the simulation converged to a desired value. In essence membrane binding probability was increased when the target $k_{on}$ was undershot and reduced when overshot, averaging around a fixed $k_{on}$. A single $k_{on}$ value was used for all membrane regions, but different $k_{off}$ values were used for placode boundary membranes and inner/side placode membranes.

The in vivo $k_{on}$ used in simulations shown in Fig. 3 was estimated using the following equation from the "diffusion plus binding" model from (Sprague et al., 2004):

$$k_{on}^* = \frac{k_{off} C_{eq}}{F_{eq}}$$

The $k_{off}$ was calculated from our FRAP experiments, and we found that the $\frac{C_{eq}}{F_{eq}}$ ratio could be deduced from parameters measured in vivo:

$$\frac{C_{eq}}{F_{eq}} = \frac{\text{[membrane]}}{\text{[cytoplasmic]}} \times \text{mobile fraction}$$

We measured the membrane to cytoplasmic ratio of mNG-Rok in stage 11 embryos and the value of the $k_{off}$ and mobile fraction were calculated from fitted FRAP curves (Fig. 3C and Supplemental Fig.3 A,B): $k_{off} = 0.149$; mobile fraction = 0.77; membrane versus cytoplasm ratio: 2.367.

Using the above equation and in vivo measured values, we estimated the $k_{on}$ to be about 0.27, using 0.3 for simulations in Fig. 3, but also testing a range of $k_{on}$ values
between 0.1 and 0.4 in the simulations (0.2, 0.3. and 0.4 are shown). Varying $k_{on}$ within this range had only minor effects on polarisation values.

In order that the membrane binding sites be capable of saturation, the cell edge pixels were subjected to a maximum occupancy value. For the final simulations this was set at a value of 1 particle per pixel. Occupancy limits of 1-3 particles per pixel were tested and overall this made little difference to the cell polarity, but had a notable effect on the bound/free ratio, as we might expect.

Although the number of particles modelled within each cell could be varied, this made little difference to the long term bound/unbound ratios, but naturally more particles gave smoother, less variant values. Typically, 2000 particles per cell were simulated. The particle simulation was started with randomly distributed particle positions within the cell interiors and progressed through 2000 unmonitored steps to equilibrate the model. Thereafter analyses of the particle positions were made at regularly spaced intervals for a further 10,000 steps. At each sample point the counts of bound and unbound particles were recorded for the regions marked in Figure 3G and later averaged for the whole simulation. Particle number had a slight effect on polarisation values, with smaller particle numbers leading to slightly higher polarisation values (polarisation of 1.90 for 125 particles versus polarisation of 1.48 for 2000 particles, with $k_{on}$ of 0.3 and the measured values of $k_{off}$ for both situations).

Python code to perform the 2D cell particle simulations, generating both regional counts and pixmap images, is available at: https://github.com/tjs23/memodis
Supplemental Movie 1. Dynamics of Crumbs and Myosin at the salivary gland placode boundary.
Time-lapse movie of an embryo with endogenously tagged Crumbs (green, Crumbs-GFP) and endogenously tagged Myosin II heavy chain (magenta, Zipper-YFP). Frames are 3 min apart.

Supplemental Movie 2. Example of FRAP at a salivary gland placode boundary junction.
Time-lapse movie of an embryo with endogenously tagged Rok (mNG-Rok), showing a bleach and recovery at the boundary of the placode, quantification in Figure 3.

Supplemental Movie 3. Example of FRAP at a salivary gland inner cell junction.
Time-lapse movie of an embryo with endogenously tagged Rok (mNG-Rok), showing a bleach and recovery at a junction of an inner placodal cell, quantification in Figure 3.

Supplemental File 1. Reasoning for k_{off} deduction from FRAP recovery curves.
References


