Filamentation Regulatory Pathways Control Adhesion-Dependent Surface Responses in Yeast

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ABSTRACT Signaling pathways can regulate biological responses by the transcriptional regulation of target genes. In yeast, multiple signaling pathways control filamentous growth, a morphogenetic response that occurs in many species including fungal pathogens. Here, we examine the role of signaling pathways that control filamentous growth in regulating adhesion-dependent surface responses, including mat formation and colony patterning. Expression profiling and mutant phenotype analysis showed that the major pathways that regulate filamentous growth [filamentous growth MAPK (fMAPK), RAS, retrograde (RTG), RIM101, RPD3, ELP, SNF1, and PHO85] also regulated mat formation and colony patterning. The chromatin remodeling complex, SAGA, also regulated these responses. We also show that the RAS and RTG pathways coregulated a common set of target genes, and that SAGA regulated target genes known to be controlled by the fMAPK, RAS, and RTG pathways. Analysis of surface growth-specific targets identified genes that respond to low oxygen, high temperature, and desiccation stresses. We also explore the question of why cells make adhesive contacts in colonies. Cell adhesion contacts mediated by the coregulated target and adhesion molecule, Flo11p, deterred entry into colonies by macroscopic predators and impacted colony temperature regulation. The identification of new regulators (e.g., SAGA), and targets of surface growth in yeast may provide insights into fungal pathogenesis in settings where surface growth and adhesion contributes to virulence.

KEYWORDS MAPK pathways; signaling networks; expression profiling; fungal pathogenesis; microbial predator–prey relationships; pseudohyphal growth; invasive growth; biofilm; mat; complex colony; temperature control

responses. Under certain conditions, fungal cells can differentiate into filamentous or hyphal cells that can expand across, and/or penetrate into, new environments (Soll and Daniels 2016). Many fungal species can also grow in communities of biofilms or mats, which are composed of interconnected cells that attach to each other and to surfaces. One property of mats is the formation of highly organized patterns that result from adhesive contacts between cells. In pathogens, filamentous growth (Lo et al. 1997) and biofilm forma-

tion (Desai *et al.* 2014) are critical determinants of virulence. For example, cells can adhere to medical devices and grow in dense mats that are resistant to antifungal medicines (Chandra *et al.* 2001; Sudbery *et al.* 2004; Kumamoto 2005; Ramage *et al.* 2005; Nett and Andes 2015).

The budding yeast *Saccharomyces cerevisiae* is a unicellular fungal microbe, and a convenient model for studying nutrient-regulated foraging responses like filamentous growth and mat formation. These responses are best studied in "wild" strain backgrounds (such as $\Sigma 1278b$) as the responses have been lost in certain laboratory strains due to genetic manipulation (Liu et al. 1996; Dowell et al. 2010; Chin et al. 2012). During filamentous growth, yeast cells differentiate into elongated and polarized filaments that remain connected in pseudohyphae (Gimeno et al. 1992; Cullen and Sprague 2012).

At least 600 genes have been identified by genetic screens (Lorenz and Heitman 1998; Palecek *et al.* 2000) and genome-wide studies (Jin *et al.* 2008; Xu *et al.* 2010; Ryan *et al.* 2012) that play

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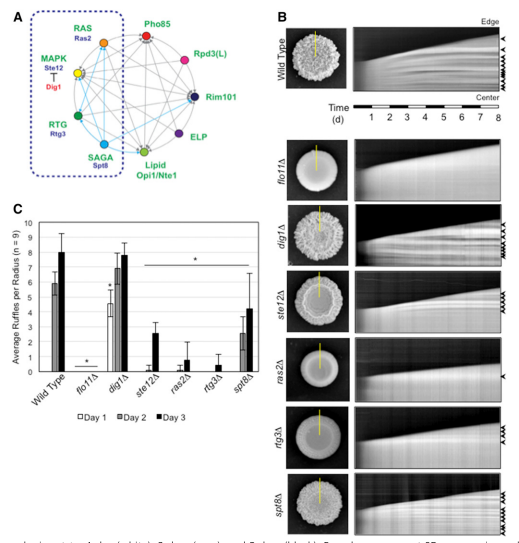


Figure 1 Functionally interconnected signaling pathways regulate mat expansion and colony pattern formation over time. (A) Circles represent signaling pathways or regulatory complexes that control filamentous growth. Arrows refer to target genes coregulated by pathways, as adapted from Chavel et al. (2014). Cyan arrows refer to functional connections identified in this study. The dashed blue box marks the part of the network on which the study is focused. (B) Timecourse experiment of colony pattern formation in wild-type cells and the indicated mutants. Colony expansion was examined over an 8-day time period (Videos S1-S7). In the left panels, colonies at the 8-day time period are shown. On the right are kymographs that show colony expansion over time. The yellow lines in the left panels refer to the region selected for kymograph analysis. The panels on the right show colony patterns that developed over time in the region selected by the yellow line, where the bottom of the figure represents the colony center and the top represents the colony perimeter (edge). Arrows mark features that are characteristic of ruffle formation, which is quantitated in (C). (C) Bar graph showing the average number of ruffles in wildtype and the indicated mutant

colonies at t = 1 day (white), 2 days (gray), and 3 days (black). Error bars represent SD among nine radii from three biological replicates. * P < 0.05 between the mutant and wild-type for the same time point.

some role in filamentous growth. A subset of these genes encode signaling pathway components that include at least four major nutrient-sensing pathways [filamentous growth MAPK (fMAPK), RAS, TOR (target of rapamycin), and SNF1], as well as pathways that regulate the response to pH (RIM101), phosphate utilization (PHO85), and mitochondrial stress [the retrograde mitochondriato-nucleus (RTG) pathway]. In addition, proteins that control the epigenetic modification of histones to alter gene expression have also been implicated in the regulation of filamentous growth (Rpd3p).

Many of the pathways that regulate filamentous growth are functionally connected through their ability to coregulate common target genes (Figure 1A). In some cases, this occurs at the level of transcription. In a pioneering study, it was shown that many of the transcription factors that control filamentous growth control each other's expression (Borneman *et al.* 2006). Transcription factors can also converge at common promoter elements. One example is the gene encoding the major cell adhesion molecule in yeast,

Flo11p (Kraushaar et al. 2015; Chan et al. 2016). The FLO11 gene contains one of the largest and most highly regulated promoters in the yeast genome, and functions as a hub where multiple transcription factors and chromatin remodeling enzymes bind (Robertson and Fink 1998; Rupp et al. 1999; Palecek et al. 2000; Pan and Heitman 2000; Kuchin et al. 2002; van Dyk et al. 2005; Barrales et al. 2008). Signaling pathways that control filamentous growth can also regulate each other's activity. The classic example comes from the discovery that the RAS pathway can regulate the activity of the fMAPK pathway (Mösch et al. 1996). It is now clear that many pathways regulate the activity of the fMAPK pathway (Chavel et al. 2010, 2014). One way this may occur is through the protein kinases of the major regulatory pathways, which can regulate each other's localization and activity (Bharucha et al. 2008).

