Cell Cycle Heterogeneity Can Generate Robust Cell Type Proportioning

Graphical Abstract

Cell cycle position is heterogeneous within a population of growing cells

Cell cycle position affects responsiveness to differentiation cues

Cell cycle perturbation is buffered by changes in responsiveness to differentiation cues

Highlights

- Dictyostelium cells break symmetry in a stochastic salt and pepper fashion
- Cell cycle position affects responsiveness to differentiation inducing signals
- Cell cycle length variation ensures cells are distributed in different cycle phases
- Perturbation of cell cycle dynamics is buffered by changes in signal responsiveness

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In Brief
Gruenheit et al. find cell cycle position heterogeneity promotes symmetry breaking during development as it helps cells respond to different differentiation cues. Buffering by changes in signal responses ensures this system is robust to perturbation of cell cycle dynamics. This illustrates how heterogeneity can be optimized to facilitate development.
Cell Cycle Heterogeneity Can Generate Robust Cell Type Proportioning

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SUMMARY

Cell-cell heterogeneity can facilitate lineage choice during embryonic development because it primes cells to respond to differentiation cues. However, remarkably little is known about the origin of heterogeneity or whether intrinsic and extrinsic variation can be controlled to generate reproducible cell type proportioning as seen in vivo. Here, we use experimentation and modeling in D. discoideum to demonstrate that population-level cell cycle heterogeneity can be optimized to generate robust cell fate proportioning. First, cell cycle position is quantitatively linked to responsiveness to differentiation-inducing signals. Second, intrinsic variation in cell cycle length ensures cells are randomly distributed throughout the cell cycle at the onset of multicellular development. Finally, extrinsic perturbation of optimal cell cycle heterogeneity is buffered by compensatory changes in global signal responsiveness. These studies thus illustrate key regulatory principles underlying cell-cell heterogeneity optimization and the generation of robust and reproducible fate choice in development.

INTRODUCTION

Tissue patterning and cell type proportioning are robust and reproducible. However, cell-cell heterogeneity can prevent cell populations from behaving in a coordinated fashion. For example, many developmental systems are regulative and can correct errors (Lawrence and Levine, 2006). Gene networks underlying cell fate choice can also buffer fluctuations (Averbukh et al., 2017; Levy and Siegal, 2008). Recent studies, however, suggest that heterogeneity can increase the spectrum of differentiation capabilities of cells in a uniform environment (Altschuler and Wu, 2010; Balázs et al., 2011). Most notably, this leads to random “salt-and-pepper” differentiation where reproducible proportions of different cell types still arise. Examples range from competence in B. subtilis (Maamar et al., 2007) to lineage specification in the mouse blastocyst (Dietrich and Hiiragi, 2007).

Although the molecular mechanisms underlying salt-and-pepper differentiation are poorly understood, general principles are emerging. First, heterogeneity is thought to prime some cells to adopt a particular lineage (Canham et al., 2010; Chang et al., 2008). For example, priming could affect the likelihood that a cell will respond to signals that trigger differentiation, even if all cells receive the signals (i.e., it affects the threshold of responsiveness) (Canham et al., 2010; Chang et al., 2008). Alternatively, in cases where differentiation is cell-autonomous and achieved in the absence of an external cue, primed cells may simply express different amounts of key regulators of the differentiation program (Maamar et al., 2007). Second, the primed state is thought to be unstable and transient (Canham et al., 2010; Filipczyk et al., 2015; Suel et al., 2006). For example, when primed cells are isolated and regrown, the heterogeneous population is rapidly reconstituted (Canham et al., 2010; Chang et al., 2008). Despite this emerging framework, it is unclear how the expression of lineage priming genes affects the threshold of responsiveness or cell fate choice at the molecular level. Furthermore, because few lineage priming genes have been identified, it is unknown how lineage priming dynamics or the number of lineage-primed cells is controlled. Addressing these questions will be crucial to understanding how this mechanism can achieve robust cell type proportioning.

Stochastic lineage priming dynamics provide one method of achieving robust developmental outcomes (Schultz et al., 2007). This is because even though the behavior of one cell may be unpredictable, the probability of a proportion of cells within a population being in a primed state can be fixed. Alternatively, there is evidence that lineage priming dynamics can be governed by an underlying oscillatory mechanism that reproducibly drives cells in and out of a primed state (Soufi and Dalton, 2016). For example, studies of human embryonic stem cell (hESC) differentiation have revealed a relationship between the cell cycle and lineage potential (Li and Kirschner,
Differentiation of hESCs is favored in the G1 phase of the cell cycle, with endoderm fate favored in early G1 and neuroectoderm fate in late G1. However, induction of neuroectoderm and endoderm occurs in response to positional signals during gastrulation, rather than in a salt-and-pepper distribution (Rossant and Tam, 2009). Consequently, how these cell culture observations relate to differentiation in vivo is unknown. Despite these limitations, these studies demonstrate that cell cycle position allows an asynchronous population of cells to exhibit differential responses to a uniform stimulus and thus provides an attractive candidate mechanism for robust salt-and-pepper differentiation in vivo.

To address this possibility, we have utilized the social amoeba Dictyostelium discoideum. Individual Dictyostelium amoebae grow and divide when nutrients are present. However, when nutrient supplies are exhausted, approximately 10^5 cells aggregate to form a multicellular mound, which undergoes a program of cell type differentiation and morphogenesis. It is thought that although growing cells are undifferentiated, they are dynamically and unstably primed toward future developmental fates (Chattwood et al., 2013). Differences in cell cycle position (Araki et al., 1994; Gomer and Firtel, 1987; Thompson and Kay, 2000a; Weijer et al., 1984), intracellular calcium concentration (Kubohara et al., 2007; Schaap et al., 1996), intracellular pH (Gross et al., 1983; Kubohara et al., 2007), and nutritional history (Leach et al., 1973; Thompson and Kay, 2000a) of growing cells have all been shown to bias cell fate choice and vary dynamically within populations of cells. Furthermore, it has been shown that these biases modulate the threshold of responsiveness to diffusible developmental signals, such as cAMP and DIF-1 (Chattwood et al., 2013; Thompson and Kay, 2000a), which only accumulate to high levels following starvation and aggregation (Kay and Thompson, 2001). Cell type differentiation thus takes place at the mound stage. DIF-1 promotes the differentiation of prestalk populations including ecmA-expressing pstAO cells and ecmB-expressing pstB cells (Keller and Thompson, 2008; Thompson and Kay, 2000b) (Figure S1). cAMP promotes the differentiation of prespore cells, which express the pspA gene. As all cells within the mound are uniformly exposed to these signals, it has been proposed that intrinsic differences in thresholds of responsiveness result in salt-and-pepper differentiation (Chattwood et al., 2013; Thompson et al., 2004b). Crucially, multicellular development in Dictyostelium is extremely robust, with cell type proportioning reproducible between multicellular aggregates and over a wide range of total cell numbers (Bonner, 1957; Ráfols et al., 2001). One possibility is that initial symmetry breaking is noisy but stabilized by feedback loops in which DIF-1 is synthesized by prespore cells and degraded by prestalk cells to ensure DIF-1 levels reach an equilibrium and maintain prestalk-prespore proportioning (Kay and Thompson, 2001). Alternatively, initial symmetry could be precise and result in near correct proportioning that only requires correction if the system is severely perturbed. We, therefore, reasoned that Dictyostelium was an ideal system to address whether variation in cell cycle position can provide sufficiently robust population-level information to generate reproducible cell type proportioning in vivo.

