1 Co-ordinated Ras and Rac activity shapes macropinocytic cups

2 and enables phagocytosis of geometrically diverse bacteria

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1 Abstract

2 Engulfment of extracellular material by phagocytosis or macropinocytosis 3 depends on the ability of cells to generate specialised cup shaped protrusions. 4 To effectively capture and internalise their targets, these cups are organised into 5 a ring or ruffle of actin-driven protrusion encircling a static interior domain. 6 These functional domains depend on the combined activities of multiple Ras and 7 Rho family small GTPases, but how their activities are integrated and 8 differentially regulated over space and time is unknown. Here, we show that the 9 amoeba Dictyostelium discoideum coordinates Ras and Rac activity using the 10 multidomain protein RGBARG (<u>R</u>CC1, Rho<u>G</u>EF, <u>BAR</u> and Ras<u>G</u>AP-containing 11 protein). We find RGBARG uses a tripartite mechanism of Ras, Rac and 12 phospholipid interactions to localise at the protruding edge and interface with the interior of both macropinocytic and phagocytic cups. There, RGBARG shapes 13 14 the protrusion by driving Rac activation at the rim whilst suppressing expansion 15 of the active Ras interior domain. Consequently, cells lacking RGBARG form 16 enlarged, flat interior domains unable to form large macropinosomes. During 17 phagocytosis, we find that disruption of *RGBARG* causes a geometry-specific 18 defect in engulfing rod-shaped bacteria and ellipsoidal beads. This demonstrates 19 the importance of co-ordinating small GTPase activities during engulfment of 20 more complex shapes and thus the full physiological range of microbes, and how 21 this is achieved in a model professional phagocyte.

1 Introduction

The capture and engulfment of extracellular material serves a number of
important cellular functions. Best understood is the clearance of pathogenic
microbes or apoptotic cells by phagocytic immune cells, but the engulfment of
fluid by the related process of macropinocytosis also plays important functions
by allowing cells to capture antigens or other factors from their environment
such as nutrients to support growth (Bloomfield and Kay, 2016; Commisso et al.,
2013; Norbury et al., 1995; Sallusto et al., 1995; Swanson and King, 2019).

9

10 To capture extracellular material, cells must encircle and isolate their target

11 within a vesicle. This can be achieved by several mechanisms, but the best

12 understood and evolutionarily widespread involves the extension of a circular

13 cup or ruffle-shaped protrusion from the cell surface to enwrap the target and

14 close to internalise it (Buckley and King, 2017; Kaplan, 1977; Swanson, 2008;

15 Veltman et al., 2016). Whilst many components of cup formation have been

16 identified, how they are co-ordinated in space and time is poorly understood.

17 Here we describe a novel mechanism used by the amoebae *Dictyostelium*

18 *discoideum* to integrate different signaling elements and form complex-shaped

- 19 protrusions that efficiently mediate engulfment.
- 20

21 Macropinocytic and phagocytic protrusions are formed by localised actin 22 polymerisation at the plasma membrane, using much of the same machinery that 23 generates pseudopods and lamellipodia during cell migration (King and Kay, 24 2019; Swanson, 2008). Whilst migratory protrusions only need the cell to define 25 a simple patch of actin polymerisation, forming a cup requires a higher level of 26 organisation, with the protrusive activity restricted to a ring encircling a static 27 interior domain. During phagocytosis this is aided by the presence of a particle to 28 act as a physical scaffold and locally activate receptors. These interactions are 29 proposed to guide engulfment by a zippering mechanism (Griffin et al., 1975; 30 Tollis et al., 2010). However, macropinocytic cups self-organise with an almost 31 identical structure in the absence of any external spatial cues (Veltman et al., 32 2016). Cup formation can therefore occur spontaneously by the intrinsic 33 dynamics of the underlying signaling.

1

2 Recent studies in *Dictyostelium* proposed a model whereby the cup interior is 3 defined by spontaneous localised activation of the small GTPase Ras and 4 consequent accumulation of the phospholipid PIP₃ (Veltman et al., 2016). This 5 patch appears to restrict actin polymerisation to its periphery to create a 6 protrusive ring. How this is achieved is unknown, but in at least Dictyostelium it 7 may depend on the activity of the PIP₃-activated Protein kinase B/Akt (Williams 8 et al., 2019). Both active Ras and PIP₃ also accumulate at cups in mammalian 9 cells (Araki et al., 2007; Marshall et al., 2001; Vieira et al., 2001) and Ras 10 activation is sufficient to drive ruffling and macropinocytosis in cancer cells (Bar-11 Sagi and Feramisco, 1986; Commisso et al., 2013). PI3K inhibition also 12 completely blocks macropinocytosis (Amyere et al., 2000; Araki et al., 1996; 13 Hoeller et al., 2013; Veltman et al., 2014) as well as phagocytosis of large particles by macrophages (Araki et al., 1996; Cox et al., 1999; Schlam et al., 14 15 2015). Ras and PIP₃ therefore play a general role in macropinosome and 16 phagosome organisation across evolution.

17

Other small GTPases are also involved in cup formation. Active Rac1 overlaps 18 19 with Ras activity in the cup interior in both macrophages and *Dictyostelium* 20 (Hoppe and Swanson, 2004; Veltman et al., 2016). Rac1 is a direct activator of 21 the SCAR/WAVE complex, which drives activation of actin polymerisation via the 22 ARP2/3 complex (Eden et al., 2002; Machesky and Insall, 1998). Consistent with 23 this, Rac1 is required for macropinosome formation in dendritic cells (West et 24 al., 2000) and optogenetic Rac1 activation is sufficient to drive ruffling and 25 macropinocytosis in macrophages (Fujii et al., 2013). Expression of 26 constitutively active Rac1 also leads to excessive actin at macropinocytic cups in 27 Dictyostelium (Dumontier et al., 2000). Therefore, whilst Ras appears to define 28 the cup interior, Rac1 is important for regulating actin protrusions, as it is does 29 during cell migration.

30

31 The presence of active Rac1 throughout the cup interior is at odds with the

- 32 tightly restricted SCAR/WAVE activity and protrusion at the extending rim
- 33 (Veltman et al., 2016). A further layer of regulation must therefore exist. This is

1 likely provided by the small GTPase CDC42 which is also required for Fc-γ-

2 receptor mediated phagocytosis and collaborates with Rac1 during engulfment

- 3 of large particles (Caron and Hall, 1998; Cox et al., 1997; Massol et al., 1998;
- 4 Schlam et al., 2015). In contrast to Rac1, active CDC42 is restricted to the
- 5 protrusive cup rim in macrophages indicating differential regulation and
- 6 functionality (Hoppe and Swanson, 2004). In *Dictyostelium* however, no clear
- 7 CDC42 orthologue has been identified.
- 8

9 Cup formation requires integrated spatio-temporal control over multiple

10 GTPases. This must be able to self-organise in the absence of external cues

- 11 during macropinocytosis, and robust enough to phagocytose physiological
- 12 targets of varying size and shape. Small GTPase activity is controlled by a large
- 13 family of proteins such as GTPase Exchange Factors (GEFs) which promote the
- 14 GTP-bound active form, and GTPase Activating Proteins (GAPs) which stimulate
- 15 hydrolysis and transition to a GDP-bound inactive state. In this study, we
- 16 characterise a previously unstudied dual GEF and GAP protein in *Dictyostelium*
- 17 that integrates Ras, Rac and lipid signaling. This provides a mechanism to
- 18 coordinate the cup interior with the protrusive rim, allowing efficient
- 19 macropinosome formation and the engulfment of diverse bacteria of differing
- 20 geometry.

21 **Results**

22 Identification of a novel BAR-domain containing protein recruited to cups

23

24 Our initial hypothesis was that cells may use the different membrane curvature 25 at the protrusive rim compared to the cup base to recognise and differentially 26 regulate cup shape. Membrane curvature can recruit specific proteins containing 27 BAR (Bin-Amphiphysin-Rvs) domains (Peter et al., 2004). These are often found 28 in multidomain proteins, including several involved in GTPase regulation and 29 trafficking (Aspenstrom, 2014). To identify candidate proteins involved in 30 macropinocytosis, we therefore searched the *Dictyostelium* genome for BAR domain-containing proteins. Excluding proteins of known localisation or 31 32 function, we systematically cloned each candidate and expressed them as both N-

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1 and C- terminal GFP-fusions in axenic Ax2 cells. Using this strategy we 2 successfully cloned 9 previously uncharacterised BAR-containing proteins and 3 observed their localisation in live cells by fluorescence microscopy. Of these, 6 4 were expressed at detectable levels (Figure 1A). 5 6 DDB G0284997, DDB G0305372 and DDB G0285851 were associated with small puncta at the plasma membrane, consistent with the well-characterised role of 7 BAR domain proteins in clathrin mediated endocytosis (Dawson et al., 2006). 8 9 DDB_G0276447 localised to intracellular vesicles too small to be 10 macropinosomes, and GFP-DDB G0272368 was exclusively observed in the 11 nucleus. Only one of the proteins tested (DDB G0269934) localised to what 12 appeared to be the protrusive regions of macropinocytic cups. 13 14 DDB_G0269934 is a 223 kDa multidomain protein and also contains Regulator of 15 Chromatin Condensation (RCC1), RhoGEF and RasGAP domains (Figure 1B). 16 DDB_G0269934 has not previously been described and due to its domain 17 organisation we will subsequently refer to it as RGBARG (RCC1, GEF, BAR and 18 GAP domain containing protein). How Ras and Rac activity are coordinated in 19 space and time to generate a 3-dimensional cup shaped protrusion is not known. 20 Combining BAR, GEF and GAP activities in a single protein potentially provides 21 an elegant mechanism to organise engulfment. Therefore the function and 22 regulation of RGBARG was investigated in detail. 23 24 Examining RGBARG-GFP dynamics by timelapse fluorescence microscopy 25 confirmed strong enrichment at the protrusive rim of both macropinocytic and 26 phagocytic cups that disappeared rapidly after engulfment (Figure 1C and D, 27 Videos 1 and 2). Co-expression with the PIP₃ reporter PH_{CRAC} -RFP that demarks 28 the cup interior confirmed RGBARG-GFP localised specifically to the periphery of 29 this signaling domain (Figure 1E-H, and Video 3). Importantly, this differs from 30 the RasGAP NF1, which localises throughout the cup interior (Bloomfield et al.,

31 2015). RGBARG may therefore play a specific role in organising cup dynamics32 and engulfment.