Yeast can also undergo mat formation (Reynolds and Fink 2001; Bojsen *et al.* 2012), where colonies expand radially across surfaces and form ruffled patterns (Granek and

Magwene 2010; Granek et al. 2011; Tam et al. 2018). Mats form wheel-spoke patterns in low-percentage agar media [0.3% agar (Reynolds and Fink 2001)], and wrinkled or ruffled colonies on high-percentage agar media [1-4% (Granek and Magwene 2010; Karunanithi et al. 2012)]. Mat expansion and patterning require contacts between cells that are also mediated by Flo11p. Regulators of mat formation have been identified by direct approaches (Reynolds 2006; Sarode et al. 2011; Váchová et al. 2011), genome-wide deletion screens (Ryan et al. 2012; Voordeckers et al. 2012; Scherz et al. 2014), overexpression screens (Cromie et al. 2017), and comparative expression profiling (Traven et al. 2012; Maršíková et al. 2017). Many of the molecular pathways that control mat formation regulate FLO11 expression. In addition, several regulators of mat formation, like the cell wall integrity sensor Wsc1p (Sarode et al. 2014) and vacuolar protein-sorting genes (Sarode et al. 2011), appear to regulate mat formation by mechanisms that do not involve Flo11p. Despite the identification of many proteins and pathways that regulate mat formation, the benefits that cells derive from forming these complex patterns remain unclear. Furthermore, the stresses encountered during colonial surface growth remain relatively unexplored.

As seen in other fungi, filamentous growth and mat formation are related responses in yeast. Both responses require Flo11p (Lo and Dranginis 1998; Reynolds and Fink 2001), and a corresponding set of regulatory proteins and pathways that control FLO11 expression (Ryan et al. 2012). These include TOR (Cutler et al. 2001; Bojsen et al. 2016), Rim101 (Li and Mitchell 1997; Sarode et al. 2011; Voordeckers et al. 2012), fMAPK (Roberts and Fink 1994; Reynolds and Fink 2001; Granek and Magwene 2010), and RAS (Gimeno et al. 1992; Granek and Magwene 2010; Zara et al. 2011; Ryan et al. 2012; Bojsen et al. 2016). The tRNA modification complex elongator (ELP) regulates all three responses (Abdullah and Cullen 2009). It has also been shown that under nutrient-limiting conditions, mats are composed of filamentous cells (Karunanithi et al. 2012). Given the abovementioned connections between filamentous growth and mat formation, we sought to further define how the network of signaling pathways that control filamentous growth might regulate adhesion-dependent surface growth.

To learn more about the regulation of surface growth in yeast, adhesion-dependent surface responses were examined from several perspectives. In one set of experiments, we developed a method for recording colony pattern formation over time by kymograph analysis, which allowed us to confirm that the major signaling pathways that control filamentous growth also regulate mat formation. A new regulator of filamentous growth, the chromatin remodeling complex SAGA, was also uncovered. Comparative expression profiling identified a large number of target genes as well as new regulatory connections between pathways. By seeking to understand why cells form adhesive connections during surface growth, we identified a role for Flo11p in protection from macroscopic predators. We also showed that colony ruffling aids

in temperature regulation. Our findings broaden the role of filamentation regulatory pathways to include the regulation of adhesion-dependent surface responses. Our findings may be relevant to studies of fungal pathogens that must tolerate growth on surfaces to become virulent.

Materials and Methods

Media and growth conditions

Yeast strains were grown and manipulated by standard methods (Sambrook et al. 1989; Rose et al. 1990). For most experiments, colonies were grown on standard YEPD (2%) or YEP-galactose (Gal) semisolid agar media (Cullen 2015a). Some experiments were performed on low-agar (0.3%) media. Glutamate sensitivity experiments used synthetic agar (dextrose, 2%; yeast nitrogen base, 6.7 g/liter; uracil, 20 mg/liter; His, 20 mg/liter; Leu, 120 mg/liter; adenine, 20 mg/liter; Lys, 60 mg/liter; Arg, 20 mg/liter; Trp, 20 mg/liter; Tyr, 30 mg/liter; Thr, 200 mg/liter; Met, 20 mg/liter; and Phe, 50 mg/liter), minimal agar (dextrose, 2%; yeast nitrogen base, 6.7 g/liter; and uracil, 20 mg/liter), and minimal + Glu (dextrose, 2%; yeast nitrogen base, 6.7 g/liter; uracil, 20 mg/liter; and Glu 20 mg/liter) media. To generate low-oxygen (5-15% oxygen) and anaerobic conditions, agar plates were incubated in GasPAK EZ Campy Pouch System (BD 260685; Becton, Dickinson and Company, Franklin Lakes, NJ) or BD GasPAK EZ Anaerobe Pouch System (BD 260683; Becton, Dickinson and Company) bags. To maintain consistent moisture levels, plates were poured and left unwrapped for 3 days at 22° to allow the evaporation of excess moisture. To generate media with reduced moisture levels, YEPD plates were left unwrapped for 10 days at 22°.

Strains and genetic manipulations

Yeast strains are listed in Table 1. Gene deletions were constructed using auxotrophic markers amplified by PCR and introduced into yeast by lithium acetate transformation by standard methods as described (Gietz and Schiest 2007). To generate the *spt8*Δ mutant, homologous recombination at the SPT8 locus was performed using the pKlURA3 (PC5225) cassette as a template. For some experiments, yeast strains were used from an ordered knockout collection (Ryan *et al.* 2012). *Caenorhabditis elegans* strains used in this study include N2 Bristol wild-type and KP4 *glr-1(n2461)* (Kaplan and Horvitz 1993). Strains were maintained at 20° under standard conditions on nematode growth media (NGM) agar plates seeded with OP50 *Escherichia coli* bacteria (Brenner 1974).

Assays for mat formation and filamentous growth

Assays for mat formation were performed as described (Reynolds and Fink 2001; Karunanithi *et al.* 2012). The plate-washing assay was performed as described (Cullen 2015b).