RESULTS

Single-Cell RNA-Seq Reveals Cell Cycle Position as the Major Driver of Cell-Cell Gene Expression Variation

In Dictyostelium, a clonal cell population can break symmetry and undergo robust cell fate choice and proportioning. We, therefore, reasoned that the major drivers of lineage priming should be detectable as variation in gene expression between growing cells. Moreover, we would expect the proportion of cells exhibiting distinct gene expression states to reflect the cell type proportions seen in development (the approximate ratio of prespore to prestalk cells is 70:30). To investigate this, we took an unbiased approach in which single-cell RNA sequencing (RNA-seq) was carried out on 81 log-phase vegetative cells to identify genes with highly variable expression patterns. Out of 11,319 expressed genes, we identified 1,619 significantly variable genes (p value = 0.01) by fitting a Michaelis-Menten model (K = 5.94). These differentially expressed genes (DEGs) revealed the presence of two large groups of cells, cluster A (25 cells) and cluster B (56 cells). For each gene, we next determined whether it could be used as a marker gene for cluster A or cluster B (or a subgroup thereof). Of the 1,619 DEGs, 1,602 genes had an AUC higher than 0.8, indicating that they are specific for either cluster A (901 genes; Figure 1A), or cluster B (701 genes; Figure 1A). 493 of these genes also show variation within cluster B, but expression within that cluster is still higher than the expression in cluster A. Because almost all significantly variable genes are also marker genes for these two clusters, this shows these genes are the main difference between the two groups. Finally, a principal-component analysis (PCA) using the 500 most variable genes also divided the cells into the same two groups (Figure 1B), and the percentage of cells in each group (31% and 69%) roughly corresponded to the ratio of prestalk and prespore cells seen during development.

To understand the molecular basis of this gene expression variation, we first performed Gene Ontology (GO) analyses. Cluster A genes showed enrichment for GO terms including “myosin II filament organization,” “mitotic cytokinesis,” and “cytokinesis” (Data S1). Marker genes for cluster B were enriched for GO terms including “oxidative phosphorylation,” “translation,” and various biosynthetic processes (Data S1). This raised the possibility that differences in cell-cycle-dependent gene expression could underlie the clusters, with cells in cluster A preparing to divide, undergoing, or having just undergone mitosis (M/S phase) and cells in cluster B actively growing (G2 phase). To test this, we used cells synchronized by cold shock to determine the Dictyostelium cell-cycle-dependent transcriptome. Cold shock likely arrests cells in late G2, because most cells undergo mitosis within 2–3 hr after cold shock release (Maeda, 1986; Strasser et al., 2012). BrdU incorporation reveals these cells then directly enter S phase (there is little or no G1 in Dictyostelium). Indeed, genes associated with M/S phase are up-regulated shortly after cold shock release (Strasser et al., 2012). Cells then undergo G2, with the second round of less synchronous mitosis typically observed around 8 hr after cold shock release. We used gene expression data from cells collected at hourly time points after cold shock release to capture one complete cell cycle. After filtering to remove poorly expressed genes, the remaining 6,121 genes were screened for significant expression peaks at different
Figure 1. *Dictyostelium* Cells Exhibit Cell Cycle Heterogeneity

(A) Gene expression heatmap for 1,602 marker genes reveals two clusters of vegetative cells. DEG analysis split the 81 vegetative cells into two groups. Expression values for each gene were mean-centered and scaled.

(B) Principal-component analysis of 81 vegetative cells. The 500 most variable genes were used for PCA. Colors correspond to highlighted cells in (A). Normal data ellipse for each cluster is shown.

(C) Cell cycle profiles of 6,228 genes. Cell cycle profiles from cold shock synchronized cells for 6,228 genes were extracted and normalized to percent expression (see STAR Methods). Genes that deviate more than 1.5 median absolute deviations (MADs) were binned according to the time point of their highest expression. Profiles for each bin were separated (a)–(h). The blue line denotes the mean.

(D) Cell cycle profile of marker genes for cluster A and cluster B. A hypergeometric test was used to identify genes from cluster A (901 genes) and cluster B (701 genes) that were over-represented in the hourly cell cycle bins described in Figure 1C. Cluster A and cluster B genes show higher expression 5–7 and 3 hr after cold shock release, respectively.

(E and F) Cell cycle length correlations. Cell cycle lengths of sister (E) or mother and daughter cells (F) were plotted. Blue line depicts the linear regression line (gray shaded area: 95% CI).

(G) Cell divisions become rapidly de-synchronized in cell populations. Starting with one cell, 1,000 population simulations were run for 96 hr, and the number of cells dividing in each hour plotted (solid line: mean, dashed lines: 95% CI). The mean converges against a value of 0.11, which is indicative of de-synchronous cell cycle lengths.
time points (see STAR Methods) and binned accordingly (Figure 1C and Data S2). GO analyses revealed strong cell cycle signatures, e.g., a significant over-representation of the terms “DNA replication,” “chromosome segregation,” and “nuclear division” 1 hr and 5–7 hr after cold shock release, which supports the idea that the majority of these cells are indeed in M/S phase or preparing to undergo mitosis (Data S2). Most importantly, using a hypergeometric test to identify over-representation, we found that cluster A genes were significantly enriched for genes with peaks in late G2 (p < 0.05; 5–7 hr after cold shock release) and then fall sharply after S phase (8 hr) (Figure 1D). It is also interesting to note that only a small number of cluster A genes have increased transcription 1 hr after cold shock release, which likely reflects the fact that cold shock specific perturbations affect the first M/S phase these cells undergo. The second M/S phase after cold shock therefore likely better reflects a normal cell cycle. Markers of cluster B, on the other hand, were significantly enriched for genes that peak 3 hr after cold shock release (p = 3.24 × 10^{-31}; Figure 1D). For example, most ribosomal proteins exhibited this transcription pattern, which is indicative of cell growth. Consequently, our single cell analyses reveal that gene expression variation between individual Dictyostelium cells is likely a consequence of a strong cell cycle signature.

Stochastic Cell Cycle Variation Results in Robust and Rapid De-synchronization of Dictyostelium Cell Populations

If cell cycle position underlies cell fate decisions in Dictyostelium, for proportioning to be reproducible, cells must be predictably distributed (on a population scale) throughout the cell cycle when differentiation is induced. Consequently, the Dictyostelium cell cycle must exhibit sufficient stochastic cell-cell variation to drive rapid population asynchrony and robust convergence to a steady-state distribution of cells in different positions in the cell cycle. To test this, we used time-lapse microscopy to measure cell cycle variation. This revealed sister cell divisions are correlated yet still somewhat variable (Figure 1E). However, no correlation was observed between mother and daughter cell cycle length (Figure 1F). Using these data, we next built a mathematical simulation to determine the number of cell divisions that would be required to completely de-synchronize a population of Dictyostelium cells that arises from a single cell (see STAR Methods). This revealed de-synchronization would occur within 40 hr, or five cell divisions (i.e., 32 cells) (Figure 1G). A single fruiting body is typically composed of more than 10,000 cells, and fruiting bodies of less than 1,000 cells can only be generated under very specific laboratory conditions (Konijn and Raper, 1961). This suggests synchrony would be lost long before cell differentiation. Therefore, if cell cycle position determined cell fate, the decision made by each cell would be unpredictable because it is based on a stochastic process. However, the average behavior of the population, due to the fast convergence to a steady-state cell cycle position distribution, would result in precise cell fate proportions.

Cell Cycle Position Results in Oscillatory Cell Fate Choice Probabilities

To address whether cell cycle position is quantitatively linked to cell fate, we developed a high-throughput live imaging method to track cells during vegetative growth and differentiation (Figure 2A). Differentiation was induced by starvation for 18 hr in conditioned medium containing a cocktail of differentiation-inducing signals. To validate this method, fluorescence-activated cell sorting (FACS) quantification was used to determine the number of cells expressing markers of the major prespore (pspA-GFP) and prestalk (ecmAO-RFP or ecmB-RFP) cell types. As expected, the markers showed a dose-dependent response to the prestalk inducer DIF-1 (Figure 2B). Furthermore, at a dose of 10 nM DIF-1, which is thought to be close to physiological levels, the number of prestalk and prespore cells was similar to those seen in vivo (16% ecmB, 22% ecmAO, and 62% pspA expressing) (Figure 2C). For most cells, it was possible to unambiguously assign cell fate because less than 1% of the cells co-expressed two markers (Figure 2C). Finally, most cells differentiated, and less than 10% of the cells expressed no marker (Figure 2C).