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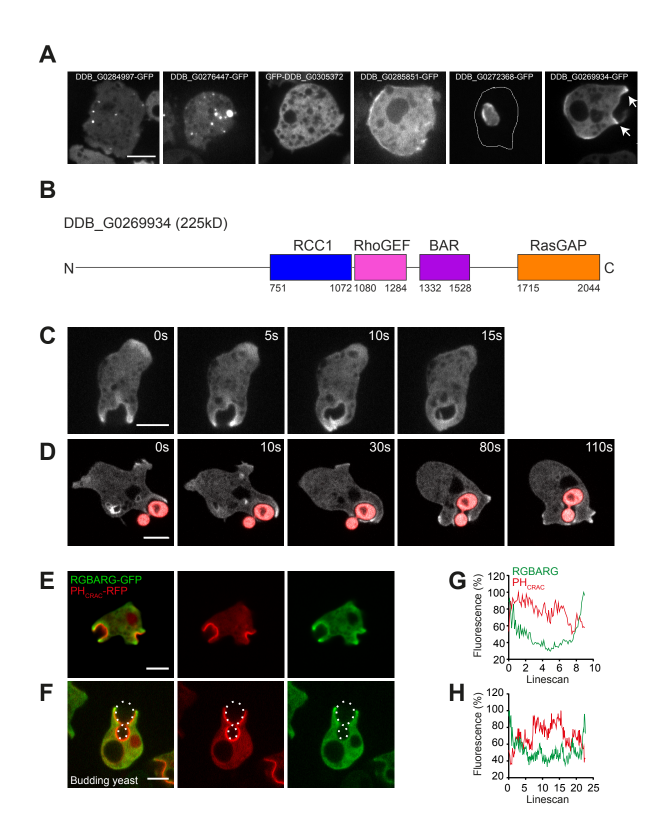


Figure 1: Identification of BAR domain proteins associated with macropinocytosis. (A) Uncharacterised BAR-domain containing proteins were expressed as GFP-fusions in Ax2 Dictyostelium cells. Images are maximum intensity projections of confocal Z-stacks. (B) Illustrates the domain organisation of DDB_G0269934/RGBARG. (C) Time series of spinning disc images of RGBARG-GFP recruitment during macropinocytosis and (D) phagocytosis of TRITC-labeled heat-killed budding yeast. (E) and (F) show the localization of RGBARG-GFP relative to the PIP₃ reporter PH_{CRAC}-RFP during macropinocytosis respectively. (G) and (H) show the relative intensity profiles of each fluorescent protein in these images. Linescans were drawn along the cup interior from protrusive tip-to-tip. All scale bars denote 5 µm.

1 RGBAR regulates cup signaling and macropinosome formation

2

To test for a functional role in engulfment a 3.6 Kb central section of *RGBARG*gene (containing the RCC1 RhoGEF and BAR domains) was deleted and replaced
with a blasticidin selection cassette by homologous recombination (Figure 2
supplement 1). Independent clones were isolated (JSK02 and 03) and shared
comparable phenotypes. In the following experiments strain JSK02 was used
unless otherwise stated with effects of *RGBARG* mutation validated by rescue
experiments.

10

11 To check for defects in macropinosome formation, cells were incubated with 12 FITC-dextran, a pH sensitive dye that is quenched at low pH. As macropinosomes acidify in under two minutes in *Dictyostelium*, and images were acquired within 13 14 the 30 minutes required to transit to neutral post-lysosomes, intracellular FITC-15 dextran is only visible in nascent macropinosomes (Figure 2A). From confocal Z-16 stacks of live cells, we found that *RGBARG*⁻ cells formed significantly smaller 17 macropinosomes than parental controls, measuring $0.5\pm0.1 \ \mu m^3$ compared to $1.5\pm0.2 \ \mu\text{m}^3$ in Ax2 (Figure 2B). This phenotype could be completely rescued by 18 19 re-expression of RGBARG-GFP. *RGBARG*⁻ cells also produced more 20 macropinosomes (2.6 ± 0.2 per cell compared to 2.2 ± 0.1 for Ax2) leading to no 21 net change in either total fluid uptake or axenic growth (Figure 2C-E). RGBARG is 22 therefore functionally important for the dynamics of macropinocytosis but not 23 essential for engulfment.

24

25 To understand why *RGBARG*⁻ cells form smaller macropinosomes we followed 26 their formation by fluorescence microscopy. Using the PH_{CRAC}-GFP reporter we 27 found in *RGBARG*⁻ cells had larger and more numerous patches of PIP₃ than 28 controls, averaging 3.8±1.1 patches per confocal section with an average length 29 of $5.5\pm2.4 \,\mu\text{m}$ compared to 1.6 ± 0.6 patches averaging $4.4\pm1.2 \,\mu\text{m}$ in Ax2 (Figure 30 3A-C). Similarly enlarged patches were also observed using the active Ras 31 reporter GFP-RBD (the Ras binding domain of Raf1, Figure 3B-C and Figure 3 32 Supplement 1A). During these experiments we also noted that *RGBARG*⁻ cells a 33 mild cytokinesis defect with $10 \pm \%$ containing >2 nuclei, compared to 5% of

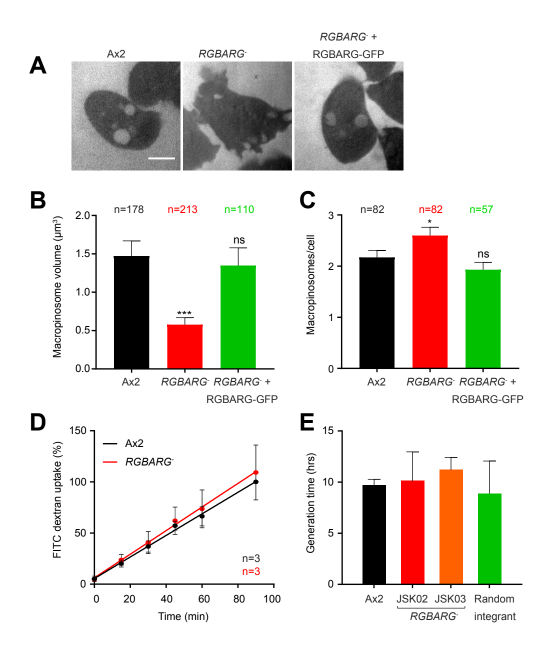


Figure 2: RGBARG- cells produce more, but smaller macropinosomes. (A) Confocal images of the indicated cell lines incubated in FITC-dextran for 10 minutes. The pH-sensitive FITC is only visible in pre-acidification macropinosomes <2 minutes after formation. The average volume of macropinosomes formed is shown in (B) and the number of macropinosomes per cell is shown in (C). n indicates the total number of macropinosomes or cells measured over 3 independent experiments. (D) Total fluid uptake measured by FITC dextran uptake over time, measured in a fluorimeter. (E) Growth rate of RGBARG mutants in axenic culture compared to the Ax2 parental cell line and a random integrant control from 3 independent experiments. All graphs show means \pm SEM, * P<0.05, *** P<0.001 as determined by Students T-test.

1 controls. This is consistent with the cytokinesis defects described in *PTEN*

2 mutants which also have excessive PIP₃ accumulation (Janetopoulos et al., 2005)

3 and all multinucleate cells were excluded from analysis.

4

5 To understand how the enlarged Ras and PIP₃ patches in *RGBARG*⁻ cells give rise 6 to smaller macropinosomes we studied their formation over time in 3D. As recently described, the macropinocytic cups of Ax2 cells form by expanding 7 8 around a defined spontaneous patch of PIP_3 (Veltman et al., 2016). These cups 9 subsequently close, usually forming one, or sometimes two, large 10 macropinosomes accompanied by termination of PIP₃ signaling on both the new 11 vesicle and the cell surface (Figure 3D and Video 4). This process is relatively 12 consistent, with each PIP₃ patch lasting an average of 150 seconds (Figure 3 Supplement 1B). In *RGBARG*⁻ cells however, whilst PH_{CRAC}-GFP still disappeared 13 from internalised vesicles, the plasma membrane domains were much more 14 15 stable. Whilst PIP₃ patches frequently split, they rarely dissipated completely 16 and often lasted longer than each of the 30 minute movies recorded (Figure 3E 17 and Video 5). It was therefore not possible to meaningfully measure the lifetime 18 of surface PIP₃ (and by extension Ras) signaling in *RGBARG*⁻ cells. As RGBARG is 19 restricted to the periphery of Ras signaling domains it appears to restrict both 20 lateral expansion of activated Ras and termination of Ras/PIP₃ signaling upon 21 cup completion.

22

23 Although extinction of PIP₃ signaling did not accompany cup closure in *RGBARG*-24 cells, numerous small vesicles could be observed continuously budding from the 25 base of the ruffles when folds of membrane collapsed in on themselves. This 26 explains why these cells form more frequent but smaller macropinosomes 27 (Figure 2). We speculate that this indicates that the entirety of the PIP₃ patch is 28 potentially fusogenic and can internalise vesicles by simply folding onto itself 29 rather than requiring a specific mechanism to orchestrate closure and fission at 30 the rim. How this might be achieved mechanistically is unclear, but is 31 reminiscent of the less organised, more ruffle-like macropinosome formation 32 observed in serum stimulated mammalian epithelial cells, or Ras transformed 33 cancer cell lines (Williamson and Donaldson, 2019).