Microscopy

Mats were examined by bright-field microscopy using an Axioplan 2 fluorescent microscope (Zeiss [Carl Zeiss], Thornwood, NY) with $5\times$, $10\times$, $20\times$, $40\times$, and $100\times$ PLAN-APOCHROMAT

Table 1 Yeast strains used in the study

| Strain ^a | Description | Reference |
|---------------------|---|------------------------------|
| PJ69- | MATa trpl-901 leu2-3,112 ura3-52 his3-200 ga14Δ ga180Δ LYS2::GALI-HIS3 GAL2-ADE2 | James <i>et al.</i> (1996) |
| $4A^b$ | met2::GAL7-lacZ | |
| PC313 | MAT a ura3-52 | Liu <i>et al.</i> (1993) |
| PC538 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 | Cullen et al. (2004) |
| PC549 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 ste20::URA3 | Cullen and Sprague (2000) |
| PC555 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 hsl7::URA3 | Cullen and Sprague (2000) |
| PC562 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 ras2::URA3 | Cullen and Sprague (2002) |
| PC611 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 ste11::URA3 | Cullen and Sprague (2002) |
| PC999 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 MSB2-HA | Chavel et al. (2010) |
| PC1029 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 flo11::KanMX6 | Karunanithi et al. (2010) |
| PC1079 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 ste12::URA3 | Karunanithi et al. (2010) |
| PC2523 | MATa ura3-52 ste4 FUS1-lacZ -::NAT FUS1-HIS3 flo8::HYG | Chavel et al. (2010) |
| PC2712 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 GAL-FLO11::KanMX6 | Karunanithi et al. (2010) |
| PC3030 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 MSB2-HA sin3::NAT | Chavel et al. (2010) |
| PC3039 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 MSB2-HA dig1::NAT | Chavel et al. (2010) |
| PC3431 | MATa ura3-52 ste4 FUS1-lacZ -::NAT FUS1-HIS3 MSB2-HA sfl1::KlURA3 | Chavel et al. (2010) |
| PC3642 | MATa ste4 FUS1-lacZ FUS1-HIS3 ura3-52 MSB2-HA rtg3::NAT | Chavel et al. (2010) |
| PC3652 | MATa ste4 FUS1-lacZ FUS1-HIS3 ura3-52 MSB2-HA rtg2::NAT | Chavel et al. (2010) |
| PC3695 | MATa ste4 FUS1-lacZ FUS1-HIS3 ura3-52 MSB2-HA rtg1::NAT | Chavel et al. (2014) |
| PC4005 | MATa ste4 FUS1-lacZ FUS1-HIS3 ura3-52 gcn5::KlURA3 | This study |
| PC3688 | MATa ste4 FUS1-lacZ FUS1-HIS3 ura3-52 opi1::NAT | Chavel et al. (2010) |
| PC4008 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 spt8::KIURA3 | This study |
| PC5090 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 nte1::NAT | Chavel <i>et al.</i> (2014) |
| PC5119 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 pho80::NAT | This study |
| PC6733 | MATa ura3-52 ste4 FUS1-lacZ FUS1-HIS3 ura3-52::pTEF2-mCherry URA3 his5::NAT mCherry::- GFP _Y -HIS | Chow et al. (2019) |
| PC6016 | MATa can1Δ::Ste2pr-spHIS5 lyp1Δ::Ste3pr-LEU2 his3::hisG leu2Δ0 ura3Δ0 | Ryan et al. (2012) |
| CB13A9c | MATa can1Δ::Ste2pr-spHiS5 lyp1Δ::Ste3pr-LEU2 his3::hisG leu2Δ0 ura3Δ0 dan1Δ | Ryan <i>et al.</i> (2012) |
| CB52G2 | MATa can1Δ::Ste2pr-spHIS5 lyp1Δ::Ste3pr-LEU2 his3::hisG leu2Δ0 ura3Δ0 sip18Δ | Ryan <i>et al.</i> (2012) |
| CB53F12 | MATa can1Δ::Ste2pr-spHIS5 lyp1Δ::Ste3pr-LEU2 his3::hisG leu2Δ0 ura3Δ0 sno4Δ | Ryan <i>et al.</i> (2012) |
| CB60E1 | MAT a can1Δ::Ste2pr-spHlS5 lyp1Δ::Ste3pr-LEU2 his3::hisG leu2Δ0 ura3Δ0 gre1Δ | Ryan <i>et al.</i> (2012) |
| CB29E8 | MATa can1Δ::Ste2pr-spHIS5 lyp1Δ::Ste3pr-LEU2 his3::hisG leu2Δ0 ura3Δ0 hsp26Δ | Ryan <i>et al.</i> (2012) |

 $^{^{}a}$ Unless indicated, strains are derived from the Σ 1278b strain background.

100×/1.4 (oil) (N.A. 0.17) objectives. Digital images were obtained with an Axiocam MRm camera (Zeiss). Axiovision 4.4 software (Zeiss) was used for image acquisition. Digital images were imported into ImageJ (https://imagej.nih.gov/ij/) in 8-bit format.

Comparative RNA sequencing analysis

To compare the transcriptional response of wild-type cells (PC538) and the $ste12\Delta$ (PC1079), $dig1\Delta$ (PC3039), $rtg3\Delta$ (PC3642), $spt8\Delta$ (PC4008), and $ras2\Delta$ (PC562) mutants, cells were concentrated (OD A₆₀₀ = 20) and spotted in 10-µl aliquots onto YEP-Gal (2% agar) for 24 hr. Cells were spotted as six colonies per plate, equidistant to each other and the plate center. All six colonies were harvested for each trial and three separate trials were compared for each strain. The entire colony surface was scraped into 500 µl of distilled water, harvested by centrifugation, washed, and stored

at -80° . RNA was harvested by hot-acid phenol-chloroform extraction as described (Adhikari and Cullen 2014). Samples were further purified using a QIAGEN RNeasy Mini Kit (catalog number 74104; QIAGEN, Valencia, CA). RNA concentration and purity was measured via NanoDrop (NanoDrop 2000C; Thermo Fisher Scientific, Waltham, MA). RNA stability was determined by running the sample on an agarose gel.

RNA sequencing (RNA-seq) was performed as described previously (Adhikari and Cullen 2014) by sequencing RNA prepared from three separate cultures. RNA-seq libraries were prepared from total RNA using the TruSeq RNA Sample Prep Kit (Illumina, San Diego, CA). Library size distributions were validated using an Agilent 2200 TapeStation (Agilent Technologies, Santa Clara, CA). Additional library quality control, blending of pooled indexed libraries, and cluster optimization was performed using Invitrogen's Qubit 2.0 Fluorometer (Invitrogen, Carlsbad, CA). RNA-seq libraries were

^b Strain from M. Johnston's laboratory.