To determine whether there is a relationship between cell cycle stage and prestalk or prespore fate choice, growing cells were filmed at low density for 12–14 hr. Under these conditions, cells have an average cell cycle length of 8.3 hr, and all cells can be tracked through at least one cell division (Video S1). Furthermore, analysis of PCNA-RFP expressing cells allowed us to confirm that the timing of the last cell division can be used to predict cell cycle position accurately. Almost all cells that underwent mitosis accumulated a PCNA-RFP dot in the nucleus within 20 min (76% within 20 min, 98% within 30 min; n = 93 cells) (Video S2). BrdU incorporation has previously revealed this coincides with late S phase (Muramoto and Chubb, 2008). This confirms previous observations that Dictyostelium cells have little or no G1 and that cells immediately enter S phase after mitosis, before cells enter G2. Once the growth medium was removed, and cell type differentiation induced, little further cell division was observed (6% of cells). Consequently, the timing of the last division relative to the addition of signals was used to define the cell cycle position of each cell when induced to differentiate. Finally, after 18 hr, a fluorescence image was taken to determine whether each cell had adopted a prespore (pspA-GFP) or prestalk (no fluorescence) fate (Video S1).

Cell type differentiation was found to follow an oscillatory pattern. Prespore cell differentiation peaked in cells that underwent mitosis 6–7 hr before removal of growth medium (i.e., induced to differentiate in mid G2) (Figure 2D). The probability of prestalk cell differentiation was highest when cells had just undergone mitosis (i.e., in S phase). Prestalk or prespore cell differentiation was not restricted to specific cell cycle phases. For example, significant numbers of prestalk cells differentiated when cells divided 2–3 hr before the addition of differentiation signals. These cells cannot be in S phase because PCNA-RFP localization measurements reveal S phase length to be 37 min on average (μ = 37.4 min, σ = 5.1 min) and a maximum of 50 min (Figure 3A). Finally, cells at the end of G2 were also slightly stalkier than cells in the rest of G2 (Figure 2D). This suggested that the probability of prestalk cell differentiation increases in G2 but only just before cells enter mitosis. Indeed, cells with a significantly longer than average cell cycle length, which are more likely to divide within the next hour, are relatively stalky. Together, these findings suggest that the cell cycle acts as a probabilistic cell fate oscillator that regulates the threshold of responsiveness to differentiation signals.
Nutritional History Affects Cell Cycle Length and Cell Fate Choice

Our findings suggest that the Dictyostelium cell cycle is an example of a dynamic system that relies on stochasticity (in cell cycle length) to ensure robust cell type proportioning. One problem, however, is that extrinsic disruption of population heterogeneity, oscillator period, or amplitude would have a deleterious effect on development. Indeed, extrinsic environmental perturbations, such as pH, temperature, or metabolic changes, affect cell fate choice. However, it is unknown whether these effects are due to changes in cell cycle dynamics or other changes in cell physiology.

To test this, we further studied the behavior of cells grown in media containing low glucose (G\textsubscript{+/C0}), which are biased toward the prestalk cell fate (Leach et al., 1973; Thompson and Kay, 2000a; Figure 4A). We found G\textsubscript{+/C0} cells have a significantly longer cell cycle than M/S G\textsubscript{2} longer than average cell cycle (t test; \(p = 6.18 \times 10^{-6}\); Figure 3A). However, not all cell cycle phases increase in length. PCNA-RFP localization revealed no difference in M/S phase length (t test; \(p = 0.56\); Figure 4A), whereas G2 was significantly lengthened (t test; \(p = 6.25 \times 10^{-6}\); Figure 3A). This finding initially appeared counterintuitive because mid-G2 cells tend to differentiate as prespore cells. However, our single cell analyses did reveal that the small number of G+ cells with a significantly longer than average cell cycle are biased toward the prestalk cell fate (Figure 2D). This raised the possibility that G\textsubscript{-} grown cells (and G+ cells with a longer than average cell cycle length) may be similar to M/S phase cells. To test this, we used RNA-seq to compare the gene expression profile of G+ and G\textsubscript{-} grown cells. This revealed 324 DEGs (Data S3). As expected, G\textsubscript{-} cells showed high expression of genes associated with gluconeogenesis and the glutamate metabolic process (e.g., fbp, pckA), rather than glycolysis. However, G\textsubscript{-} cells also exhibited changes in cell-cycle-dependent gene expression (e.g., cdc20, aurK, cdc45, top2). Most importantly, genes with peaks associated with M/S phase following cold shock release are over-represented, with a corresponding decrease in G2-associated gene expression (Figure 3B). This suggested that metabolic status affects cell cycle dynamics, with G\textsubscript{-} cells stalled in late G2 phase when M/S-associated transcription increases, thus resulting in a prestalk fate bias (Figure 3C). To confirm this, we used the single cell growth and differentiation assay to determine the lineage potential of G\textsubscript{-} grown cells at different cell cycle phases. This revealed G\textsubscript{-} grown cells with a longer than average G2 phase do exhibit a prestalk fate bias.

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Figure 3D, thus supporting the idea that the coupling of cell cycle phase to differentiation propensity underlies cell type proportioning in Dictyostelium. However, they also illustrate that cells are susceptible to extrinsic environmental perturbation of these cell cycle dynamics.

Ras Activity Rescues the Stalk Bias Caused by G– Growth without Affecting Cell Cycle Dynamics

Developmental mechanisms should result in reproducible cell type proportioning (within normal parameter space). We, therefore, hypothesized that robustness in Dictyostelium development would be facilitated if extrinsic environmental fluctuations that perturb cell cycle dynamics could be buffered. Increases or decreases in buffering activity would therefore be expected to rescue or exacerbate the effects of growth in the absence of glucose, but without altering cell cycle dynamics. Previous studies suggested that the putative Ras-GEF gefE might act in this way. For example, gefE knockout results in a sporey bias in G+ cells, and can rescue the stalk bias caused by G– growth (Figure 4; Chattwood et al., 2013), but without affecting growth.
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Consequently, regulation of the levels of Ras activity provided a
standout candidate for a buffering system.

The Dictyostelium genome encodes 11 putative Ras homo-
logs, but only three of these (rasC, rasD, and rasG) are expressed
at appreciable levels during multicellular developmental stages,
when cell fate choice is made (Rosengarten et al., 2015).
To further explore the connection between Ras signaling, cell fate
choice, and buffering, we compared the effects of knocking out
gefE with other genetic manipulations that should result in
Ras pathway attenuation or hyper-activation. First, we gener-
ated rasC, rasD, and rasG gene knockouts in the AX3 genetic
background used in this study. We found that rasD− knockout
mutant cells phenocopied the gefE− mutant and exhibited a
spore bias in chimera or forced wild-type cells to adopt stalky
fates (Figure 4A). As previously reported, rasC− mutant cells
are blocked during aggregation when clonally developed (Lim
et al., 2001) or when in the majority in chimera. However, this
defect is non-cell-autonomous, and labeled rasC− mutant cells
efficiently enter chimeric development when mixed with a major-
ity of wild-type cells. This revealed that rasC− mutant cells, unlike
the rasD− mutant, do not exhibit a cell fate bias and are distrib-
uted evenly throughout chimeric slugs. Finally, we tested the ef-
fects of knocking out rasG. rasG− mutant slugs are smaller than
wild-type when developed clonally or in the majority. Most
importantly, however, rasG− mutant cells actually show the
opposite phenotype to the rasD mutant, as they exhibit a stalk
bias, forcing wild-type cells to adopt the prespore cell fate.
This is consistent with the fact that rasG− mutant cells express
higher levels of activated RasD (Khosla et al., 2000).
Alternatively, this may be because rasG− mutant cells, like G− cells,
exhibit a longer cell division time than wild-type cells (Tuxworth
et al., 1987). These findings thus support the idea that Ras family
members play different roles during Dictyostelium development
and that RasD activity is dependent on GefE. Indeed, gefE−
mutant cells exhibit decreased RasD activation, whereas RasC
or RasG are unaffected (Chattwood et al., 2013).