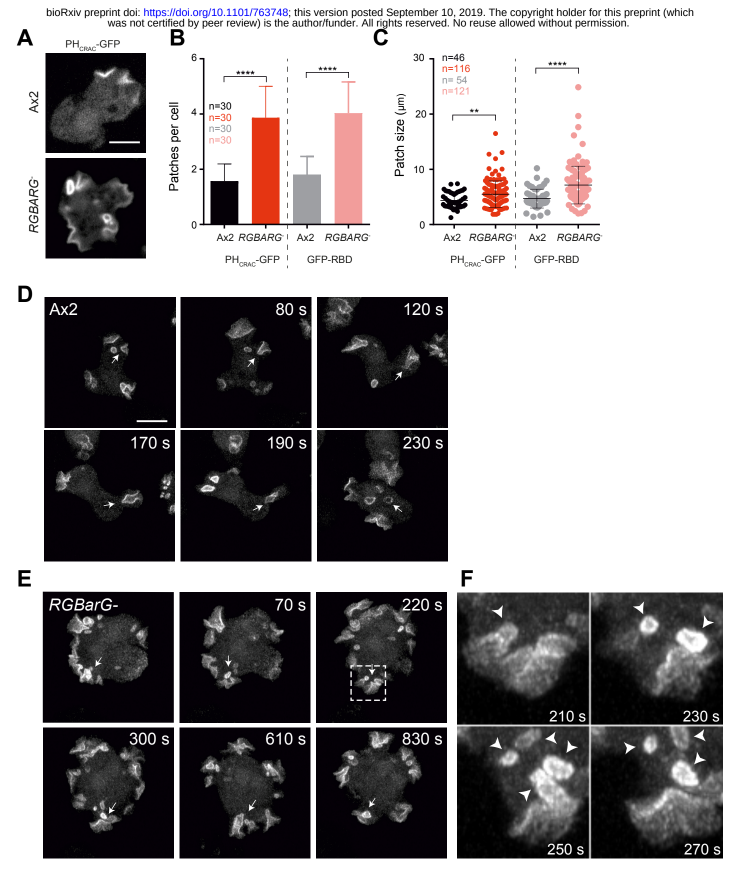


Figure 3: Dynamics of macropinosome formation in RGBARG- cells. (A) Membrane localization of the PIP_{3} probe PH_{CRAC} -GFP. Images are single confocal sections taken by spinning disc microscopy. The number of patches of PH_{CRAC} -GFP or the active Ras probe GFP-RBD per cell in single plane through the middle of each cell is quantified in (B). The average size of each patch is shown in (C). n is the total number of cells or patches measured over 3 independent experiments. Error bars show the mean ± standard deviation, ** P<0.01, *** P<0.001, Mann-Whitney T-test. (D) and (E) Time series of a maximum intensity projection through the entire depth of PH_{CRAC} -GFP expressing cells. (D) Shows Ax2 cells, indicating the formation, closure and subsequent extinction of PH_{CRAC} -GFP patch at the cell surface (arrow). (E) Shows an equivalent movie of RGBARG- cells, where the PH_{CRAC} -GFP patch remains after closure (arrow). (F) Is an enlargement of the boxed area in (E), showing multiple vesicles forming from a single, large PH_{CRAC} -GFP patch (arrow-heads). All scale bars indicate 5 µm.

1

GEF, GAP and BAR domain interactions each contribute to RGBARG localisation

4

Localisation of RGBARG at the interface between the cup interior and the
protrusive rim is likely to be critical to effectively control the shape and
dynamics of these domains during engulfment. This will position its RhoGEF
activity where protrusion is promoted and its RasGAP activity where it can
restrain expansion of the interior, leading to the organised cup formation
observed in Ax2 cells and absent in *RGBARG* - mutants.

11

12 To dissect the mechanisms of RGBARG recruitment we tested the effect of 13 deleting each protein domain in turn. To quantify RGBARG enrichment across the cup, linescans from cup tip to tip were normalised to non-protruding regions 14 15 of the same cell and averaged across multiple cells (Figure 4A and B). GFP-fused to the cyclic AMP receptor (cAR1-GFP) localises uniformly to the plasma 16 17 membrane and was used as a control (Figure 4C and D). This method confirmed 18 RGBARG-GFP was enriched 3-fold at the protruding edges of macropinocytic 19 cups and allowed us to quantify how each protein domain contributes to 20 recruitment at the cup (Figure 4 and supplement).

21

22 Removal of the RCC1 domain had no effect on localisation and was able to fully 23 rescue the ability of *RGBARG*⁻ cells to form large macropinosomes (Figure 4E-H). 24 In contrast, deletion of either the RhoGEF or BAR domains caused RGBARG to 25 become uniformly cytosolic and did not rescue (Figure 4E-H). In the absence of 26 the RasGAP domain however, RGBARG was still recruited to the plasma 27 membrane but was much more broadly distributed throughout the cup and 28 significantly less enriched at the protruding rim (Figure 4G). RGBARG∆GAP-GFP 29 was also unable to rescue macropinosome formation (Figure 4H). Co-expression 30 of PH_{CRAC}-RFP confirmed that RGBARG∆RasGAP-GFP was no longer excluded 31 from PIP₃, and therefore active Ras domains (Figure 4 supplement 2A). RasGAP 32 interactions therefore restrict RGBARG to the periphery of the cup interior 33 domain.

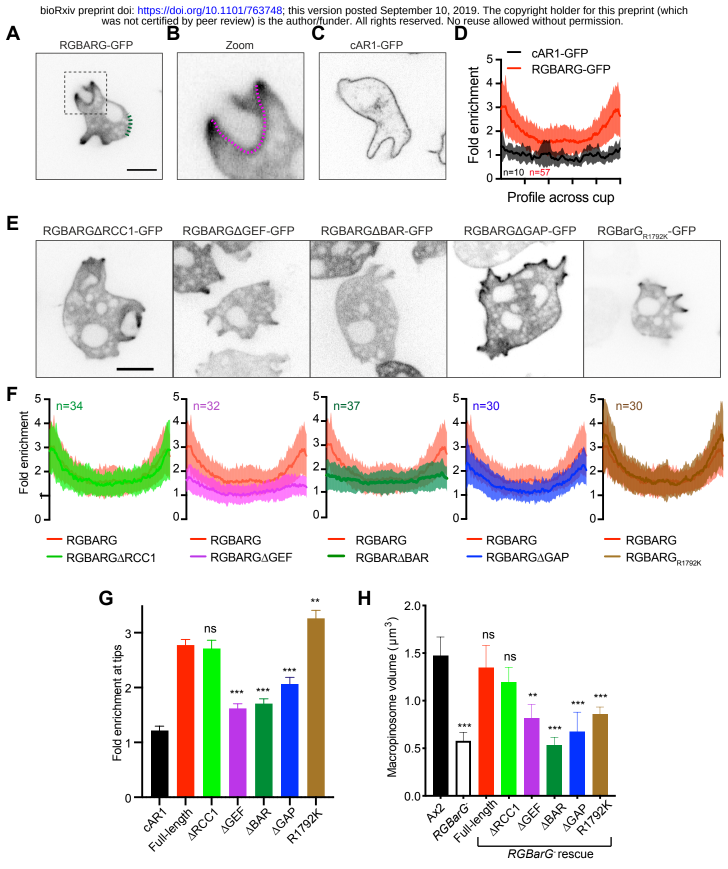


Figure 4: Multiple interaction regulate RGBARG recruitment at the cup. Full-length RGBARG or mutants lacking each domain in turn were expressed as GFP-fusions in RGBARG- cells. (A) Shows an example of full-length RGBARG-GFP. Enrichement was measured relative to the average intensity of a non-protrusive membrane region (green dotted line). The boxed region is enlarged in (B), showing an example of the line measured from the rim along the cup interior. (C) Shows the uniform localization of cAR1-GFP used as a control. (D) Averaged, normalized linescans along the cup from multiple cells, demonstrating a 3-fold enrichment of RGBARG-GFP at the cup rim and uniform cAR1-GFP concentration. (E) Shows representative images of cells expressing RGBARG-GFP with the domains indicated delected, as well as the R1792K point mutation that inactivates RasGAP activity. The averaged intensity of each construct across the cup is shown in (F), with the profile of the full-length protein from (D) in red for comparison. Values plotted are the mean \pm standard deviation. (G) The enrichment at the protruding rim of each construct measured by averaging the first 10% of each individual linescan and calculating the mean and SEM across each group. (H) The ability of each construct to rescue large macropinosome formation in RGBARG- cells was determined by measuring the size of nascent FITC dextran-containing macropinosomes by confocal microscopy, as in Figure 2A. >100 macropinosomes over 3 independent experiments were measured. Bars denote mean macropinosome volume \pm SEM, ** P<0.005 Mann-Whitney T-test.

1 2 To confirm the role of the RasGAP interactions in restricting RGBARG 3 localisation, we also made a point mutant in the conserved arginine responsible 4 for stabilising the transition from Ras-GTP to Ras-GDP(Bos et al., 2007). This 5 mutation (R1792K) is predicted to disrupt GAP activity but still allow Ras 6 binding and completely rescued RGBARG exclusion from the cup interior, and 7 slightly but significantly increased enrichment at the cup tip (Figure 4F-G and 8 supplement 2B). However, despite localising to the protruding rim, 9 RGBARG_{R1792K}- GFP did not rescue the cup organisation of *RGBARG*⁻ cells, which 10 still produced enlarged PIP₃ patches and small macropinosomes (Figure 4H and 11 Supplement 2). RGBARG is therefore an active RasGAP and this domain also 12 provides spatial information to position RGBARG to the periphery of the active 13 Ras/PIP₃ patch where it can prevent its expansion and shape the forming cup. 14 15 The data above show that RGBARG integrates spatial cues from its RhoGEF, BAR 16 and RasGAP domains for correct positioning and cup organisation. To identify 17 the relevant binding partners and contribution of each domain, we also 18 expressed them individually fused to GFP. Whilst RhoGEF-GFP expressed too 19 poorly to observe its localisation, both the RCC1 and GAP domains were 20 completely cytosolic (Figure 5A and C). In contrast, the BAR domain alone was 21 sufficient for strong recruitment throughout the plasma membrane (Figure 5B). 22 This was blocked by including either of the adjacent RhoGEF or RasGAP domains 23 (Figure 5D and E), suggesting that BAR-domain binding may also be regulated by 24 intramolecular interactions. In contradiction of our initial hypothesis however, 25 BAR-GFP was not enriched at areas of curvature or protrusion. The BAR domain 26 therefore appears to drive general recruitment to the plasma membrane rather 27 than recognising curvature at cups. 28 29 As the BAR domain of RGBARG does not concentrate at specific membrane 30 shapes, we investigated its lipid binding specificity by lipid-protein overlay. 31 Incubation of lysates from cells expressing BAR-GFP with PIP strips indicated

- 32 binding to all PIPs with two or more phosphates (Figure 5F). This was confirmed
- 33 by PIP array, indicating a slight selectivity for PI(3,4)P₂ (Figure 5 Supplement

1 A). Given this broad ability to bind all highly phosphorylated phosphoinositides
 it is likely that this BAR domain generally recognises their high negative charge
 rather than specific phosphate configurations. This supports a mechanism
 whereby highly-phosphorylated PIPs recruit RGBARG to the plasma membrane
 via its BAR domain where additional interactions with the RhoGEF and RasGAP
 domains further restrict its position and activity to the protruding edges of
 forming cups.