**Figure 1**

A. Ventral view of early stage 11 and lateral view of stage 12 salivary gland placode.

B. mNeonGreen-CRISPR knock-in into Rok locus.

C. Salivary gland placode with mNeonGreen-Rok expression.

D. Posterior spiracles with mNeonGreen-Rok expression.

E. Eye disc with mNeonGreen-Rok expression.

F. Live cross-section view showing Rok expression.

G. Ventral midline with dorsal closure and eye disc highlighted.

H. Live ventral midline with eye disc highlighted.

I. Live eye disc showing Rok expression.

J. Zipper-YFP and Crumbs-GFP expressions.
Sidor et al._Figure S1
Sidor et al._ Figure 2
Sidor et al._ Figure S2

A: en-Gal4 x UAS-Crb

αCrumbs

mNG-Rok

A': apical slices 1 to 4

B: en-Gal4 x UAS-Crb

αCrumbs

mNG-Rok

B': sub-apical slices 5 to 10

C: en-Gal4 x UAS-Crb

C': apical

basal
**Figure 3**

**A** Ventral midline with boundary and side junctions.

**B** Fluorescence recovery over time for boundary junctions.

**B’** Fluorescence recovery over time for inner junctions.

**C** Graph showing fluorescence recovery over time for boundary, inner, and side junctions.

**D** Distribution of dissociation rate constants ($k_{OFF}$) for boundary, inner, and side junctions.

**E** Diagram showing diffusion in the membrane and cytoplasm.

**F** Diagram illustrating the dissociation rate constants for high and low Crb conditions.

**G** Table showing the average number of membrane-bound particles per junction for different simulations.

Sidor et al._Figure 3
**E**

<table>
<thead>
<tr>
<th>Ratio</th>
<th>k\textsubscript{off} placode/ k\textsubscript{off} boundary</th>
</tr>
</thead>
<tbody>
<tr>
<td>1:1</td>
<td>boundary: k\textsubscript{off} = 0.1  \ placode: k\textsubscript{off} = 0.1</td>
</tr>
<tr>
<td>5:1</td>
<td>boundary: k\textsubscript{off} = 0.1  \ placode: k\textsubscript{off} = 0.5</td>
</tr>
<tr>
<td>2.6:1</td>
<td>boundary: k\textsubscript{off} = 0.057  \ placode: k\textsubscript{off} = 0.149</td>
</tr>
</tbody>
</table>

**Simulation Parameters**

- k\textsubscript{on} = 0.2
  - k\textsubscript{off}[placode] = 0.1  \\ k\textsubscript{off}[boundary] = 0.1
  - polarisation = 1.00
- k\textsubscript{on} = 0.3
  - k\textsubscript{off}[placode] = 0.1  \\ k\textsubscript{off}[boundary] = 0.1
  - polarisation = 1.00
- k\textsubscript{on} = 0.4
  - k\textsubscript{off}[placode] = 0.1  \\ k\textsubscript{off}[boundary] = 0.1
  - polarisation = 1.00
**Sidor et al._ Figure 4**

A **Rok (1391 aa)**

- Kinase domain
- Coiled-coil domain
- SBD
- RBD
- PH domain

<table>
<thead>
<tr>
<th></th>
<th>Full-length</th>
<th>Rok ΔCC-SBD</th>
<th>Rok ΔRBD</th>
<th>Rok ΔPH</th>
</tr>
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<tr>
<td><strong>B</strong></td>
<td>B'</td>
<td><strong>C</strong></td>
<td><strong>D</strong></td>
<td><strong>E</strong></td>
</tr>
<tr>
<td><strong>G</strong></td>
<td>G'</td>
<td><strong>G''</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>H</strong></td>
<td>H'</td>
<td><strong>H''</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**F**

* ns
* * P<0.05
** P<0.001
**** P<0.0001
Figure 5

A. 
- **en-Gal4 UAS-aPKC<sup>CAAX</sup>**

B. 
- **Rok (1391 aa)**
  - Kinase domain
  - Coiled-coil domain
  - SBD RBD
  - PH domain
- **Putative phosphorylation sites**
  - **Region 1:**
    - Dm Rok: KLAINEPRKXK-D-X-RA
    - Hs ROCK1: KLAINEPRKXK-D-X-RA
  - **Region 2:**
    - Dm Rok: S1365, T1366, T1369
    - Hs ROCK1: P1365, T1366, T1369

C. 
- **en-Gal4 UAS-mCherry<sup>myr</sup>, UAS-Pak1<sup>myr</sup>**

D. 
- **en-Gal4 x UAS-mCherry<sup>myr</sup>, UAS-Pak1<sup>myr</sup>**

E. 
- **Fold increase in counts over no substrate ctrl**
  - aPKC
  - Pak1

*CRISPR P-mutant S/T to A mutations
Dm Rho-kinase (isoform PA)