^c Strains from an ordered deletion collection labeled with CB followed by the plate number and location.

pooled (21-plex) and clustered onto a flow cell lane using an Illumina cBot. Sequencing was performed using an Illumina HiSeq 2500 in Rapid Mode employing a paired-end, 50-base read length (PE50) sequencing strategy.

Image analysis and base calling were performed using Illumina's Real Time Analysis v1.18 software, followed by "demultiplexing" of indexed reads and generation of FASTQ files, using Illumina's bcl2fastq Conversion Software v1.8.2 (http://support.illumina.com/downloads/bcl2fastq_conversion_software_184.html). For analysis of the RNA-seq data, reads of low quality were filtered out prior to alignment to the reference genome (*S. cerevisiae* assembly R64-1-1, Ensembl release 75) using TopHat v2.0.9 (Trapnell et al. 2009). Counts were generated from TopHat alignments for each gene using the Python package HTSeq v0.6.1 (Anders et al. 2015). Genes with low counts across all samples were removed.

For comparisons of mutant to wild-type colony samples, differentially expressed genes were identified using the Bioconductor package DESeq2 (Love et al. 2014) with the apeglm package to estimate t-prior shrinkage (Zhu et al. 2018). We employed the IHW package to weight hypotheses and optimize power (Ignatiadis et al. 2016). Differential expression was defined as $|\log_2(\text{ratio})| \ge 0.585 \ (\pm 1.5 \text{ fold})$ with P < 0.01. Genes were categorized as upregulated [logFC] (fold change) > 0.585, P-value < 0.01], down-regulated (logFC < -0.585, P-value < 0.01), or insignificant (logFC < -0.585 or logFC > 0.585, P-value > 0.01). Differentially expressed pathway-specific genes were visualized in principal component analysis (PCA), volcano plot, and Venn diagram figures using ggplot2. Pathway-specific gene expression was visualized using the Kyoto Encyclopedia of Genes and Genomes (Kanehisa and Goto 2000), and the Bioconductor package pathview (Luo and Brouwer 2013). Classification of targets was based on gene ontology (GO) terms and descriptions in the Saccharomyces Genome Database (http:// www.yeastgenome.org). For all other comparisons, edgeR v3.18.1 (Robinson et al. 2010) was used. A false discovery rate (FDR) method was employed to correct for multiple testing (Reiner et al. 2003). Differential expression was defined as $|\log_2(\text{ratio})| \ge 0.585 \ (\pm 1.5 \text{ fold})$ with *P*-value < 0.01.

GO term analysis

GO term analysis (Ashburner *et al.* 2000) was performed using the GO enRIchment analysis and visuaLizAtion tool (GOrilla) (Eden *et al.* 2007, 2009; Mi *et al.* 2017) using the two unranked lists mode. The Gorilla database had been last updated January 12, 2019. The background list was all of the ORFs identified during the analysis. The target list for genes regulated by all five regulators was identified as genes whose $|\log_2 FC| > 0.585$ and *P*-value < 0.01 in all mutant sets. The target list for genes regulated by both RAS and RTG was identified as genes $|\log_2 FC| > 0.585$ and *P*-value < 0.01 in only the $ras2\Delta$ and $rtg3\Delta$ sets. Enriched GO terms were identified by having $P < 10^{-3}$ and FDR < 0.05. Values for

background and target sets, and enriched GO terms, are in Supplemental Material, Table S2.

Quantitative PCR analysis

Differential gene expression was confirmed by quantitative (q) real-time PCR analysis as described previously (Chavel et al. 2014). cDNA libraries from RNA samples were generated using iScript Reverse Transcriptase Supermix (catalog number 1708840; Bio-Rad, Hercules, CA). qPCR was performed using iTaq Universal SYBR Green Supermix (catalog number 1725120; Bio-Rad) on the Bio-Rad CFX384 Real-Time System. Primers were ordered from Sigma ([Sigma Chemical], St. Louis, MO) and are listed in Table S3. FCs in expression were determined by calculating $\Delta\Delta$ Ct (Livak and Schmittgen 2001) using ACT1 mRNA as the housekeeping gene for each sample. RNA was prepared from at least three samples and the average of at least three biological replicates was recorded. Statistical significance was determined by the Student's t-test.

Time-lapse photography

Cells were grown on YEP-Gal semisolid agar media at 22° for 8 days. Photographs were taken using a Nikon D3000 (Nikon, Garden City, NY) digital camera at 30-min intervals using automatic exposure without flash. Graph paper (0.25 cm) was glued to the plate bottom for scaling and later image stabilization; these were cropped out in the final images. Images were imported into ImageJ as an image stack for image stabilization using Kang Li's image stabilizer plug-in (http://www.cs.cmu.edu/~kangli/code/Image_Stabilizer.html). Stabilized image series were saved in video format. Kymographs were generated using the reslice tool in ImageJ (https://imagej.nih.gov/ij/).

Ruffles were identified as light bands flanked by dark bands. The number of ruffles per time point was counted manually at 1-day intervals for each kymograph. Statistical analysis was carried out for each time point using an unpaired Student's *t*-test.

Evaluation of phosphorylated Kss1 levels

Samples were harvested at specific distances from the colony perimeter at t = 0. To determine P-Kss1p levels at the growing edge of a colony, the outermost millimeters of representative colonies were harvested at 1, 2, 3, and 4 days. To determine the changes in P-Kss1p levels before and after the colony ruffles, samples were harvested from regions of the colony 2 mm from the starting colony edge from representative colonies before and after ruffling.

Samples were evaluated by SDS-PAGE analysis. Immunoblots were performed as described (Cullen 2015a). P-Kss1p was detected with phospho-p44/42 primary antibodies (catalog number 4370S; Cell Signaling Technology, Danvers, MA) and anti-rabbit HRP secondary antibodies (catalog number 111-035-144; Jackson ImmunoResearch, West Grove, PA). Total Kss1p was detected by anti-Kss1p antibodies (SC-6775-R; Santa Cruz Biotechnology, Dallas, TX) and with

anti-rabbit HRP secondary antibodies (catalog number 111-035-144; Jackson ImmunoResearch). Loading control Pgk1p was identified by anti-Pgk1p primary antibodies (catalog number 459250; Invitrogen) and with anti-mouse HRP secondary antibodies (catalog number 1706516; Bio-Rad).