8

9 To identify the targets of the RhoGEF domain, we performed co-

10 immunoprecipitations with a library of recombinant GST-tagged small GTPases.

11 The *Dictyostelium* genome contains an expanded set of Rac small GTPases, but no

12 Rho or CDC42 subfamily members (Vlahou and Rivero, 2006). Of these only RacH

13 and RacG bound the RhoGEF domain of RGBARG with no detectable binding to

14 other Racs, including Rac1 which has previously been implicated in cup

- 15 formation (Dumontier et al., 2000).
- 16

17 Whilst RacH is involved primarily in endocytic trafficking and localises

18 exclusively to intracellular compartments (Somesh et al., 2006a), RacG localises

19 to the plasma membrane and is enriched at the protruding rim of phagocytic

20 cups (Somesh et al., 2006b). Previous studies have also shown that

21 overexpression of wild-type or constitutively active RacG also promotes

22 phagocytosis, indicating a potential interaction with RGBARG (Somesh et al.,

23

2006b).

24

25 Consistent with previous reports, we found loss of RacG had no significant effect 26 on macropinocytosis, with mutants forming normal sized active Ras patches and 27 macropinosomes (Somesh et al., 2006b)(Figure 5 supplement 1B and C). When 28 we measured RGBARG-GFP recruitment, its association with cups was more 29 uniform and was only enriched 1.8±0.6 fold at the rim in *RacG*- cells compared 30 to 2.6±0.7 fold isogenic controls (Figure 5H-L). This indicates that RacG and 31 RGBARG functionally interact in vivo and partly contribute to RGBARG 32 localisation. However, the remaining signals or functional redundancy with other

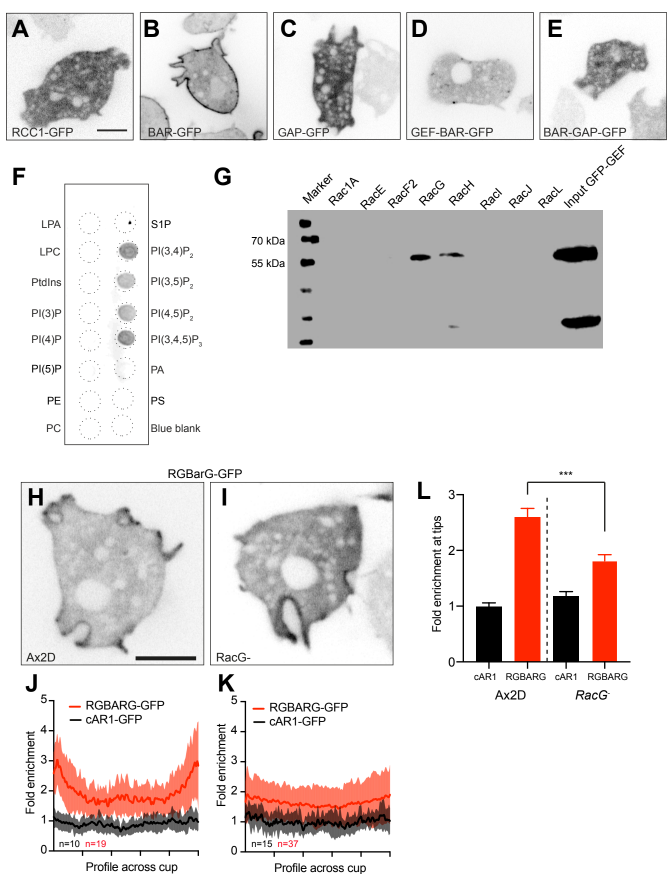


Figure 5: Binding specificity of the BAR and GEF domains. (A-E) Individual, or combinations of domains from RGBARG were expressed as GFP fusions in RGBARG- cells. Images shown are single confocal sections. (F) Lipid binding specificity of BAR-GEF by lipid overlay assay using whole cell lysate from BAR-GFP expressing cells. (G) Rac binding specificity of the RhoGEF domain determined by co-immunoprecipitation of GEF-GFP by a library of purified GST-Rac's bound to beads. (H) and (I) Confocal images of full-length RGBARG-GFP localisation in RacG- cells and their parental background strain Ax2D. The average profile (± standard deviation) of RGBARG-GFP along the cup relative in Ax2D and RacG- cells relative to cAR1-GFP is shown in (J) and (K) respectively. (L) Enrichment of RGBARG-GFP and cAR1 at the cup tip in each cell line. Bars indicate mean ± SEM, *** P<0.005 Mann-Whitney T-test. All scale bars indicate 5 µm.

1 Racs is sufficient for partial RGBARG recruitment and apparently normal

- 2 engulfment in the absence of RacG.
- 3

4 Combined, our data indicate that RGBARG uses a coincidence detection 5 recruitment mechanism to direct cup formation: BAR domain binding to 6 negatively charged phospholipids directs the protein to the plasma membrane 7 whilst additional interactions with RacG and active Ras synergise to constrain 8 RGBARG to the cup rim. This tripartite regulation ensures that RGBARG is 9 accurately positioned to exert its RhoGEF and RasGAP activities at the interface 10 between cup interior and protrusion and organise engulfment. 11 12 RGBARG is a highly active dual specificity Ras/Rap GAP 13 14 Forming a complex 3-dimensional shape is likely to require multiple regulators 15 and RGBARG is not the only RasGAP involved in macropinocytosis in 16 Dictyostelium. Axenic strains (including the Ax2 parental strain used in this 17 work) also harbour mutations in NF1 that enhance fluid uptake via enlarged 18 active Ras patches and subsequent formation of larger macropinosomes 19 (Bloomfield et al., 2015). However, whilst mutations in NF1 were reported in 20 each of 10 independently isolated axenic strains, RGBARG mutations were 21 completely absent ((Bloomfield et al., 2015) and G. Bloomfield, personal 22 communication, May 2019). Our attempts to generate RGBARG mutants in non-23 axenic strains were also unsuccessful. This indicates that disruption of NF1 but 24 not *RGBARG* is sufficient to support axenic growth, and that they serve different 25 functions. 26

27 To better understand the differences between NF1 and RGBARG, we compared 28 the specificity and activities of their RasGAP domains. The Dictyostelium genome 29 encodes 14 Ras subfamily members of which RasB, RasG and RasS are the most 30 important for macropinocytosis (Chubb et al., 2000; Hoeller et al., 2013; 31 Junemann et al., 2016; Khosla et al., 2000). Overexpression of RasD can also

- 32 partially compensate for loss of RasG and S (Khosla et al., 2000). The small
- 33 GTPase Rap, a close relative of Ras, has also been implicated in macropinosome

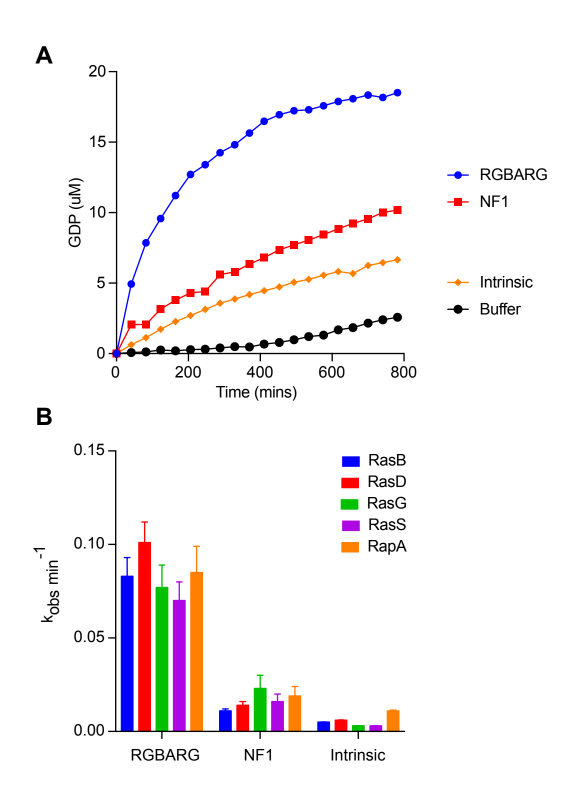


Figure 6: RasGAP activity of RGBARG and NF1. (A) Stimulation of GDP released from GTP-loaded RasG upon addition of recombinant RasGAP domains from either RGBARG or NF1, compared to the intrinsic GAP activity of the GTPase or GTP in buffer. (B) GAP activity of NF1 and RGBARG against a library of Ras superfamily members. Average of 3 independent experiments performed as in (A) in parallel. Bars indicate mean ± standard deviation.