To confirm this idea, we also generated strains in which constit-
utively active G12T versions of RasC, RasD, or RasG were ex-
pressed under the control of a tetracycline-inducible promoter.
This allowed expression to be transiently induced during growth.
When labeled wild-type cells were mixed with cells transiently
induced to express activated Ras, expression of constitutively
active RasDG12T resulted in a strong stalky fate bias (i.e., oppo-
site to the rasD− knockout). We also found that conditional
expression of constitutively active RasG[G12T] or RasC[G12T]
tended to push cells toward the prespore fate. However, the effect
was much weaker than that seen for RasDG12T. These findings sup-
port earlier studies that showed Ras proteins can exhibit a de-
gree of functional overlap when overexpressed (Khosla et al.,
2000). However, when knockout and overexpression studies are
taken together, our results suggest that GefE acts through
RasD to promote prestalk cell fate choice. This idea is supported
by monolayer culture assays, where rasD− mutant cells formed
fewer stalk cells than wild-type, and transient activation of
RasDG12T increased stalk cell differentiation (Figure 4B).

Chimeric development and monolayer differentiation assays
suggest that GefE specifically controls RasD to ensure correct
fate choice. Therefore, we next tested whether these effects
were due to changes in the response threshold (which is consist-
tent with a buffering mechanism), rather than changes in cell cy-
cle dynamics. The doubling time of wild-type cells (7.4 hr) was
indistinguishable from gefE− (7.8 hr), rasD− (7.3 hr), or RasD[G12T]
(7.8 hr) cells (Figure 4C). Furthermore, we found that the lengths
of M/S and G2 phases were indistinguishable in gefE− mutant
and wild-type cells (Figure 4D; Student’s t test, total: p = 0.27;
M/S: p = 0.18; G2: p = 0.28), whether cells were grown in the
presence or absence of glucose (Figure 4D). Together, these re-
results suggest that nutritional history affects cell cycle phase
lengths and thus the amount of time cells spend in a signal sen-
sitive state. In contrast, the effects of altering RasD activity are
due to changes in the cellular response threshold to differenta-
tion-inducing signals. Indeed, we found that gefE− knockout af-
fects the responsiveness to prestalk inducing signals during all
phases of the cell cycle (Figures 4E and 4F).

Cell Cycle Dynamics and Ras-Dependent Signal
Threshold Are Negatively Coupled
Changes in cell cycle dynamics or RasD-dependent changes in
the response threshold to differentiation signals can affect cell
fate choice and proportioning. Therefore, we next tested
whether changes in cell cycle dynamics that increase the num-
ber of prestalk cells result in a compensatory decrease in
RasD activity (i.e., favoring prespore cell differentiation) to
attempt to rebalance cell fate proportioning. RNA-seq gene

Figure 4. RasD Activity Affects Cell Fate Choice but Not Cell Cycle Dynamics

(A) Ras activity and nutrition affect cell fate choice in chimeric development. Left panel: gefE− cells exhibit a prespore and tip bias (black arrows) in chimera with wild-type cells. G− cells predominantly sort to the collar and back in chimera with G+ cells. gefE− G− cells are also found in the spore region (red arrows), which results in partial rescue of the G− effect (Chattwood et al., 2013). Middle panel: labeled wild-type cells were mixed with unlabeled rasC−, rasD−, or rasG− cells. rasD− cells exhibit a spore bias (black arrows), forcing wild-type cells to adopt prespore fates (white arrows). rasG− cells exhibit a stalk bias. rasC− cells do not develop when in the majority. They co-aggregate with wild-type cells in the majority but do not show a cell fate preference. Right panel: labeled wild-type cells were mixed with unlabeled cells induced to express constitutively active rasC[G12T], rasD[G12T], or rasG[G12T]. RasD[G12T] expression resulted in the strongest effect, forcing cells toward the collar and back fate (i.e., opposite to the rasD− or gefE− mutant phenotype).

(B) RasD activity and nutrition affect stalk cell differentiation in monolayer cultures. Decreased Ras activity (rasD− or gefE−) significantly reduced stalk cell differentiation at 10-nM DIF-1 (3 replicates). In contrast, increased RasD activity (AX3♂gefE[G12T]) significantly increased the percentage of prestalk cells (t test: p = 0.0088; p = 0.0012; p = 0.0001).

(C) Cell cycle length is not affected by RasD activity. The growth rate of wild-type AX3, gefE−, rasD− mutant, and RasD[G12T] cells was measured in culture. Error bars are standard error of mean of 3 replicates.

(D) Cell cycle phase lengths are unaffected by RasD activity, PCNA localization was used to determine S and G2 length. There is no significant difference in S phase lengths, whereas total cell cycle and G2 phase lengths of G− grown cells in wild-type and gefE− are significantly longer.

(E and F) Stalk cell propensity decreases in gefE− G+ and gefE− G− cells. gefE− G+ cells (E) that divided up to 2 hr before induction have a higher propensity to become prespore cells. gefE− G− cell (F) propensity to become prespore cells is higher than AX3 G− cells at almost all time points or cell cycle phases.
Developmental time

relative expression rrgA:rigA

percent stalk cells

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expression profiling was first used to identify genes associated with high RasD activity (wild-type) or low RasD activity (gefE−). This revealed 45 DEGs (Figure 5A and Data S3). We next used these genes to determine whether Ras activity was increased or decreased when cells were grown in prestalk biased G− or prespore biased G+ conditions. We found a highly significant proportion of RasD-dependent genes (36 out of 45; hypergeometric test; p = 4 × 10−15) were also affected by nutritional history (>1.5 × increase or decrease in expression). Most importantly, when the level of expression of these genes was compared, prestalk-biased G−-grown cells were more similar to prespore-biased low RasD gefE− cells (Figure 5A). Therefore, these data suggest that when cells are subjected to growth in conditions that affect the cell cycle and make cells more likely to differentiate as prestalk cells, they also exhibit a gene expression profile that is associated with the prespore-biased low RasD gene state. This behavior is consistent with a cell fate buffering mechanism.

To further test this idea, we studied the behavior of representative Ras-dependent genes from the putative buffering network. rigA and rrgA were chosen because they exhibit representative (average) changes in response to gefE mutation and glucose (Figure 5A). Furthermore, their relatively high levels of vegetative expression in wild-type cells (Data S3) enabled us to confirm these data by different methods (see below). For example, we examined relative transcript abundance and thus gefE-dependent gene activity in single cells by RNA-fluorescence in situ hybridization (FISH). This revealed that the relative expression of both genes is similar in most cells (Figures 5B and 5C). However, a significant number of cells express one gene at higher levels. In wild-type, this tends to be rigA (Figures 5B and 5C). In RasD(G12T) cells, the number of cells expressing high levels of rigA increases further, whereas in gefE− and rasD− mutant cells the number of cells expressing high levels of rigA is increased (Figures 5B and 5C).