Effect of S. cerevisiae cell-cell adhesion on pharyngeal uptake by C. elegans

Wild-type yeast cells expressing GFP (PC6733) were grown in synthetic or YEPD liquid media at 30° for 16 hr. Cells were washed twice in M9 buffer (KH₂PO₄, 3 g/liter; Na₂HPO₄, 6 g/liter; NaCl, 5 g/liter; and 1 mM MgSO₄). As previously described (Bois *et al.* 2013), adult *C. elegans* were transferred into suspensions of *S. cerevisiae*. After 45 min, *C. elegans* were removed and mounted onto 2% agarose pads, and immobilized with 10 mM sodium azide. Slides were examined with an Axioplan 2 fluorescent microscope (Zeiss) at 40×. Individual *S. cerevisiae* cells were counted. Independent replicates were performed on three separate days.

Experiments involving yeast colonies exposed to entry by C. elegans

Yeast cells were grown in YEPD liquid media at 30° for 16 hr. For interactions between C. elegans and S. cerevisiae colonies, 200- μ l aliquots of cells were dispensed onto NGM (Wood 1988) agar media and grown as colonies at 22° for 72 hr. Under this condition, yeast colonies formed Flo11p-dependent patterns. OP50 E. coli (Brenner 1974) were taken from stock cultures, and 200 μ l aliquots were dispensed onto NGM agar media and grown at 22° for 3 days.

To examine colony penetration, adult and fourth-stage larva (L4) *C. elegans* were transferred from stock OP50 plates to experimental plates around the yeast or *E. coli*. Mat penetration times were determined by measuring the time from a worm nose first contacting the colony to the tail fully entering the mat, up to 100 sec. The number of stalls and reversals was determined by counting the number of incidents when a worm's tail would stop forward movement or reverse. The percent of time moving forward was determined by recording the amount of time that a worm was moving forward divided by the total time required to penetrate a mat. Independent replicates were performed on at least three separate days.

To examine the pharyngeal uptake of *S. cerevisiae* by *C. elegans*, wild-type (PC538) and $flo11\Delta$ (PC1029) yeast were transformed with a plasmid containing GFP-2xPH (PC2560) [CS189 (Stefan *et al.* 2005) provided by the Emr laboratory (Cornell University)]. Cells were grown in YEPD liquid media at 30° for 16 hr and 200- μ l aliquots were spotted onto NGM agar media. Colonies were grown at 22° for 3 days. Worms were transferred directly into colonies and left for 45 min before mounting and imaging. Independent replicates were performed on three separate days.

Videos were captured using an AmScope MD35 camera and AmScope image capture software. The camera was inserted into the eyepiece of a Zeiss SteREO Discovery V8. Dot plots for worm entry times were generated in ggplot2 (Wickham 2016). Statistical analysis was performed using the Student's *t*-test.

Infrared imaging

Infrared images were taken using FLIR A325sc (FLIR Systems, Wilsonville, OR) and captured using FLIR ResearchIRMax4 (FLIR Systems), provided by the Sustainable Manufacturing And Robotic Technology center at the University at Buffalo. Average and coolest temperatures for each colony were measured using FLIR ResearchIRMax4 (FLIR Systems).

An insulated housing unit was built to mount the thermal imaging camera above a stage for agar plates. Yeast cells were grown in YEPD liquid media at 30° for 16 hr. Next, 10- μ l aliquots were spotted onto YEP-Gal agar media. Colonies were grown for 72 hr at 30°. For imaging, plates were transferred from the incubator to the mount in a photographing area with an ambient temperature of 22°. Plate lids were removed to allow imaging. Images were taken immediately after removing the lid.

Data availability

All strains are available upon request. The Gene Expression Omnibus (GEO) accession number for raw sequencing data is GSE115657. A comparison with previously published expression profiling data sets for cells grown in liquid culture has been described [GEO accession number GSE61783 (Adhikari and Cullen 2014)]. Supplemental material available at FigShare: https://doi.org/10.25386/genetics.8066615.

Results

A signaling network regulates adhesion-dependent surface responses in yeast

Multiple signal transduction pathways regulate filamentous growth (Figure 1A) (Chavel *et al.* 2010, 2014). A large number of genes that control invasive growth, pseudohyphal formation, and mat formation show significant overlap based on a genome-wide analysis using deletion mutants in the filamentous strain background (Ryan *et al.* 2012). Thus, we examined and compared how the signaling network that regulates filamentous growth impacts adhesion-dependent surface responses in yeast. We sought to directly compare key pathways in the network (MAPK and RAS) to less well-characterized pathways (RTG) and to a newly identified chromatin remodeling complex (SAGA). All of the pathways were studied in reference to a key target adhesion molecule, Flo11p.

Some strains (e.g., Σ 1278b) exhibit ruffled colony morphology when grown on surfaces. This phenotype is dependent on cell adhesion contacts mediated by Flo11p (Reynolds and Fink 2001; Karunanithi *et al.* 2012). To evaluate colony pattern formation, we performed a time-course experiment followed by kymograph analysis, which can reveal changes in local features over time (Kaksonen *et al.* 2003). Wild-type cells were spotted on 2% agar media (YEP-Gal), and ruffle

formation was examined by photographing colonies at 30-min intervals over an 8-day period (Video S1). In a series of experiments, kymographs were generated along multiple radii of expanding colonies to measure the extent of ruffle formation [Figure 1B, wild-type (PC538), yellow line corresponds to one kymograph at right]. Ruffles were identified as light horizontal bands flanked by dark bands and could be counted manually by this method (Figure 1B, wild-type, black arrows). Measuring the banding pattern from separate sections of the colony allowed us to assess the extent of ruffle formation over time (Figure 1C, wild-type). As has been previously demonstrated (Granek and Magwene 2010), colonies lacking the adhesion molecule Flo11p did not form ruffles (Video S2), which was also evident by kymograph analysis [Figure 1, B and C and Figure S1, $flo11\Delta$ (PC1029)]. Therefore, kymograph analysis permitted the numerical assessment of pattern formation in yeast colonies. In principle, kymograph analysis may allow quantification of colony pattern formation in other microbes.