1	formation (Inaba et al., 2017). We therefore measured the specific GAP activity
2	from both NF1 and RGBARG against each small GTPase.
3	
4	Consistent with the inability of $RGBARG_{R1792K}$ to rescue the knockout phenotype, we
5	found that the RasGAP domain of RGBARG was highly active against all the
6	GTPases tested (Figure 6). The RasGAP domain of NF1 was also active against each
7	Ras tested, but with 75% less activity than RGBARG in each case. RGBARG is
8	therefore a more potent RasGAP in vitro, but the lack of specificity for particular Ras
9	isoforms for both RGBARG and NF1 indicates their functional differences are likely
10	imparted by additional factors such as their cellular localisation and dynamics.
11	
12	Loss of RGBARG improves phagocytosis of large objects
13	
14	The data above demonstrate that RGBARG is important during the spontaneous
15	self-organisation of macropinocytic cups. As RGBARG also localises to
16	phagocytic cups and engulfment of solid particles such as microbes uses much of
17	the same machinery, we also investigated how RGBARG contributes to
18	phagocytosis.
19	
20	Disruption of NF1 has previously been shown to increase the size of particles
21	that Dictyostelium can engulf (Bloomfield et al., 2015). As RGBARG also affects
22	the size of the PIP_3 domains that define the cup interior we first tested the ability
23	of <i>RGBARG</i> - cells to phagocytose different sized beads. Whilst disruption of
24	RGBARG had no effect on phagocytosis of 1 μm diameter beads, engulfment of 4.5
25	μm beads was enhanced 3-fold, with an average of 2.2 ± 0.4 beads engulfed per
26	cell after 1 hour, compared to 1.0 ± 0.4 in control (Figure 7A and B). Enhanced Ras
27	activation therefore appears to be generally beneficial for the engulfment of
28	large spherical targets.
29	
30	Surprisingly, although expression of RGBARG-GFP from an extrachromosomal
31	vector fully rescued macropinosome formation (Figure 2A-C), this reduced the
32	ability of <i>RGBARG</i> ⁻ cells to engulf 4.5 μ m beads to 63% of control levels (Figure
33	7B). This effect was even more severe upon expression of domain deletion

13

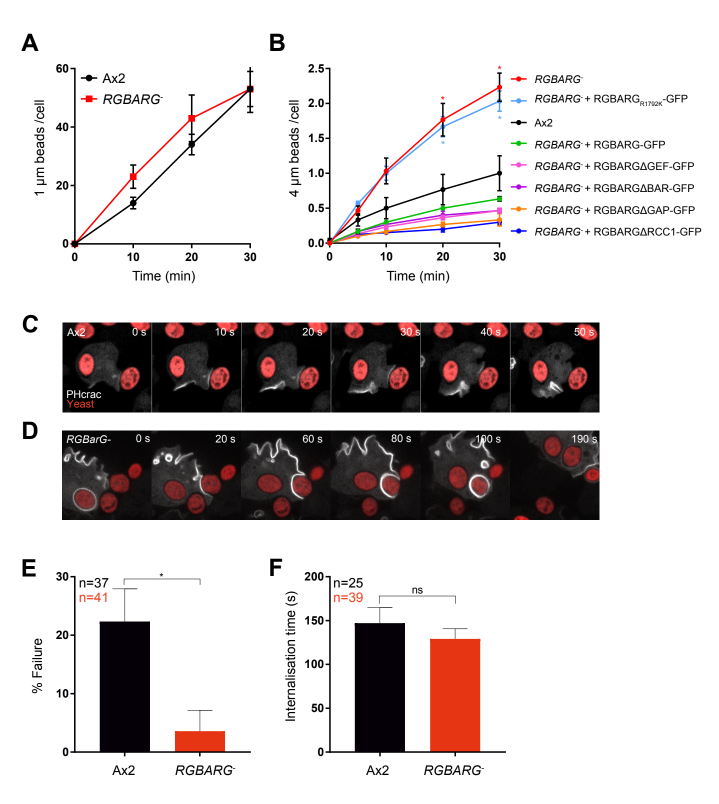


Figure 7: Phagocytic defects in RGBARG- cells. (A) Ax2 and RGBARG- cell have comparable rates of phagocytosis of fluorescent 1 μ m beads. (B) Phagocytosis of 4 μ m beads by Ax2 cells, RGBARG- cells, or RGBARG- cells expressing full-length or mutant RGBARG-GFP. Phagocytosis measured by flow cytometry in 3 independent experiments. Phagocytosis of TRITC-labelled yeast was directly observed by spinning disc confocal microscopy by either (C) Ax2 or (D) RGBARG- cells expressing PHCRAC-GFP. (C) Shows an example of a failed engulfment. (E) The relative frequency of phagocytosis failure after cup formation (indicated by PHCRAC-GFP recruitment). The time from initial contact to complete engulfment in successful phagocytic events is shown in (F). n indictates total number of phagocytic events over 3 independent experiments. All values plotted are mean \pm standard deviation. * P<0.01 Students T-test.

1 mutants including the Δ BAR, Δ GEF and Δ GAP constructs which do not localise 2 properly and have no deleterious effect on macropinosome formation. This 3 indicates a dominant negative effect, most likely due to sequestration of binding 4 partners. In contrast, expression of RGBARG_{R1792K} had no inhibitory effect on 5 *RGBARG*⁻ cells. This only differs from RGBARG-GFP in its RasGAP activity 6 indicating that mislocalisation or overexpression of this domain is sufficient to 7 inhibit engulfment of large targets. 8 9 To better understand how loss of RGBARG affects phagocytosis we performed 10 time-lapse microscopy of cells expressing PH_{CRAC}-GFP engulfing TRITC-labeled 11 yeast. Engulfment occurred rapidly in both cell types but failed at a frequency of 12 \sim 20% in Ax2 cells with the PIP₃ patch dissipating and the yeast escaping from 13 the cell (Video 6). Whilst the time for successful engulfment was no significantly 14 altered by loss of *RGBARG* (129±11 minutes in mutants vs 147±18 minutes in

15 Ax2), capture was much more robust with a failure rate of only 4±5 % compared

- 16 to 22± 7% (Figure 7C-F and Video 7). The main influence on the phagocytic
- efficiency of large targets in *RGBARG*⁻ cells thus appears to be increased cup
 stability and enlarged Ras signaling patch rather than rate of protrusion around
 the object.
- 20

Spatial regulation of Ras by RGBARG is important for phagocytosis of elongated targets

23

To be effective, phagocytic cells must be able to engulf microbes with differing
physical properties such as shape, size, stiffness and surface chemistry. As
RGBARG is important for the organisation and stability of phagocytic and
macropinocytic cups, we also investigated its role during the engulfment of
different bacteria.

29

Phagocytosis was measured by the ability of *Dictyostelium* cells to reduce the
 turbidity of a bacterial suspension over time. Whilst disruption of *RGBARG* had

32 no effect on the ability to clear a suspension of *Klebsiella aerogenes*, engulfment

33 of *Escherishia coli* was substantially reduced (Figure 8A and B). This was fully

1 rescued by re-expression of RGBARG-GFP. Therefore, although loss of RGBARG 2 has no effect on the engulfment of 1 μ m beads and is beneficial for the uptake of 3 large beads and yeast, it causes a specific defect in engulfment of some bacteria.

4

5 The most obvious physical difference between *K. aerogenes* and *E. coli* is their 6 shape (Figure 8C and D). Both have similar short axes but *K. aerogenes* average 7 3.2 µm in length whilst *E. coli* have an average long axis of 5.4 µm. Previous work 8 investigating phagocytosis of different shaped beads by macrophages concluded 9 that complex elongated shapes are more difficult to engulf (Champion and 10 Mitragotri, 2006). We therefore hypothesised that the selective phagocytosis 11 defects of *RGBARG*⁻ cells was due to the target shape. To test this, we measured 12 the ability of *RGBARG*⁻ cells to engulf an additional elongated rod-shaped bacteria 13 (GFP-expressing *Mycobacteria smegmatis*, 3-5 µm long) by flow cytometry. The ability of *RGBARG*⁻ cells to engulfing these bacteria was again reduced by 75% 14 15 (Figure 8E), again correlating with an inability to phagocytose elongated targets. 16

17 The data above are consistent with a role for RGBARG in enabling the engulfment of elongated bacteria. However, the bacterial strains used will also differ in other 18 aspects such as their surface components, phagocytic receptor activation and 19 20 stiffness. To directly test the importance of RGBARG in engulfing targets of 21 different shape we therefore stretched 3 µm latex beads to generate oblate 22 ellipsoids of conserved volume and surface chemistry (Ho et al., 1993).

23

24 To measure relative phagocytosis in the same experiment, cells were incubated 25 with a 1:1 mix of spherical and stretched beads (2.6x aspect ratio) and the 26 number of engulfed beads of each shape quantified by microscopy. The ability of 27 Ax2 cells to engulf ellipsoid particles was reduced by 30% compared to spheres 28 (P<0.01, T-test). However, whilst *RGBARG*⁻ cells engulfed the spheres with similar 29 efficiency, the number of stretched beads taken up was reduced by over 70% 30 (Figure 8C, P<0.01, T-test). These effects were again rescued by re-expression of 31 RGBARG-GFP, but not RGBARG_{R1792K}-GFP demonstrating a key role for the 32 RasGAP activity in mediating phagocytosis of elongated particles.

33

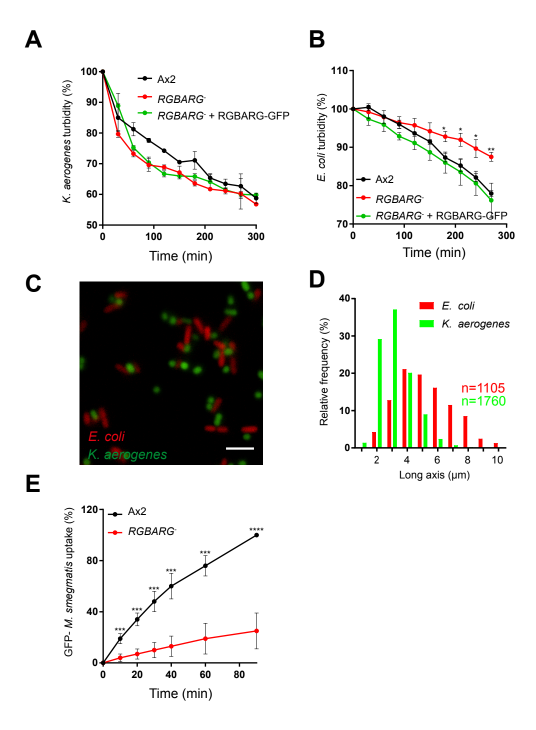


Figure 8: RGBARG- cells have selective defects in phagocytosis of bacteria. Phagocytosis of (A) K. aerogenes or (B) E. coli was measured by monitoring the decreasing turbidity of a bacterial suspension after addition of Dictyostelium. (C) show wide-field fluorescence microscopy images of GFP-expressing K. aerogenes mixed with RFP-expressing E. coli, demonstrating their different shape and size. The length of each bacteria was quantified automatically and is plotted in (D). (E) Phagocytosis of GFP-M. smegmatis, an alternative elongated bacteria, measured by flow cytometry. Values plotted in (A), (B) and (E) are mean \pm standard deviation of 3 independent experiments. *P<0.05, **P<0.01, ***P<0.005, Student's T-test.