Although RNA-seq was performed on growing cells, we found that genes in the Ras-D-dependent gene network are maximally expressed during multicellular stages when cell fate choices are made (Figure 5D). Similarly, GO annotation reveals enrichment for plasma membrane and transmembrane proteins (p = 0.03), as well as secreted proteins (p = 0.01), which is consistent with a role in cell-cell signaling. Therefore, we next tested whether rrgA and rigA are required for normal cell fate choice. Knockout mutant lines were created in otherwise wild-type cells (i.e., “normal” RasD activity), cells in which the level of RasD activity was reduced through disruption of gefE− or elevated through expression of constitutively activated RasD(G12T). The behavior of these cells was tested in monolayer cell culture or in chimeric development with wild-type cells. Knockout of rigA in a gefE− background (which normally results in higher expression of rigA) is sufficient to rescue the prespore cell fate bias (tip and prespore in chimeric slugs) seen in low gefE− mutant cells (Figures 5E and 5F). Similarly, knockout of rrgA in an AX3<sup>rasD</sup>(G12T) background (which normally results in higher expression of rigA) is sufficient to rescue the prestalk cell fate bias (collar and back in chimeric slugs) caused by expression of constitutively active RasD<sup>G12T</sup> (Figures 5E and 5F). In both cases, no phenotype is seen in rrgA or rigA single mutants (Figure 5). This finding is not specific to these genes. We also generated knockouts in two other Ras-D-dependent genes and found their changes in expression in response to gefE gene disruption (rrgB increases, and rigB decreases) could also predict their effects on cell fate choice. Therefore, consistent with their patterns of expression, genes in this network are required for normal cell fate choice when cells are forced into low or high RasD states.

**Natural Variation in Cell Cycle Dynamics**

**Changes in Ras-Dependent Gene Expression**

Gene expression changes in response to glucose deprivation or RasD levels suggest that Ras-D-dependent gene expression provides a buffering mechanism to counter the effects of extrinsic variation on the cell cycle and thus cell fate. We, therefore, tested whether these effects are specific to nutritional history or also seen in response to other environmental perturbations. For example, drugs that affect intracellular pH have previously been shown to affect cell fate choice (Gross et al., 1983; Kubohara et al., 2007). Hence, we tested whether growth at different pHs also affects cell fate choice. When compared to G+ cells grown in “normal” pH−6.8 medium, pH−7.5 cells exhibit a sporey cell fate bias in chimera. This contrasts with nutritional perturbation, where G− cells exhibit a stalky bias (Figure 6A). qPCR revealed that while stalky G− growth resulted in a sporey gene expression profile (increased rrgA versus rigA), sporey pH−7.5

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**Figure 5. Ras Activity-Dependent Gene Expression Is Coupled to Changes in Cell Cycle Dynamics**

(A) RNA-seq reveals reciprocal genes expression changes in response to GefE and nutrition. RNA-seq was performed on AX3 G+, AX3 G−, and gefE− G+ cells. Ras-dependent genes were identified by comparing gene expression of gefE− G+ and AX3 G+ samples. A highly significant proportion of these 45 DEGs (80%) also show a >1.5 × FC in expression between AX3 G+ and AX3 G− (hypergeometric test, p = 4 × 10−15). Comparison of log2 FCs of the 36 overlapping genes shows that stalky nutritional bias results in sporey low Ras gene activity. Candidate genes are highlighted in red.

(B) RNA-FISH of rrgA and rigA. rrgA and rigA are heterogeneously expressed, and expression levels depend on RasD activation.

(C) Quantification of rrgA and rigA RNA-FISH. The relative expression of each gene in each cell was calculated as an index of expression of rrgA/rrgA + rigA. Values of 1 or 0 represent only rrgA or rigA expression, respectively.

(D) Ras-dependent genes are maximally expressed during multicellular development. Expression profiles of Ras-D-dependent genes were determined from RNA-seq data at different time points during development.

(E) rrgA and rigA mutant behavior in stalk cell induction monolayer assays. rrgA knockout in AX3 does not significantly affect stalk cell differentiation, whereas the rrgA−/gefE double mutant rescues gefE− mutant prespore bias (see supplementary Mendeley data for p values). rigA knockout in AX3 does not significantly affect stalk cell differentiation, whereas the rrgA/AX3<sup>rasD</sup>(G12T) strain shows significantly reduced stalk cell differentiation compared to AX3<sup>rasD</sup>(G12T). Plotted values are the mean of three replicates, and error bars depict the SEM.

(F) Behavior of Ras-D-dependent gene knockouts in chimeric development. Labeled cells were developed in a 50:50 ratio with unlabeled AX3 cells. The rrgA and rrgB single mutants do not significantly affect cell fate choice. The rrgA−/rrgB− knockout in the gefE− mutant rescue the tip and prespore bias seen in the gefE− single mutant. The dicitelBase: DDB122407655 and rigB single mutants do not significantly affect cell fate choice. The rigA−/AX3<sup>rasD</sup>(G12T) and rigB−/AX3<sup>rasD</sup>(G12T) strains show significantly reduced collar and back stalk cell differentiation bias compared to AX3<sup>rasD</sup>(G12T).
**Figure 6. Ras Activity Buffers Extrinsic Fluctuations that Affect Cell Cycle Dynamics**

(A) Altering nutrition or pH during growth affects cell fate choice. RFP labeled or unlabeled cells were grown in G+ medium at pH 6.8 and 7.5 or in G− medium at pH 6.8. Chimeric development revealed cells at more alkaline pH are biased to the prespore cell fate, whereas low glucose results in a stalky bias.

(B) Altering nutrition or pH during growth affects Ras network activity. Expression of rrgA and rigA was measured in G+ pH 6.8 and 7.5 and G− medium pH 6.8 by qPCR (3 replicates each). In stalky G− cells, the expression of the low Ras activity reporter rrgA increases relative to high Ras activity reporter rigA expression. This trend is reversed in sporey alkaline pH. Error bars depict SEM.

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growth resulted in a stalky gene expression profile (increased rigA versus rrgA) (Figure 6B).

The directionality of gene expression changes in response to pH or levels of glucose is consistent with a buffering mechanism. We therefore next tested whether these gene expression changes are required for cell type proportioning in response to changes in extrinsic conditions. Knockout of rrgA was found to have no effect when cells were grown under normal G+ conditions (Figures 6C and 6D) in chimeric development and monolayer differentiation. However, when rrgA knockout cells were grown in G− conditions (i.e., when rrgA gene expression is normally induced), clear defects were seen. In chimeric development, G− mutant cells were found to adopt the stalky collar and back fate in chimeric slugs, even when mixed with G−-grown wild-type cells (Figure 6Cl). Furthermore, mutant cells were hypersensitive to the effects of G− growth and showed an increased propensity to differentiate as prestalk cells in monolayer cell culture (Figure 6D).

The observed requirement for rrgA in specific growth conditions is consistent with the idea that changes in GefE/RasD activity lessen the effects of cell cycle perturbations. Therefore, we next determined whether changes in GefE/RasD network activity could also be seen under normal physiological conditions. For example, a small number of cells show a significantly longer than average cell cycle (i.e., exhibit “natural” cell cycle perturbation) even when grown in G+ conditions. These cells also have a slight tendency to become stalk cells (Figure 3C). We, therefore, tested whether they also decrease RasD network activity (e.g., increase rrgA expression) in an attempt to compensate. For this, we visualized and quantified gene network activity in living cells. A dual reporter strain was generated in which the promoters of rigA and rrgA were used to drive GFP and RFP expression, respectively. This showed similar behavior to that seen by RNA-seq and RNA FISH (Figure 6E). We next used live cell imaging to simultaneously monitor cell cycle dynamics and levels of rrgA/RFP expression in single cells. The cell cycle length of each cell that initiated rrgA/RFP reporter gene expression was measured. Because translation and folding of RFP protein are likely to introduce a delay in the system, we also measured the length of the previous cell cycle. These data were then compared to the cell cycle length of the population as a whole. Cells that activated rrgA reporter expression, and thus down-regulated Ras activity, were found to have recently exhibited a significantly longer cell cycle than the average of the general population (Figures 6F and 6G). Together, these data strongly support the idea that coupling of the signal response threshold through the Ras gene network to the cell cycle dynamics serves to buffer extrinsic fluctuations in a cell fate oscillator and maintain robust cell fate proportioning.