Kymograph analysis was next applied to mutants lacking key components of the major signaling pathways that regulate filamentous growth. As expected (Roberts and Fink 1994; Rupp et al. 1999; Reynolds and Fink 2001; Granek and Magwene 2010), the fMAPK pathway was required for colony ruffling. Specifically, kymograph analysis showed that colonies lacking a negative regulator of the fMAPK pathway, Dig1p (Cook et al. 1996; Tedford et al. 1997; Bardwell et al. 1998; Olson et al. 2000; Breitkreutz et al. 2003; Kusari et al. 2004; Chou et al. 2006; van der Felden et al. 2014), formed more ruffles than wild-type after 1 day of growth [Figure 1, B and C, $dig1\Delta$ (PC3039), Figure S1, and Video S3], though this difference was less apparent over longer time periods. Conversely, loss of the fMAPK transcription factor, Ste12p, led to a defect in colony ruffling at the examined time points [Figure 1, B and C, $ste12\Delta$ (PC1079), Figure S1, and Video S4]. The fMAPK pathway was also required for mat formation on 0.3% agar (Figure S1; $ste12\Delta$ and $dig1\Delta$; Figure S2 provides examples of other fMAPK regulators; $ste11\Delta$ and $ste20\Delta$). Also as expected (Gimeno et al. 1992; Rupp et al. 1999; Granek and Magwene 2010; Zara et al. 2011; Ryan et al. 2012), the RAS pathway was required for colony ruffling [Figure 1, B and C, ras2Δ (PC562) and Video S5] and mat formation (Figures S1 and S2). Thus, kymograph analysis can be used to evaluate the roles of signaling pathways in regulating aspects of mat growth and colony pattern formation.

We next tested whether other pathways that regulate filamentous growth also regulated adhesion-dependent colonial responses. The mitochondria-to-nucleus RTG pathway regulates invasive growth (Chavel *et al.* 2010; González *et al.* 2017). We found that the RTG pathway was also required for colony ruffling [Figure 1, B and C, $rtg3\Delta$ (PC3642), Figure S3, and Video S6] and mat formation on 0.3% agar (Figures S1 and S2).

We next tested the role of other proteins that control filamentous growth. Pho85p regulates filamentous growth

(Chavel *et al.* 2010) and was also required for mat formation (Figure S1 and S2). Rpd3p (Chavel *et al.* 2010; Voordeckers *et al.* 2012), lipid regulators Opi1p and Nte1p (Chavel *et al.* 2014), and Snf1p (Cullen and Sprague 2000; Voordeckers *et al.* 2012) were also required for mat formation (Figures S1 and S2). Based on these results, we conclude that many of the major signaling pathways that regulate filamentous growth also regulate mat formation and colony patterning.

We hypothesized that the chromatin remodeling complex SAGA (Koutelou et al. 2010) might also regulate adhesiondependent surface growth. The hypothesis was based on the fact that a component of SAGA, Gcn5p (Georgakopoulos and Thireos 1992; Sterner and Berger 2000), is required for filamentous growth (Chavel et al. 2014). Moreover, SAGA components have previously been shown to control aspects of colonial patterning (Voordeckers et al. 2012). Spt8p, a SAGA component (Winston et al. 1987), also regulates filamentous growth. Kymograph analysis of the $spt8\Delta$ mutant showed a defect in ruffle formation at 2 and 3 days [Figure 1, B and C, spt8 Δ (PC4008), Figure S3, and Video S7]. The defect was subtle by kymograph analysis but obvious in the time-lapse analysis. The $spt8\Delta$ mutant was also defective for colony ruffling and mat pattern formation (Figure S1). Therefore, we include SAGA as a regulator of filamentous surface responses in yeast (Figure 1A, cyan).

Expression profiling of colony surface growth identifies new targets

To explore how filamentation signaling pathways might regulate surface growth, comparative RNA-seq was performed in several mutants that disrupt the main pathways that regulate filamentous growth (dashed blue box in Figure 1A, pathway diagrams can be found in Figure S4, A–D). RNA was prepared from wild-type and mutant colonies with the following genotypes: $ste12\Delta$ (fMAPK), $dig1\Delta$ (fMAPK), $ras2\Delta$ (RAS), $rtg3\Delta$ (RTG), and $spt8\Delta$ (SAGA). RNA was prepared from colonies under conditions that favored pattern formation (YEP-Gal). Each mutant showed the expected colony patterning. Specifically, the $ras2\Delta$ (PC562), $rtg3\Delta$ (PC3642), $ste12\Delta$ (PC1079), and $spt8\Delta$ (PC4008) mutants were less ruffled than wild-type colonies (PC538), and the $dig1\Delta$ (PC3039) mutant was more ruffled than wild-type colonies (Figure S4E). After sequencing, PCA of the RNA-seq data showed close clustering of strain replicates, while strains differentiated into their replicate clusters (Table S1).

Comparative RNA-seq analysis between the wild-type colonies and each mutant was performed with the DESeq2 package in R (Love *et al.* 2014). Differential gene expression was defined by $|\log_2 FC| > 0.585$ and P-value < 0.01. By this method, 1833 genes were differentially expressed in at least one of the mutants tested (Figure S4F) and represented 29% of the ORFs in the genome (Lin *et al.* 2013). The annotated data set can be found in Table S1. To identify the most differentially regulated targets between the mutant and wild-type colonies, volcano plots showing all differences in gene expression were generated for each mutant. Individual genes

were distributed by change in expression (x-axis, log₂FC) and significance of change in expression [y-axis, $-\log_{10}(P$ value)] (Figure 2A). Many of the genes whose expression is known to be induced during filamentous growth were uncovered during the initial analyses (Figure 2A and Table S1). These included targets of the fMAPK pathway: FLO11 (Rupp et al. 1999) (Figure 2A, $dig1\Delta$ and $ste12\Delta$; Table S1, $dig1\Delta$ and $ste12\Delta$), YLR042C (Roberts et al. 2000) (Figure 2A, $ste12\Delta$; Table S1, $dig1\Delta$ and $ste12\Delta$), CLN1 (Madhani et al. 1999) (Table S1, $dig1\Delta$ and $ste12\Delta$), PGU1 (Madhani et al. 1999; Roberts et al. 2000) (Figure 2A, $dig1\Delta$ and ste12 Δ ; Table S1, dig1 Δ and ste12 Δ), SVS1 (Roberts et al. 2000) (Figure 2A, $dig1\Delta$ and $ste12\Delta$; Table S1, $dig1\Delta$ and ste12 Δ), KSS1 (Table S1, dig1 Δ and ste12 Δ) (Roberts et al. 2000), and MSB2 (Cullen et al. 2004) (Table S1, $dig1\Delta$ and $ste12\Delta$). The abovementioned fMAPK pathway targets YLR042C, PGU1 (Roberts et al. 2000), FLO11 (Rupp et al. 1999) are also Ras2p-dependent (Table S1, $ras2\Delta$), potentially through its regulation of the fMAPK pathway (Mösch et al. 1996, 1999; Chavel et al. 2010). Similarly, targets of the RTG pathway were identified in the wild-type- $rtg3\Delta$ data set: CIT2, CIT1, IDH1, IDH2 (Liu and Butow 1999) (Figure 2A, $rtg3\Delta$; Table S1, $rtg3\Delta$), and *DLD3* (Liu and Butow 2006) (Table S1, $rtg3\Delta$). Genes regulated by SAGA were identified in the wild-type-spt8 Δ data set [ADH1, ARG1, BDF2, CTT1, FBA1, GRE2, PGK1, TDH3, and PHO84 (Basehoar et al. 2004; Huisinga and Pugh 2004)] (Table S1, $spt8\Delta$).