- 1 How phagocytic cups organise and adapt their cytoskeleton to engulf targets of
- 2 differing geometry is very poorly understood. Our data demonstrate that in
- 3 *Dictyostelium*, a tripartite recruitment mechanism operates to precisely position
- 4 a RasGAP and RhoGEF domain-containing protein at the interface between the
- 5 protrusive rim and static interior domains of phagocytic and macropinocytic
- 6 cups. We propose a model whereby this organises the cup by regulating the
- 7 balance between protrusion and expansion of the interior in order to both
- 8 efficiently form macropinosomes and facilitate engulfment of geometrically
- 9 diverse targets.

1 Discussion

2

3 In this study we have identified a new component and mechanism used by cells 4 to organise their protrusions into the 3-dimensional cup shapes required to 5 engulf extracellular fluid or particles. Consistent with previous studies, our data 6 support a model whereby cup formation is guided by the formation of a 7 protrusive rim encircling a static interior domain (Veltman et al., 2016). We 8 show that in *Dictvostelium*, RGBARG provides a direct link between the Ras and 9 Rac activities that underlie these different functional domains, providing a novel 10 mechanism to co-ordinate cup organisation in space and time. 11 12 RGBARG is not the only RasGAP in *Dictyostelium* involved in macropinosome 13 formation. However, it is the only RasGAP in the Dictyostelium genome to also 14 possesses a RhoGEF domain. RGBARG is therefore unique in its ability to 15 integrate the activities of both GTPase families. There are no human proteins 16 with an identical domain structure to RGBARG, and whilst most classical 17 RasGAPs are found in multidomain proteins, none also contain a classical 18 RhoGEF domain (Bos et al., 2007). A screen for RhoGAPs involved in phagosome 19 formation in macrophages identified 3 proteins (ARHGAP12, ARHGAP25 and 20 SH3BP1), but although they all contain PIP₃ binding (PH) or BAR domains, none 21 contain domains that link to other GTPase families (Schlam et al., 2015). The 22 oncogene TIAM1 contains both RhoGEF and Ras-binding domains however, and 23 BAR domains are found in conjunction with GAP or GEFs in several other 24 proteins. Therefore whilst mammalian cells also need to coordinate Ras and Rac 25 activity during cup formation, this is likely achieved via multiple proteins, 26 potentially in a complex.

27

Multiple GAP's and GEF's collaborate to shape protrusions into cups. This is
apparent in the different roles played by RGBARG and NF1. Both are important
negative regulators of Ras, but whilst RGBARG is specifically enriched at the cup
rim, NF1 appears to be present thoughout the cup (Bloomfield et al., 2015).
RGBARG and NF1 therefore play different functional roles; whilst disruption of

1 NF1 leads to an increase in the volume of fluid taken up and can facilitate axenic

2 growth (Bloomfield et al., 2015), RGBARG appears more important for cup

3 structure and shape. We therefore speculate a model whereby NF1 acts to

4 generally suppress Ras activity and regulate the spontaneous excitability of

- 5 active Ras patches, whilst RGBARG operates at their periphery to restrict their
- 6 expansion and stimulate protrusion via Rac.
- 7

8 This model is doubtless overly simplistic, and other RasGAP's also contribute to 9 shaping active Ras dynamics. For example, the IQGAP-related protein IqgC was 10 also recently shown to have RasGAP activity and localise throughout the interior 11 of macropinocytic and phagocytic cups in *Dictyostelium* (Marinovic et al., 2019). 12 In contrast to our findings with NF1 and RGBARG however, IggC is reported to 13 have specific RasGAP activity against RasG. As the different Ras isoforms are 14 non-redundant (Khosla et al., 2000), IqgC adds a further layer of regulatory 15 complexity shaping the dynamics of engulfment.

16

17 Whilst the regulation of Ras signaling and the static interior domain is becoming 18 clearer, how protrusion is regulated during engulfment is less well understood. 19 In mammalian cells, several studies indicate that actin dynamics and protrusion 20 at the cup is regulated by the combined activities of Rac1 and CDC42 (Cox et al., 21 1997; Massol et al., 1998; Schlam et al., 2015). Rac1 and CDC42 are differentially 22 activated with active Rac1 throughout the cup and CDC42 activation earlier and 23 more restricted to the rim (Hoppe and Swanson, 2004). Whilst Dictyostelium 24 does not possess a clear CDC42 orthologue, we find that RGBARG specifically 25 interacts with the atypical Rac isoforms RacG and RacH. Currently, no direct 26 effectors of either protein are known, and RacG does not interact with the Rac-27 binding domain of PAK commonly used as a probe for active Rac1 (Somesh et al., 28 2006b). RacG therefore has at least partly distinct effectors to Rac1. Nonetheless, 29 in cell-free assays, RacG is able to induce actin polymerisation via the ARP2/3 30 complex, although whether this is dependent on SCAR/WAVE or other WASP 31 family members is not known (Somesh et al., 2006b). RacG therefore appears to

1 be at least partly responsible for defining the protrusive rim, possibly through

2 some coincidence-detection mechanism with active Rac1.

3

4 Whilst RacG has no clear direct orthologue in mammalian cells, it is most similar to Cdc42 in protein sequence. Whilst constitutively active Rac1 induces the 5 6 formation of lamellipodial-type protrusions (Dumontier et al., 2000), constitutively active RacG and Cdc42 both induce filopodia (Nobes and Hall, 7 8 1995; Somesh et al., 2006b). We therefore speculate that RacG and Cdc42 are 9 functional orthologues, and the mechanism by which these small GTPases 10 integrate with active Ras to restrict the localisation of RhoGEF and RasGAP 11 proteins such as RGBARG is a general device used to define the protrusive cup 12 rim. 13 14 The involvement of both RacG/Cdc42 and Rac1 family GTPases indicates a 15 complex relationship between filopodial and lamellipodial type protrusions 16 during cup formation. Whilst most studies in *Dictyostelium*, RAW macrophages, 17 dendritic cells and cancer cell lines describe macropinosome formation from

18 smooth, sheet-like projections or cups (Swanson, 2008; Veltman et al., 2016;

19 West et al., 2000; Williamson and Donaldson, 2019), it was recently shown that

20 RAW macrophages can also form macropinosomes by a more filopodial "tent-

21 pole"-type mechanism where protrusion is driven by actin-rich spikes (Condon

et al., 2018). Whether this is a general mechanism, or represents a shift in the

23 balance of filopodial vs lamellipodial regulatory proteins in these cells is unclear.

However it is probable that filopodial and lamellal cup formation are non-

25 exclusive extremes of a continuum - much as they are during cell migration.

26

The multi-layered regulation of small GTPases is particularly important when
cells are challenged to engulf particles or microbes of different shapes. This is
critical for amoebae to feed on diverse bacteria or immune cells to be able to
capture and kill a wide range of pathogens, but how cells adapt to different target
geometries is very poorly understood (Champion and Mitragotri, 2006;
Champion and Mitragotri, 2009). To our knowledge, *RGBARG*⁻ cells are the first
mutants described with a geometry-specific defect in phagocytosis, underlining

- 1 the importance of co-ordinating and balancing Ras and Rac activities. This again
- 2 differs from the role of NF1, as the *NF1*-deficient Ax2 strain used in this study is
- 3 able to efficiently engulf and grow on a wide range of bacteria including
- 4 elongated strains such as *E. coli* (Buckley et al., 2019). It is still not known how
- 5 other regulatory elements or cytoskeletal components adapt to differing shapes,
- 6 but it seems likely that large-scale rearrangements are necessary to
- 7 accommodate different targets.
- 8
- 9 In summary, we describe a mechanism to co-ordinate the activity of Rac and Ras
- 10 family GTPases during engulfment in *Dictyostelium*. The proteins that mediate
- 11 this co-ordination in mammalian cells remain unknown. However, we propose a
- 12 general model by which spatial signals and effectors from multiple small
- 13 GTPases integrate to shape the protrusions that form macropinocytic and
- 14 phagocytic cups, enabling cells to engulf diverse targets.

1 Methods

2 Dictyostelium culture and molecular biology

3 Unless otherwise stated, Dictyostelium strains were derived from the MRC-Ax2 4 axenic strain provided by the Kay laboratory and were routinely cultured in filter 5 sterilised HL-5 medium (Formedium) at 22°C. RacG mutants and corresponding 6 parental strain (from the Devreotes group, Johns Hopkins, Ax2D) were kind gifts 7 from Francisco Rivero (University of Hull)(Somesh et al., 2006b). Growth rates 8 were measured by seeding cells at 0.5×10^5 /ml in HL-5 and counting cell number 9 twice daily for three days. Growth rate was then calculated by fitting an 10 exponential growth curve using Graphpad Prism software. Cells were transformed by electroporation: 6 x 10⁶ cells were resuspended in 0.4 mls of ice 11 12 cold E-buffer (10 mM KH₂PO₄ pH 6.1, 50 mM sucrose) and transferred to 2 mm electroporation cuvette containing DNA (0.5 µg for extrachromosomal plasmids, 13 14 15 μ g for knockout vectors). Cells were then electroporated at 1.2 kV and 3 μ F 15 capacitance with a 5 Ω resistor in series using a Bio-Rad Gene Pulser II. After 24 16 hours transformants were selected in either 20 µg/ml hygromycin (Invitrogen), 17 10 μg/ml G418 (Sigma) or 10 μg/ml blasticidin (Melford).