A Simple Model for Cell Cycle Control of Cell Fate

Our data suggest that a noisy cell cycle oscillator controls cell fate choice in Dictyostelium, while coupling of cell cycle dynamics to response thresholds confers robustness in the face of extrinsic environmental perturbation. In order to gain insights into the underlying mechanism, we used our highly quantitative data to build a model for cell cycle control of cell fate choice. The model was based on three key experimental observations. First, the probability of stalk cell differentiation is highest in cells that have just undergone mitosis. Therefore, the model assumed the existence of a cell cycle-associated factor (CCAF) that determines the stalk cell differentiation propensity (χ), with a propensity of 1, indicating that it will almost certainly adopt the stalk fate. Second, χ declines gradually following mitosis. Hence, the model assumed the breakdown of CCAF is gradual with rate λ, rather than stepped. The propensity of any cell to adopt the stalk fate was then modeled as a function of the time since mitosis (Figure 7A; STAR Methods). Finally, χ rises in cells that are at the end of G2 but is still significantly lower than cells that have just divided. Consequently, the model assumed the existence of a checkpoint at the end of G2 and that CCAF levels rise rapidly between this checkpoint (ς) and mitosis. We found that the inclusion of this checkpoint ς before mitosis, after which CCAF levels rise to saturation levels instantaneously and are then held constant until after mitosis, resulted in a slight rise in χ in cells with a longer than average cell cycle length (as they are more likely to be close to this checkpoint). This cell fate model could also be coupled to the previously described stochastic model that we constructed for cell cycle length variation in cell populations and thus also takes into account the observed correlation of cell cycles between sister cells. Thus, having created a population of cells using this stochastic model, we were able to assign fates to each cell based on the probability given by the propensity function in Figure 7A. Figure 7B demonstrates the excellent fit between the experimentally observed behavior of wild-type cells grown under G+ conditions and the model.

To further test the model, we addressed whether it could also predict the behavior of G−-grown cells. Experimental data suggest they exhibit a 1–2 hr block at a G2/M checkpoint, resulting in a high propensity to adopt the stalk fate. We fitted the data for G− cells to the model, which resulted in a longer delay between ς and mitosis. This model reflected the observed data well, and G− cells blocked in late G2 exhibited a higher likelihood of stalk cell differentiation than G+ cells in late G2. This is because cells
blocked in late G2 have a higher probability to have passed the checkpoint and reached saturating CCAF levels. The model also predicted that the rate of decay of CCAF was the same as that seen for G+ cells (Figure 7C). Finally, we also addressed whether the model could help explain why lowering Ras activity decreases stalk bias. Simply changing the amount of CCAF that accumulated at mitosis allowed the experimental data to fit the model (Figure 7D). Therefore, while this simple model may not fully capture the true behavior of CCAF, it is able to recapitulate most experimentally observed features of the cell fate system.

**DISCUSSION**

**Buffering or Mistake Correction?**
Developmental systems must be robust to perturbation by extrinsic variation that affects signaling dynamics or biochemical reaction rates. In addition, intrinsic stochastic variation is an inevitable consequence of gene expression kinetics. Mistakes can be corrected through cell-cell communication if information about the relative proportions of the cell types is relayed. Our results suggest that the underlying circuits can also be buffered to ensure a robust and reproducible initial outcome. This may be because the cell cycle must exhibit stochastic variation to generate a sufficiently de-synchronized population of cells. This does, however, render the cell cycle susceptible to perturbations caused by shifts in external environmental conditions, such as nutritional availability. Here, we find that *Dictyostelium* cells have evolved an elegant solution, in which changes in the cell cycle result in compensatory changes in signal sensitivity and thus the probability of cell type differentiation (Figure S2). In this, extrinsic environmental changes that affect cell cycle dynamics trigger changes in a RasD responsive gene network. *In silico* analyses reveal this network is enriched in genes that have the potential to affect cell signaling responses, such as transmembrane and plasma membrane proteins. One of these, rigA, encodes a protein with sequence similarity to human histidine rich glycoprotein, which has been shown to modulate different signaling pathways (Jones et al., 2005). Another, rrgB, is related to components found in the yeast spindle pole body (centrosome in *Dictyostelium* and mammalian cells), which is now widely accepted to serve as a hub for the integration and coordination of signaling pathways, including cell cycle-related signaling (Arquint et al., 2014). Finally, it is interesting to note that RhgB encodes a rhesus-like glycoprotein and thus a putative ammonium transporter,
given long-standing observations linking ammonium and intracellular pH in cell fate choice and responses to cAMP and Dif-1 in Dictyostelium (Gross, 2009).

**Linking the Cell Cycle to Cell Fate Choice**

Links between cell cycle progression and cell fate decisions are well established. Recently, however, there has been renewed interest in the possibility that the phase of the cell cycle in which cells receive differentiation cues could affect fate choice. Much of this recent work has focused on differentiation events that occur in G1 (or indeed different phases of G1). However, a cell cycle-based mechanism has the potential to generate further cell type complexity if other phases of the cell cycle also affect fate choice. In fact, Dictyostelium cells have little or no G1. Instead, we find that prestalk cell differentiation is favored in S phase and early G2, and prespore cell differentiation in mid G2. It should also be noted that our studies reveal the potential for more complexity because multiple prestalk lineages are favored during S phase/early G2. It will be interesting to determine whether this is due to different windows of opportunity within this phase of the cell cycle or whether another level of regulation (perhaps stochastic) is layered on top of the cell cycle.

While our studies reveal that different phases of the cell cycle can affect lineage choice, the underlying mechanism is still poorly understood. Indeed, there are many differences between cells at different cell cycle stages, including cell size, nuclear volume, and DNA content. The activity of cell cycle regulators also rises and falls sharply as cells transition from one state to another, and these could directly affect regulators of developmental transcription. Indeed, we find that the decision of a cell to enter mitosis results in an abrupt shift in cell fate propensity. If this is due to the accumulation of a cell cycle regulatory complex, our studies will aid in its identification because they reveal the kinetics of its activity. Furthermore, due to the ease with which genes that influence developmental decisions can be isolated in Dictyostelium and tested for effects on the cell cycle or cell signaling, studies of this system will likely pave the way for a better understanding of this process.

**STAR★METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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**SUPPLEMENTAL INFORMATION**

Supplemental Information includes two figures, two videos, and three data files and can be found with this article online at https://doi.org/10.1016/j.develcel.2018.09.023.