Confirmation of other targets by qPCR analysis has been performed in related studies including: targets of fMAPK related to the fungal cell wall, *OCH1*, *PRY2*, *FLO10*, and *TIP1* (Chow *et al.* 2018); *SUC2* and *YLR042C*; and *GIC2* (Aditi Prabhakar, personal communication). Therefore, comparative RNA-seq analysis identified many genes expected based on previous or parallel studies.

One question we sought to address was the extent of target gene overlap among the signaling pathways that regulate filamentous growth. We found that while most genes were regulated by one pathway (Figure S4G, 742/1833 or 59%), coregulated targets were seen in almost every combination of regulators, including targets coregulated by all five regulators (Figure S4F). This result also demonstrated that the SAGA complex coregulated targets with the fMAPK, RAS, and RTG pathways (Figure S4F), which further validates it as a major regulator of filamentous growth (Figure 1A, cyan arrows). Additionally, Spt8p regulated a target of the RIM101 pathway, NRG1 (Chavel et al. 2014) (Figure 1A, cyan arrow), and a target of lipid biosynthesis, INO1 (Chavel et al. 2014) (Figure 1A, cyan arrow). SAGA did not coregulate targets of the PHO85, RPD3(L), or ELP pathways. Taken together, these data suggest that one function of this network of signaling pathways is to regulate a large number of genes (the more nodes in the network, the greater the number of differentially expressed genes); a second function is to coregulate key targets to amplify target gene expression.

One possible explanation for the regulatory overlap is cross feedback among the pathways. To explore this aspect of signaling network connectivity, we examined whether genes encoding pathway components were themselves targets of other pathways that regulate filamentous growth. The RTG pathway regulated the expression of components of fMAPK (MSB2), RAS (TPK2), and SAGA (SFG73). fMAPK regulated genes encoding components of its own pathway (MSB2, TEC1, and KSS1) and the RAS pathway (BCY1). The RAS pathway regulated one of its own effectors (TPK1) and the fMAPK pathway (TEC1). SAGA regulated the fMAPK pathway (MSB2 and TEC1) and itself (SUS1) (Figure S5, A and B). These results support and extend a previous study, which showed that the activity of the fMAPK is subject to regulation by other filamentation regulatory pathways (Chavel et al. 2010). Interestingly, not all of the interactions would be expected to result in positive feedback. For example, the $rtg3\Delta$ mutant showed a \sim 1.9-fold increase in TPK2 gene expression (Table S1), which is a PKA subunit and component of the RAS pathway. These data indicate that coregulation of targets may be the result of feedback among the pathways that regulate filamentous growth.

To identify functionally relevant targets whose expression was amplified by multiple pathways that regulate filamentous growth, we focused on genes that were differentially expressed in all regulatory mutants (Figure 2B and Table S2, all mutants). GO term analysis (Ashburner et al. 2000) of the coregulated genes showed significant enrichment in cell adhesion, cell wall constituents, and fungal cell wall proteins (Table S2, all mutants). One of the targets that was coregulated by all five regulators, and that was included in many of the above enriched GO terms, was the gene that encodes the major cell adhesion molecule Flo11p (Figure 2B and Table S2, all mutants). FLO11 is known to be regulated by a large number of proteins and pathways (Rupp et al. 1999; Barrales et al. 2008). We confirmed that fMAPK, RAS, RTG, and Spt8p controlled FLO11 expression by qPCR (Figure 2C). These data fit with the key role that Flo11p plays in regulating adhesion-dependent responses in yeast. Another target regulated by all five mutants, which also had the largest net change in regulation, was PGU1, the gene that encodes a secreted pectinase that is also expressed during filamentous growth (Madhani et al. 1999; Cullen 2015a). This result was confirmed by qPCR analysis (Figure 2D). Taken together, our results are consistent with the idea that multiple signaling pathways and protein complexes coordinately regulate target gene expression during colonial surface growth (Figure 2A). One function of the network is to control the expression of FLO11, a major regulator of adhesiondependent responses in yeast (Figure 2, B and C).

A regulatory connection between the RAS and RTG pathways

In an analysis of highly significant changes in expression in the $ras2\Delta$ and $rtg3\Delta$ sets ($P < 10^{-12}$) (Figure S6), 41 targets were in common and 93% of these targets (39/41) showed the same regulation (both up or down) (Figure S6). For example, citrate synthase, *CIT3*, was among the significant,