18

19 BAR domain contain proteins were identified by multiple BLAST searches using 20 Dictybase (www.dictybase.org) (Fey et al., 2013). Coding sequences were then 21 amplified by PCR from vegetative Ax2 cDNA adding compatible restriction sites 22 for subcloning into the BglII/SpeI sites of the N- and C- terminal GFP-fusion 23 Dictyostelium extrachromosomal expression vectors pDM1043 and pDM1045 (non-axenically selectable versions of the pDM modular expression system 24 25 (Veltman et al., 2009)). Truncation and point mutants of RGBARG were also 26 generated by PCR and expressed using pDM1045. The *RGBARG* (DDB G0269934) knockout construct was generated by PCR fusion of \sim 1Kb 5' and 3' 27 28 recombination arms with the floxed blasticidin selection cassette from 29 pDM1079, as described in detail in (Paschke et al., 2018). After transformation, 30 independent clones were obtain by dilute plating in 96 well plates. Disruption of 31 the RGBARG locus was screened by PCR from genomic DNA isolated from 1 x 10⁶ 32 cells lysed in 100 µl 10mM Tris- HCl pH8.0, 50 mM KCl, 2.5mM MgCl1, 0.45% 33 NP40, 0.45% Tween 20 and 0.4 mg/ml Proteinase K (NEB). After 5 minutes

1 incubation at room temperature, the proteinase K was denatured at 95°C for 10

2 minutes prior to PCR analysis. The Ras binding domain (RBD) of PAK1-GFP

3 construct used as an active Ras reporter was a gift from Gareth Bloomfield.

4

5 **Macropinocytosis assays**

6 To measure bulk fluid uptake 2mg/ml FITC-dextran (70kDa; Sigma) was added to cells at 5 x 10^6 /ml in shaking culture. At each timepoint, 500µl of cells were 7 8 removed and added to 1ml ice-cold KK2 (16.5mM KH₂PO₄, 3.8mM K₂HPO₄, pH6.1). Cells were then pelleted at 7,000 x *g* for 30 seconds, washed once in KK2 9 10 and frozen. Pellet were then lysed in 200µl 50mM Na₂PO₄ pH9.3, 0.2% Triton 11 X100) and fluorescence measured on a plate reader at 485 excitation/520nm 12 emission. Fluorescence was then normalised to total protein in an additional 13 sample and calculated as a percentage of wild-type cells at 90 minutes. 14 15 Macropinosome volume was measured by incubating cells for 5 minutes in 16 0.1µg/ml FITC-dextran and obtaining Z-stacks on a spinning disc confocal

17 microscope. FITC is pH-sensitive and the sensitivity was set so only new non-

18 acidified macropinosomes were visible. For analysis, individual cells were

19 cropped out, randomised, and volume calculated from manually measuring the

20 maximum diameter of each macropinosome in each cell, assuming they were21 spherical.

22

23 Phagocytosis assays

24 Phagocytosis of fluorescent beads was measured by flow cytometry as

25 previously described in detail (Sattler et al., 2013). Briefly, 1 or 4.5 μm diameter

26 YG-carboxylated polystyrene beads (Polysciences Inc) were shaken with 2 x 10⁶

27 *Dictyostelium* /ml at ratios of 200:1 and 10:1 respectively. 500 μl samples were

28 removed at each timepoint and added to 3 ml ice-cold Sorenson sorbitol buffer

29 (SSB; 15 mM KH₂PO₄, 2 mM Na₂HPO₄, 120 mM Sorbitol) containing 5mM sodium

30 azide. Samples were then centrifuged at $100 \ge g$ for 10 minutes, pellets

31 resuspended in SSB and analysed on an Attune NxT flow cytometer (Life

32 Technologies). Analysis was performed using FloJo software as described

33 (Sattler et al., 2013).

1 2 To measure uptake of GFP-expressing *M. smegmatis* by flow cytometry, the 3 bacteria were grown to an OD_{600} of 1, pelleted by centrifugation at 10,625 x g for 4 4 minutes and resuspended in 1ml HL5 medium. Bacteria clumps were then 5 disrupted by passing through a 26-guage needle several times, before adding a 6 1/10th volume of bacteria to a *Dictyostelium* culture and processing as above. 7 8 To measure phagocytosis of bacteria by decreasing turbidity, an overnight 9 bacterial culture in LB was diluted 1:25 and grown at 37°C until an OD₆₀₀ of 0.7 10 before pelleting and resuspension in SSB at an OD_{600} of 0.8. This was then added 11 to an equal volume of *Dictyostelium* at 2×10^7 cells/ml in SSB at room 12 temperature and shaken in flasks. The OD_{600} was then measured over time. 13 14 Phagocytosis and TRITC labeling of heat killed S. cerevisiae was performed 15 essentially as previously described (Rivero and Maniak, 2006). Dictyostelium at 16 1 x 10⁶ cells/ml in HL5 were seeded in glass-bottomed microscopy dishes (Mat-17 tek) and left for 1 hour prior to addition of a 5-fold excess of yeast. After 30 18 minutes, the fluorescence of extracellular yeast was guenched by addition of 0.2 19 mg/ml trypan blue and images of multiple fields of view taken on a wide-field 20 microscope scoring >100 cells per condition. 21 22 **Microscopy and image analysis** 23 Live cell imaging was performed in glass-bottomed microscopy dishes (Mat-Tek) 24 with cells seeded the preceding day in filtered HL-5 medium, unless otherwise 25 stated. Spinning disc images were captured using a Perkin-Elmer Ultraview VoX 26 spinning disk microscope with a UplanSApo 60x oil immersion objective (NA 1.4) 27 and Hammamatsu C9100-50-EM0CCD camera. Laser scanning confocal images 28 were obtained using a Zeiss LSM880 Airyscan confocal equipped with a Fastscan 29 detector, and a 63x 1.4 NA objective. Images were acquired in fastscan mode and 30 deconvolved by airyprocessessing using Zen black software (Zeiss). 31 32 Image analysis was performed using ImageJ (https://imageJ.nih.gov) with 33 average plots of protein enrichment across cups generated using a custom script

1 in Igor Pro (Wavemetrics). For this, confocal images were captured and linescans 2 of GFP-fluorescence intensity measured from the protrusive tip to tip. The 3 average signal from a 1-2 μ m non-protruding region of the cell was also 4 measured as was the local background outside each cell. Local background was 5 subtracted and signal across the cup divided by the non-protruding membrane 6 signal to give fold-enrichment. To compare enrichment across multiple cups, 7 normalised linescans were extrapolated over 1000 points and averaged. 8 Enrichment at the cup tip was measured by the average of the first 100 points of 9 the profile for each cup. 10 The bacterial long axis was measured automatically from widefield images of 11 12 either GFP or RFP expressing bacteria in ImageJ. Individual bacteria were 13 identified by thresholding and long axis measured using the Feret's diameter 14 function. 15 16 Western blotting and Lipid overlay assays 17 Western blotting was performed by standard techniques, separating proteins by 18 SDS-PAGE and probing using a custom rabbit polyclonal antibody to GFP (gift 19 from Andrew Peden). Endogenously biotinylated proteins were used as a loading 20 control, using Alexa680-labelled streptavidin (Life Technologies)(Davidson et al., 21 2013). Blots were imaged LiCor odyssey SA fluorescence gel imager. 22 23 For lipid overlay assays, 1 x 10⁷ Ax2 cells expressing the BAR domain fused to 24 GFP (pCB114) were washed once in SSB, lysed in 600 µl RIPA buffer (50mM 25 TrisHCl pH7.5, 150mM NaCl, 0.1% SDS, 2mM EDTA, 0.5% sodium deoxycholate, 26 1 x HALT protease inhibitors (Thermo Fisher), 0.5% Triton X100) and left on ice 27 for 45 minutes. Insoluble material was then removed by centrifugation at 15,871 28 x g for 20 minutes at 4°C. PIP strips or Arrays (Echelon Biosciences) were 29 blocked in 3% fatty acid-free bovine serum albumin (BSA) in TBS-T (20mM Tris 30 base, 150mM NaCl, 0.05% Tween20, pH7.2). Samples were then diluted in TBS-T 31 and incubated with the strips for 1 hour at 22°C, before washing and processing 32 as for Western blotting. 33

1 GAP and GEF biochemistry

2 Interactions with recombinant GST-Rac isoforms were performed as described 3 previously (Plak et al., 2013). Dictyostelium cells expressing GST-Rac bait 4 proteins and GFP-fused to the GEF domain of RGBARG were expressed and lysed 5 in 2mls of buffer (10mM Na2HPO4 pH7.2, 1% Triton X100, 10% glycerol, 150mM 6 NaCl, 10mM MgCl₂, 1mM EDTA, 1mM Na₃VO₄, 5mM NaF) including protease 7 inhibitor cocktail (Roche). Lysates were mixed with glutathione sepharose beads 8 (GE Healthcare) and incubated overnight at 4°C. Unbound proteins were washed 9 away with PBS, and bound proteins detected by Western blot using an anti-GFP 10 antibody (SC9996). 11 12 For GAP activity measurements, His-NF1 GAP domain (AA 2530-3158) and MBP-

13 His-RGBAR GAP domain (AA 1717-2045) were produced and isolated from *E.*

14 coli Rosetta cells. His-NF1 GAP was purified using a HisTrap excel - affinity

15 column (GE Healthcare) and eluted in buffer containing 50mM Tris, 50mM NaCl,

16 5% Glycerol, 3mM β -mercaptoethanol and 200mM imidazole, pH7.5. MBP-His-

17 RGBAR GAP, was purified by Maltose Binding Protein Trap (MBPTrap) - affinity

18 column (GE Healthcare) and eluted in 20mM Tris, 200mM NaCl, 5% Glycerol

19 1mM β-Mercaptoethanol and 10mM Maltose, pH7.5. Proteins were further

20 purified by size exclusion chromatography (Sephacryl 16/60, GE Healthcare) and

stored in 50mM Tris, 50mM NaCl, 5mM DTT, and 5mM MgCl2, pH7,5.1.

22

 $23~~1\mu M$ of the indicated Ras proteins with and without equal amount of indicated

24~ GAP domain was incubated with 50 μM of GTP at 20°C in 50 mM Tris pH 7.5, 50 $\,$

25 mM NaCl and 5mM MgCl₂. At each timepoint the GDP content of the samples was

26 analysed by a HPLC (Thermo Ultimate 3000): a reversed phase C18 column was

27 employed to detect GDP and GTP content (in %) as previously described (Eberth

and Ahmadian, 2009). Linear rates of GDP production (first 4-8 timepoints) were

29 calculated using GraFit 5.0 (Erithacus software).