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**AUTHOR CONTRIBUTIONS**

The project was devised and overseen by C.R.L.T. The manuscript was written by C.R.L.T., S.L.C., and N.G. Computational and bioinformatic analyses were carried out by N.G. Modeling was carried out by J.L. and W.M.S., and T.K. Chimera and RNA-seq experiments were carried out by K.P., W.M.S., C.A., and K.N. Ras reporter gene characterizations were carried out by S.K., E.J.J., W.v.Z., and K.N.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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**STAR METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and Virus Strains** | | |
| Klebsiella aerogenes | Dicty Stock Center | dictyBase: DBS0349838 |
| **Chemicals, Peptides, and Recombinant Proteins** | | |
| HLS5 medium | Formedium | HLG0102 |
| KK2 | This study | N/A |
| Oxoid purified agar | ThermoFisher | LP0028B |
| G418 | Sigma | 00000004727878001 |
| conditioned media | This study | N/A |
| DIF-1 | ENZO life sciences | BML-GR324-0100 |
| stalk medium | This study | N/A |
| EDTA | Sigma | E6758 |
| HCR probe sets and fluorophore-labeled HCR hairpins | Molecular Technologies | N/A |
| **Deposited Data** | | |
| SOLID sequencing data AX3 G+ / G-; gefE- G+/G- | This study | SRA: SAMN07834373 - SAMN07834380 |
| Illumina HiSeq 4000 81 AX3 single cells | This study | SRA: SAMN07833758 -SAMN07833838 |
| Datasets for Figures 1, 2, 3, 4, 5, and 6 and three additional figures | This study | https://doi.org/10.17632/rvny4rmfpp.1 |
| **Experimental Models: Organisms/Strains** | | |
| Dictyostelium discoideum AX3 | Dicty Stock Center | dictyBase: DBS0235539 |
| AX3 cells co-transformed with pspA-GFP and ecmAO-RFP | This study | N/A |
| AX3 cells co-transformed with ecmB-GFP and ecmAO-RFP | This study | N/A |
| AX3 cells co-transformed with ecmB-GFP and pspA-RFP | This study | N/A |
| AX3 cells transformed with PCNA-RFP intra-chromosomal reporter | This study | N/A |
| AX3 gefE- cells transformed with PCNA-RFP intra-chromosomal reporter | N/A | N/A |
| AX3 cells transformed with dual rrgA/rigA promoter reporter construct | This study | N/A |
| AX3 gefE- cells transformed with dual rrgA/rigA promoter reporter construct | This study | N/A |
| AX3 rrgA knockout | This study | N/A |
| AX3 rigA knockout | This study | N/A |
| AX3 dictyBase: DDB_G0292996 knockout | This study | N/A |
| AX3 dictyBase: DDB_G0281385 knockout | This study | N/A |
| AX3 dictyBase: DDB_G0293434 knockout | This study | N/A |
| AX3^ras^G12T | This study | N/A |
| AX3^ras^G12T | This study | N/A |
| AX3^ras^G12T | This study | N/A |
| **Software and Algorithms** | | |
| novoalignCS V1.04.01 | NovoCraft | N/A |
| HTseq-count | https://htseq.readthedocs.io/ | N/A |
| NGS QC toolkit | https://ccbr.github.io/Pipeliner/Tools/NGS_QC_Toolkit.html | N/A |
| bowtie2 | http://bowtie-bio.sourceforge.net/bowtie2/index.shtml | N/A |

(Continued on next page)
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Christopher R.L. Thompson (christopher.thompson@ucl.ac.uk).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Strain Growth, Maintenance, and Development

Dictyostelium discoideum AX3 strains were cultured on lawns of Klebsiella aerogenes or in HL5 medium with (G+) or without (G−) 86 mM glucose. Cells were grown for 2–4 days for G− phenotypes. All cultures were maintained at log phase (1–4 × 10^6 cells/ml) during this period. For development, amoebae were washed with KK2 (16.1 mM KH₂PO₄, 3.7 mM K₂HPO₄) and deposited onto KK2 plates containing 1.5% purified agar (Oxoid) at a density of 3.5 × 10⁶ amoebae/cm². Plates were incubated for 14–16 hours at 22°C in a dark, moist box then removed and allowed to complete development in the light. For chimeric development, amoebae of two different genotypes, where one strain was labelled by expression of GFP or RFP, were mixed and developed together. Images were taken after approximately 16 hours to determine any sorting bias. Stalk cell differentiation in monolayers was quantified using the cAMP removal assay described in (Thompson et al., 2004a). All mutants were generated in the AX3 background used in this study by the insertion of a blasticidin resistance cassette into the coding sequence of the gene by homologous recombination. Constructs for gefE (Chattwood et al., 2013), rasC (Lim et al., 2001), rasD (Wilkins et al., 2000) and rasG (Bolourani et al., 2006) disruption have previously been described. Gene deletions were generated for rrgA (dictyBase: DDB_G0268600), rigA (dictyBase: DDB_G0274655) and rrgB (dictyBase: DDB_G0270640). For rigB (dictyBase: DDB_G0272238), the blasticidin resistance cassette was introduced at position 1,724,548 of chromosome 2. Double mutants were generated by transient expression of Cre recombinase before transformation with second knockout vector. For induction of constitutively active Ras proteins, cells were grown in the presence of doxycycline for 6 hours, before washing and plating for development.

METHOD DETAILS

RNA-Seq Sample Preparation and Sequencing

RNA libraries from AX3 and gefE− cells grown in G+ or G− media (two replicates each) were sequenced on the SOLiDv4 platform resulting in 50bp single-end reads. For single cell sequencing, log phase cells were grown in tissue culture dishes before being harvested and resuspended at a density of 1×10⁶ cells/ml in HL5 for capture on a Fluidigm C1 flow cell using a medium size chip. This resulted in the capture of 81 individual cells, which were sequenced on an Illumina HiSeq 4000 resulting in 100bp paired-end reads (150bp insert). Raw reads from (Strasser et al., 2012) were downloaded from the Gene Expression Omnibus using accession number GSE30368 and processed in the same way as the other Illumina data.

Cell Type Differentiation in Low-Density Culture

AX3 cells were co-transformed with pspA-GFP and ecmAO-RFP, ecmB-GFP and ecmAO-RFP, or ecmB-GFP and pspA-RFP cell type specific reporters (Parkinson et al., 2009). Clonal lines were selected and maintained in 20 μg/ml G418. For quantification of cell type induction, log phase cells were seeded in 10 cm tissue dishes at a density of 1×10⁶ cells/ml in 9 ml conditioned media with or without 10nM DIF-1 (minimum of ten plates per experiment) to induce differentiation. Conditioned media was made by plating cells at a density of 1×10⁶ cells/ml in stalk medium (10mM MES, pH6.2, 1mM CaCl₂, 2mM NaCl, 10mM KCl, 200μg/ml streptomycin sulphate) containing 5mM cAMP for 16 hours. After 16 hours, the media was collected and any cells removed by centrifugation. Freshly prepared conditioned media was prepared for each experiment performed. After 20 hours of incubation in conditioned media with or without DIF-1, cells were harvested from plates in 1ml KK2 plus 20mM EDTA. The proportion of fluorescent cells was determined by FACs (>10,000 cells per experiment). The data from all strains containing a given reporter gene were averaged, resulting in
two biological replicates for each reporter construct. In addition, each induction was carried out on three separate occasions, resulting in a total of six biological replicates per cell type reporter. From this data, we were able to infer the percentage of cells that either do not differentiate or are non-transformed (100 – (pspA+ecmA+ecmB)). For tracking of growth and differentiation, log phase cells were seeded at a density of $4 \times 10^3$ cells/ml in 750 μl H5L medium in a 3 cm glass bottomed imaging dish. Cells were filmed in multiple positions for 12-14 hours (i.e. until cell reached a density of approximately $1 \times 10^4$ cells/ml) at a frame rate of one image every 5 minutes to allow cell tracking. The growth media was then removed and the cells carefully washed twice with KK2 (16.1mM KH2PO4, 3.7mM K2HPO4). 750μl of conditioned media plus 10nM DIF-1 was added to the washed cells to induce differentiation and cells were filmed for a further 16-20 hours at a frame rate of one image every 5 minutes. A final fluorescence image was captured to observe reporter gene expression. Cells were tracked by hand using the Fiji software from the beginning of the movie. The frame when a cell last divided prior to addition of conditioned media was recorded. Cells were tracked throughout the remainder of the movie and their final cell fate was recorded, as determined by reporter gene expression. The cell cycle stage at the time of induction was inferred from the amount of time between the addition of conditioned media, and the last cell division before induction. The same method was used for tracking the growth and differentiation of G- cells, but these cells were conditioned prior to the experiment by growing the cells in log phase in H5L without glucose for 48 hours.