MPAGRETIVK QRSMDVERRR RAN TLEREMR DP TICNVDC LLDTVSA LV SCDHE SL RRL
K NIEQYAAK Y K PLAMQINQL RMVNDVFHFI KLIGAGAFGE VQLVHRHKS S QYVAMKRLLK
FEMMKRPDSA FFWEERHIMA HASE NWIQL HAFQDQAKYL YMVMDMPMPG DI SLMGDYD
IPEKWAFIYT MEVVLALTDI HMGFVHRD V KPDNMILLDS Y GHLKLADFGT CMR MGANGQV
VSSNAVGTPD YSPEVQLQ SQGV D N EY G Y EMLFG ETPFY AD SLVG T YGK
IMDHKNSLF PP E V ESEQA KAL IRAF TDL RTQRLG YG1 EDI KAH PFFR NDTW SF DNI R
ESVPPVVP E LQD DTRNRNE DIEREDEKPE VFPVPKGF D G NHP LFGI T GY DQYLD S S
TVDA EKEAN VANGSAASNN HGHGHNHRHR PSNSNELKRL EALLERERGR SEALEQ DAG
LRQQIE UI TL REAE LQ R IAS E Y EKDLAR LQ HN YKV AMQ K V EQ EIELR KKT EAL VETQRN
LENEQXT RLR D L N I ND K V VS LEKQLLE MEQ SYKTETEN TQ KLKHNAELD FTVKS QEEKV
RDMVDMIDTL QKH KELG QE NAELQALVQ EKNLRSQLE M HKEAENKMQ TLINDIERTM
CREQKQEDN R ALLEK ISDL EK AHAGLDFE LKAAQRGYQ QQ EVK A H QETEK SRLV SREEAN
LQEVKALOS LNEEK SARIK ADQH SQ EKER QLS MLSVDYR QIQ LRLQKLE GRE CRQ ESEK
AALOQQLQQE HSKR NAL S LSH S SE VAH LRS RENQLQK ELSTQEA RK RFEEDTLQLK
STHHEALANN RELQA Q L A E QCF S RLYKTQ ANEN REE SAE RL SKIEDL E R V SLK HQVQ
VAVARADSEA LR S I AEETV AD LEK E KTIK E EL KD F VMK HRNEINAK EA ALATLKEAEN
EHKHKLGOKA AEYEDLVQOH KK QQ EELAL M R S KD E EITK LLDKCK NEVL LI KQUA VN KLA
E VMNR RD D L P K QNK KAR T E AELR KKEK E M RRLQELSEQE RDKF NQL L L K H DL QQLCAE
EQQLQK QMV E M ID CK A TEI E NLQSKL NETA SLSS AD N DE PQ SQH S L S L TQDSVF G W L
SVPKQNR RR R GH W KQ YV YI V SS RK IF YN SDID KHNT TD A VL IL D LS K Y Y H V R S V T QD G
V IR A D A KEIP R IF Q LLYAGE GASH RP D EQS Q L D V S L H G N C N E R P T G IV HKGHEF V H IT
YHMPT ACEVC PKL WMHMF KP PAAYE CK R CR N KIH K HVDK HDPLAPCKLN HDP R S A R DML
LLA T PED O S LWVAR LLKR I QK SY KAA SY N NSTD G KI SP O RK S B Y KPV A NVN QR S
ALIPANSILK

→ predicted Pak1 sites score>0.8 (DISPHOS results)
→ predicted Pak1 sites score<0.8 (DISPHOS results)
→ predicted aPKC sites (NetPhos3.1)
→ putative sites for both aPKC and Pak1
→ peptides tested in vitro
→ conserved residues between human and Drosophila that are phosphorylated in EpH4 cells
→ bold and underlined residues mark the RBD and PH domain

Sidor et al. _Figure S4_
Sidor et al._Figure 6

A

mNG-Rok[Pmut]

B mNG-Rok ctrl

C mNG-Rok[Pmut]

D

<table>
<thead>
<tr>
<th></th>
<th>mNGRok ctrl</th>
<th>mNGRok mutant</th>
</tr>
</thead>
<tbody>
<tr>
<td>embryonic lethality</td>
<td>5.9% (16/269)</td>
<td>12.1% (33/273)</td>
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<tr>
<td>pupal lethality</td>
<td>0% (0/79)</td>
<td>0% (0/75)</td>
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E late stage 11

E' late stage 11

E'' late stage 11

E‴ late stage 11

E‴ late stage 11

E‴ late stage 11

E‴ late stage 11

F late stage 11

F′ late stage 11

F″ late stage 11

F‴ late stage 11

F‴ late stage 11

F‴ late stage 11

F‴ late stage 11

G

J

<table>
<thead>
<tr>
<th></th>
<th>mNGRok ctrl</th>
<th>mNGRok mutant</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;st13 gland phenotype</td>
<td>0.0% (0/21)</td>
<td>23.3% (7/31)</td>
</tr>
</tbody>
</table>

H late stage 11

H′ late stage 11

H″ late stage 11

H‴ late stage 11

I late stage 11

I′ late stage 11

I″ late stage 11

I‴ late stage 11

mNG-Rok kinase domain

coiled-coil domain

SBD

RBD

PH domain

region 1: 2 sites mutated

region 2: 11 sites mutated

<0.0001

<0.0001

<0.0001
Sidor et al. Figure S5