30

31 Ellipsoid beads generation and phagocytosis

32 $3\mu m$ unmodified non-fluorescent polystyrene beads (Polysciences Inc.) were

33 embedded in polyvinyl alcohol (PVA) film (Sigma Aldrich) and stretched as

1 previously described (Ho et al., 1993). Briefly, 2.8 mls beads of bead solution 2 were added to 20mls 25% w/w dissolved PVA solution and poured into a 10.5 x 3 10.5 cm plastic mold to create a film. These were cut into 3 x 2cm strips, marked 4 with a grid to follow deformation and placed in a custom stretching device as 5 described in detail in (Ho et al., 1993). Films were the placed in a 145 °C oil bath 6 to soften beads and film and slowly pulled to the desired length. After cooling below the glass transition Tg temperature, the beads were extracted from the 7 central region where the grid was deformed evenly. This part was cut into small 8 9 pieces and rotated in 10 mls of a 3:7 mix of isopropanol:water overnight at 20°C 10 to dissolve. Beads were aliquoted and twice heated at 75°C for 10 minutes and 11 washed in isopropanol: water. Beads were then washed twice in 12 isopropanol:water at 22 °C, before two washes in water. The amount of stretch 13 was measured by imaging on an inverted microscope and manually measuring 14 their length in Image]. 15 16 To measure phagocytosis, equal numbers of stretched and unstretched beads 17 were mixed, sonicated and incubated at a 10-fold excess to cells at 1×10^6 /ml, 18 shaking in HL5. After 30 minutes, 500 µl samples were added to 3 ml SSB with

19 5mM sodium azide to detach unengulfed beads. Cells were washed in ice-cold

20 SSB, transferred to a microscopy dish and allowed to adhere for 10 minutes

21 before imaging and the number of each shape bead internalised quantified

22 manually.

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Supplementary videos 1 2 3 Video 1: RGBARG-GFP is enriched at the tips of macropinocytic cups. Movie of 4 *RGBARG*- cells expressing RGBarG-GFP in axenic culture. 5 6 Video 2: RGBARG-GFP is enriched at the tips of phagocytic cups. *RGBARG*- cells 7 expressing RGBarG-GFP engulfing a TRITC-labelled budded yeast. 8 9 Video 3: RGBARG-GFP localises to the periphery of PIP3 patches during macropinocytosis. 3D movie of cells macropinosome formation in *RGBARG*- cells 10 11 expressing RGBARG-GFP and PH_{CRAC}-RFP. 12 **Video 4:** PIP₃ dynamics and cup formation in Ax2 cells. Maximum intensity 13 14 projection of Ax2 cells expressing PH_{CRAC}-GFP, showing removal of PIP3 15 signalling after macropinosome formation is complete. 16 17 Video 5: RGBARG- cells have larger and more persistent PIP3 patches. Maximum 18 intensity projection of cells expressing PH_{CRAC}-GFP. 19 20 Video 6: Phagocytosis of TRITC-labelled yeast by Ax2 cells expressing PH_{CRAC}-21 GFP. 22 23 Video 7: Phagocytosis of TRITC-labelled yeast by RGBARG- cells expressing 24 PH_{CRAC}-GFP.

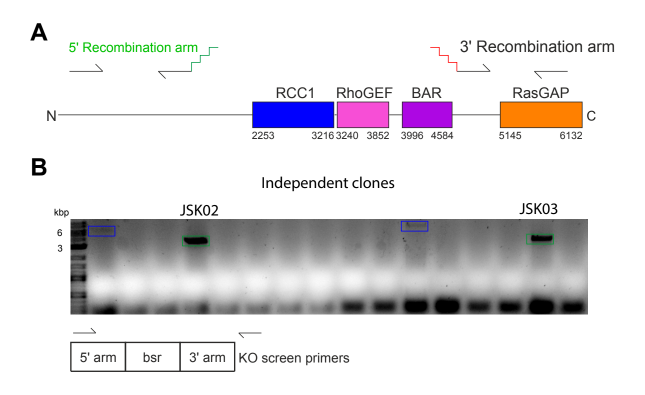


Figure 2 supplement: Disruption of DDB_G0269934. (A) Schematic of the genomic locus indicating the position of the regions encoding each domain, and the 5' and 3' recombination arms amplified by PCR. These were attached either side of a blasticidin selection cassette by fusion PCR and used to transform Ax2 cells and delete ~3 kbp of the gene. (D) PCR screen of transformants, using one primer within the 5'recombination arm and another after the 3' arm. Clones with DDB_G0269934 disrupted will give a product of 3.1 kbp (green box), the wild-type locus is 6.1 kbp (blue boxes).

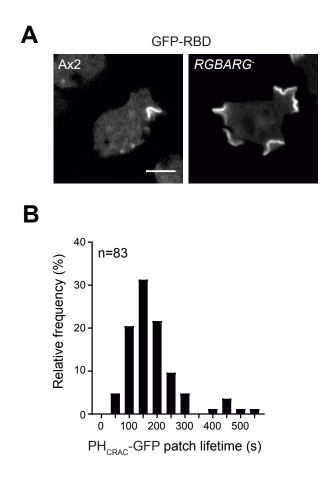


Figure 3 supplement: (A) Distribution of the active Ras probe RBD-GFP in wild-type and RGBARG- cells. Images are single confocal planes, bar indicates 5 μ m. (B) Histogram of PHCRAC-GFP patch lifetime in Ax2 cells from maximum projection movies. Lifetime was measured from the first frame an independent patch was visible to when it was completely removed from both the surface and any internalized vesicle.

Α								
RGBARG(225kD)		754	RCC1		RhoGEF	BAR -		GAP 2044
RGBARG∆RCC1(189kE))	751		<mark>.</mark>	RhoGEF	BAR -	-	GAP 1722
RGBARG∆GEF(201kD)		751	RCC1)72		BAR -		GAP 1834
RGBARG∆BAR(202kD)			RCC1	- F	RhoGEF		Ras	GAP
RGBARG∆GAP(188kD)			RCC1	- F	080 1284	BAR	1517	1846
RGBARG _{R1792K} (225kD)		751	RCC1	R	080 1284	1332 1528 BAR 13321529	R1792K	3AP 2044
RCC1(35kDa)		-	RCC1	321				
RhoGEF(22kDa)				<mark>F</mark>	RhoGEF			
BAR(22kDa)						BAR		
RasGAP(36kDa)						1 197	······ <mark>Ras</mark> i 1	GAP 329

Figure 4 Supplement 1: Schematic of the RGBARG truncation and point mutants used in this study.

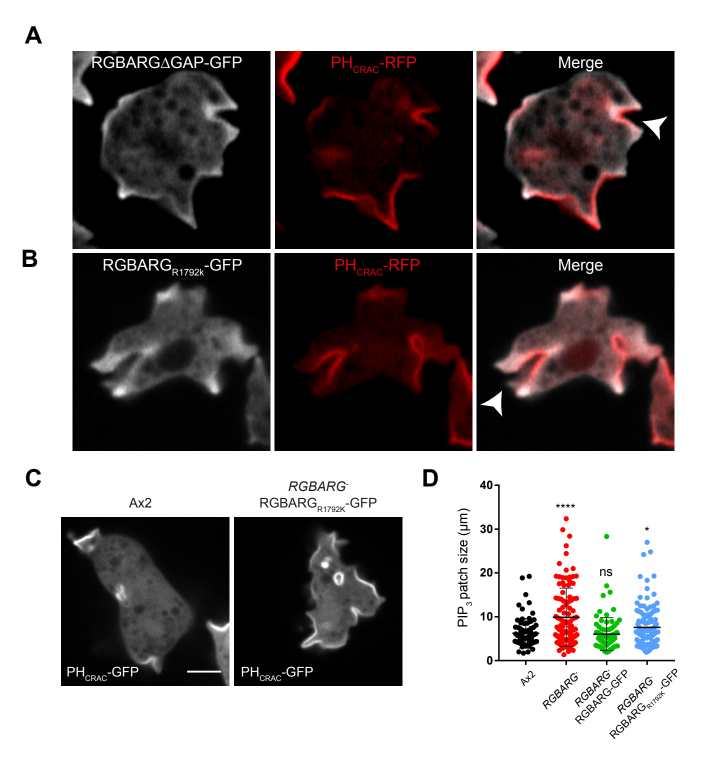


Figure 4 Supplement 2: (A) Co-expression of RGBARGdRasGAP-GFP and PHCRAC-RFP in RGBARG- cells indicating that the RasGAP domain helps excluded RGBARG-GFP from PIP3 rich regions of the cell. (B) The GAP activity inactivating point mutant R1792K localizes normally and is excluded from the base of protruding cups (arrowhead). (C) PHCRAC-GFP localization in Ax2 and RGBARG- cells expressing RGBARGR1792K-GFP, demonstrating that PIP3 dynamics are not rescued by this construct. PHCRAC-GFP patch size is quantified in (D).

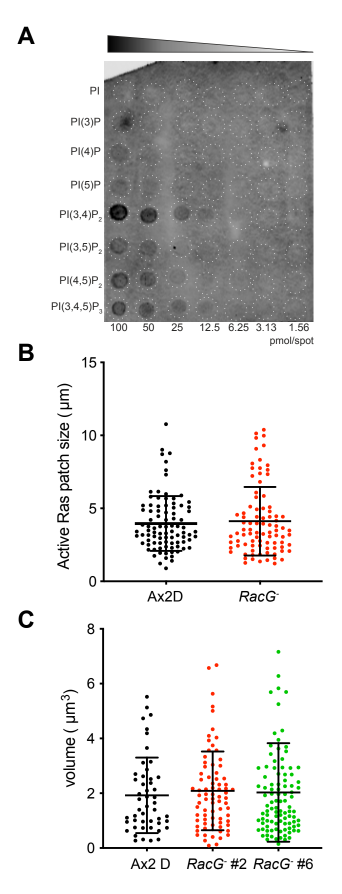


Figure 5 Supplement: (A) PIP array analysis of BAR-GFP binding showing a moderate preference for PI(3,4)P2 in this assay. (B) Active Ras signaling during macropinosome formation in RacG- cells and their parental cell line Ax2D. Patch size was quantified from single confocal planes of cells expressing RBD-GFP. (C) Volume of macropinosomes formed by RacG- and control cells. Measured by imaging cells taking up FITC dextran. Two independent RacG- clones were analysed over 3 independent experiments.