**Assaying Cell Cycle Lengths**

Cells were grown in G+ or G- media and imaged every 4 min for 16 hours. The time between two cell divisions was manually extracted by tracking single cells. Sister cells were tracked and their cell cycle was measured if a cell divided into exactly two cells. Mother and daughter cells were noted if the next division of the daughter cell occurred before the end of the movie.

**Assaying Cell Cycle Phase Lengths**

Cell cycle phase lengths were assessed using a PCNA-RFP intra-chromosomal reporter (Muramoto and Chubb, 2008). Cells were seeded at $1 \times 10^4$ cells/ml in filter-sterilised H5L with or without 75mM glucose and grown for 48 hours (growing to 3-5x10^6 cells/ml).

**Assaying Ras Network Activity**

To generate a dual rrgA/rigA promoter reporter construct, the promoters of each gene were amplified and cloned into the XhoI/BglII sites of either pDM323 (rrgA) or pDM324 (rigA). The pDM323- rigA:GFP construct was digested with XhoI/HindIII, and blunt end cloned into the NgoMIV site of pDM324- rigA:RFP. Constructs were electroporated into vegetative AX3 and getE+ cells, and selected with 10-20μg/ml G418. For time-lapse microscopy, the same protocol as used to follow PCNA localisation was used. For RNA-FISH, we performed single-molecule hybridization chain reaction (smHCR) using the In Situ HCR v2.0 protocol for tissue sections on slides provided by Molecular Technologies. In brief, we fixed vegetative cells in -20C methanol for 1 min. Fixed cells were treated with 0.5% TritonX-100/PBS for 10 min. HCR probe sets and fluorophore-labeled HCR hairpins were purchased from Molecular Technologies. Samples were hybridized overnight with 2 nM of each HCR probe. HCR amplification was performed overnight with 60 nM of each HCR hairpin (conjugated to Alexa488 or Alexa594 to visualize rrgA and rigA mRNA respectively). We manually segmented the boundaries of cells and measured the mean signal intensity of Alexa488 and Alexa594 using ImageJ (NIH).

**Mathematic Modeling**

**Stochastic Model for Cell Cycle Variation**

In the model, after mitosis, the cell cycle lengths of each pair of daughters are drawn from a joint probability distribution. We denote the length of the daughters’ cell cycles $\tau_1$ and $\tau_2$ respectively. $\tau_1$ is drawn from a Gamma distribution with parameters $a$ and $b$ for the shape and rate respectively. The difference in times $\tau_1$ and $\tau_2$ is then assumed to be a mean-zero Laplacian random variable with standard deviation $\sigma$, so that

$$\tau_1 \sim \text{Gamma}(a,b), \quad (\tau_2 - \tau_1) \sim \text{Laplace}(\sigma).$$

The mean and variance of these cell cycle times can be computed to be equal to

$$E(\{\tau_1, \tau_2\}) = \frac{a}{b}, \quad \text{Var}(\{\tau_1, \tau_2\}) = \frac{2a + b^2\sigma^2}{2b^2}.$$

The variance $\sigma^2$ of the sister-sister difference in cell length is set to be equal to the usual unbiased estimator of the variance found in the data. Likewise, unbiased estimators of the mean and variance of the cell cycles across the whole data set is then used in turn to ascertain the values of $a$ and $b$.

**Stochastic Model for Cell Fate**

We construct a model to link the position of a cell in the cell cycle with its fate. In particular, we propose a simple function to represent the propensity that a given cell will become a stalk cell. This is a function of $t$, the current time, and where $t_1$ and $t_n$ are the times of the last and next mitosis event for the cell respectively.

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The values of the parameters $\gamma$, $\lambda$, and $\alpha$ are fitted to the observed data regarding cell fate. In particular, we aim to match the behaviour seen in each scenario presented in Figures 7B–7D. A least squares fit for $\gamma$ and $\lambda$ is used to match the exponential decay part of the function to the first six data points of each of these datasets. The value of $\alpha$ was determined by assuming all cells which have passed the checkpoint have stalk propensity equal to $\min(\gamma,1)$. In hours 7-11+, the proportion of cells in the last $\alpha$ minutes of their cell cycle was calculated, and then a value for $\alpha$ was determined in each case using a least squares fit to the data. The values found for each of these variables can be seen in each figure.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**RNA-Seq Re-processing**

For the SOLID data, three pre-processing steps using the SOLID-pre-processor were included in the downstream analysis to remove poor quality sequences. Firstly, all transcripts were trimmed to 40bp from the 3’ end. Next, any sequences containing an unresolved base (-1 quality score, missing colour call) were also removed, as the sequence after this base call is likely to be wrong or ambiguous at best. In addition, sequences that contained 3 or more bases with a quality score of less than 22 were removed, as these bases are highly likely to be wrong, which would affect the reliability of the read towards the 3’ end. The resulting high quality reads were mapped against the 30.04.2012 *Dictyostelium discoideum* reference genome sequence (after masking the inverted repeat on chromosome 2) using novoalignCS V1.04.01 (parameters used: -l 25 -H 20 -r E 2 -s 5 -t 99 -n 40 -o SAM -k -c 8). Counts for each gene were then determined using HTseq-count and the option intersection-nonempty, while strand information was taken into account. For the Illumina data, paired-end reads from all samples were quality checked and filtered using the IlluQC_PRLl.pl script (v2.3) from the NGS QC toolkit and a read length cut-off of 70% together with a quality score cut-off of 20. In all samples more than 90% of the reads were retained, which were then mapped against the 30.04.2012 *Dictyostelium discoideum* reference genome sequence version (after masking the inverted repeat on chromosome 2) using bowtie2 version 2.0.0-beta5 and very-sensitive end-to-end mapping parameters requesting the best out of ten alignments. Resulting samfiles were filtered for uniquely mapped reads, sorted and converted into bamfiles using samtools (0.1.18 (r982:295)). The annotation file available at Dictybase was then used to count reads mapping to annotated genes using HTseq-count and the option intersection-nonempty, while strand information was taken into account.

**RNA-Seq Data Analyses and Statistical Tests**

Single cell data: Normalised expression profiles for each cell were correlated and cells with a low total read count (less than 10^6.2 reads) and low average correlation to all other cells (less than 0.4) were excluded from the analysis resulting in 81 cells. Genes with low average read counts were also excluded resulting in 11,320 genes. Normalised read counts were then log transformed using DESeq2 (v. 1.16.1; and a PCA was used to identify two clusters of cells. Marker genes for each cluster (701 and 901) were identified with low average read counts were also excluded resulting in 11,320 genes. Normalised read counts were then log transformed using DESeq2 using an adjusted $p$-value of 0.05 and a foldchange cutoff of 1.5.

Cell cycle: Normalised read counts for the technical replicates were summed, while read counts for the biological replicates were averaged. Cell cycle profiles for each gene were determined by firstly computing the percentage of the expression for each time point and gene by dividing the normalised read counts for each time point by the sum of all normalised read counts. Then, outliers with expression of 1.5 x MAD (median absolute deviation) higher than the median were identified. Genes were clustered according to the highest outlier per gene while excluding genes with low average read counts and/or no outliers from the analysis. Over- and underrepresentation of DEGs or marker genes in specific cell cycle clusters was determined using the hypergeometric test implemented in the phyper function in R.

GO term enrichment: Enriched GO terms were identified using the statistical overrepresentation test (release 20160715) implemented on http://pantherdb.org using a $p$-value cutoff of 0.05 and the ‘GO biological process complete’ database as annotation data set.

**DATA AND SOFTWARE AVAILABILITY**

Raw reads from all samples were uploaded to the NCBI short read archive (see Files S1 and S3 for accession numbers). Raw and processed data for Figures 1, 2, 3, 4, 5, and 6 were uploaded to Mendeley data (https://doi.org/10.17632/rvny4rmfpp.1).