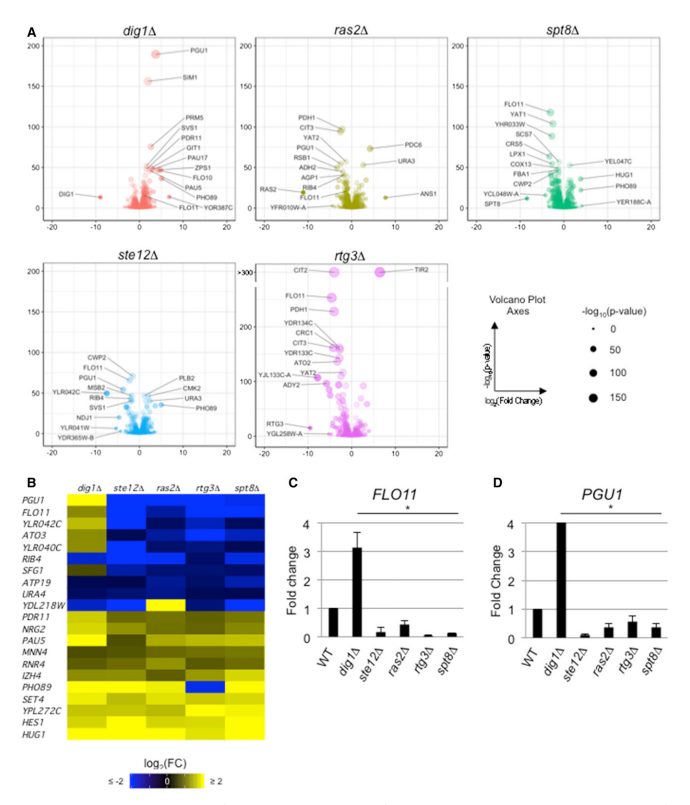


Figure 2 Comparative RNA-seq analysis identifies common and unique targets of signaling pathways that regulate - adhesion-dependent surface growth growth. (A) Volcano plots showing comparative RNA-seq between the indicated mutant and WT. Dot size is $-\log_{10}(P\text{-value})$. Volcano plot *x*-axis is $\log_2(FC)$; *y*-axis is $-\log_{10}(P\text{-value})$. Labeled dots are 10 targets with lowest *P*-value or five targets with highest Ifold changel. *FLO11* has also been labeled on $dig1\Delta$ and $ras2\Delta$ plots. *CIT2* and *TIR2* have calculated *P*-values < 10^{-300} in the $rtg3\Delta$ mutant, as reflected in the break in the *y*-axis. (B) Heat map showing targets with $\log_2(FC)$ > 0.585 and *P*-value < 10^{-3} . (C) Bar graph showing fold change in *FLO11* mRNA levels, normalized to *ACT1* with WT values set to 1, in the indicated mutants by qPCR analysis by the $\Delta\Delta$ Ct quantitation method. The experiment was performed in triplicate and error bars represent the SD between experiments. * P < 0.05 for all differences compared to WT. (D) Bar graph showing fold change in *PGU1* mRNA levels. See (C) for details. $\log_2(FC)$, $\log_2(Fold Change)$; qPCR, quantitative PCR; RNA-seq, RNA sequencing; WT, wild-type.

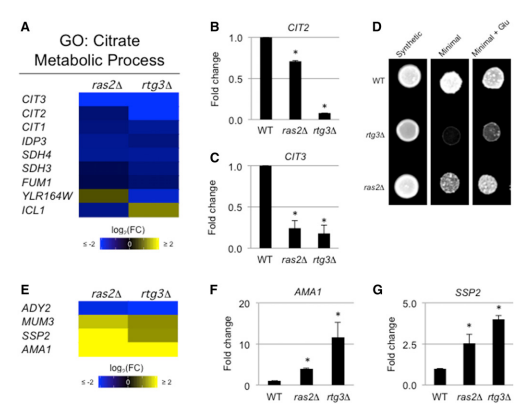


Figure 3 Common and unique target genes of the RAS and RTG pathways. (A) Heat map of targets coregulated by the RAS and RTG pathways with the citrate metabolic process GO term. (B and C) Bar graph showing fold change in (B) CIT2 and (C) CIT3 mRNAs in ras2\Delta and $rtg3\Delta$ relative to WT. See Figure 2C for details. (D) Strains spotted onto synthetic media, minimal media, or minimal media with glutamate. Cells were grown for 48 hr. (E) Heat map of targets coregulated by RAS and RTG associated with sporulation. (F and G) Bar graphs showing fold change in (F) AMA1 and (G) SSP2 mRNAs in $ras2\Delta$ and $rtg3\Delta$ relative to WT. See Figure 2C for details. Glu, glutamate; GO, gene ontology; log₂(FC), log₂(Fold Change); WT, wild-type.

differentially expressed genes in both $ras2\Delta$ and $rtg3\Delta$ (Figure 2A). Other citrate synthases, CIT1 and CIT2, were also coregulated by both the $ras2\Delta$ and $rtg3\Delta$ mutants (Figure 3A and Table S1). We confirmed that RAS and RTG coregulated CIT2 and CIT3 by qPCR analysis (Figure 3, B and C). Because of the extent of coregulation, especially the coregulation of all citrate synthase isoforms (Graybill et al. 2007), the full set of RAS and RTG coregulated targets was examined further.

As expected, the RTG pathway regulated mitochondrial gene targets (Table S1). Many of the genes were also regulated by the RAS pathway (Table S1; PDH1, YAT1, and YAT2) (Schmalix and Bandlow 1993; Epstein et al. 2001; Swiegers et al. 2001). Moreover, several hallmark RTG targets were also regulated by the RAS pathway (Table S1; CIT1, CIT2, CIT3, ATO2, and ATO3) (Suissa et al. 1984; Kim et al. 1986). In addition to coregulation of cell wall components (i.e., FLO11), GO term analysis identified significant enrichment of the citrate metabolic process by the overlapping RAS and RTG targets (Figure 3A and Table S2), and the citric acid cycle (Figure S7). Because RAS and RTG coregulated genes that are involved in the citric acid cycle, we hypothesized that RAS might be required to support that mitochondrial role. The mitochondria are critical for glutamate biosynthesis (Liu and Butow 1999; Magasanik and Kaiser 2002) and RTG is required for growth in medium lacking glutamate [Figure 3D, $rtg3\Delta$ (Liao and Butow 1993; Small et al. 1995; Chavel et al. 2014)]. More specifically, a loss of citrate synthases results in glutamate auxotrophy in S. cerevisiae (Kim et al. 1986), so we hypothesized that a RAS pathway mutant would be sensitive to limiting glutamate. Like RTG, the RAS pathway was also

required for growth in this condition (Figure 3D, $ras2\Delta$). The growth defects of cells lacking an intact RAS or RTG pathway were bypassed by the addition of glutamate (Figure 3D). Therefore, the RAS pathway plays a role in this mitochondrial process.

Similarly, the RAS pathway regulated genes that are required for sporulation (Table S1) and a subset of these genes were coregulated by the *RTG* pathway [Figure 3E, *ADY2* (Rabitsch *et al.* 2001), *AMA1* (Coluccio *et al.* 2004), *SSP2* (Sarkar *et al.* 2002), and *MUM3* (Engebrecht *et al.* 1998)]. Several of the genes were confirmed by qPCR analysis (Figure 3, F and G). Therefore, the RAS and RTG pathways coregulate functionally related genes and may be more functionally connected than had been previously thought.

Nonreciprocal target genes are regulated by opposing transcriptional regulators of the fMAPK pathway

Ste12p is one of the transcription factors that regulates the fMAPK pathway (Liu *et al.* 1993). The transcriptional repressor Dig1p inhibits Ste12p-dependent transcription (Cook *et al.* 1996; Tedford *et al.* 1997; Bardwell *et al.* 1998; Olson *et al.* 2000; Breitkreutz *et al.* 2003; Kusari *et al.* 2004; Chou *et al.* 2006; van der Felden *et al.* 2014). Although the transcriptional outputs of the pathway can be complex resulting in nonuniform target gene regulation (Breitkreutz *et al.* 2003), Ste12p and Dig1p were expected to regulate a common set of targets in a reciprocal manner (Figure 4A). We unexpectedly found that the pattern of gene regulation by the two proteins could not be solely explained by reciprocal regulation. In an analysis of the most significant changes in

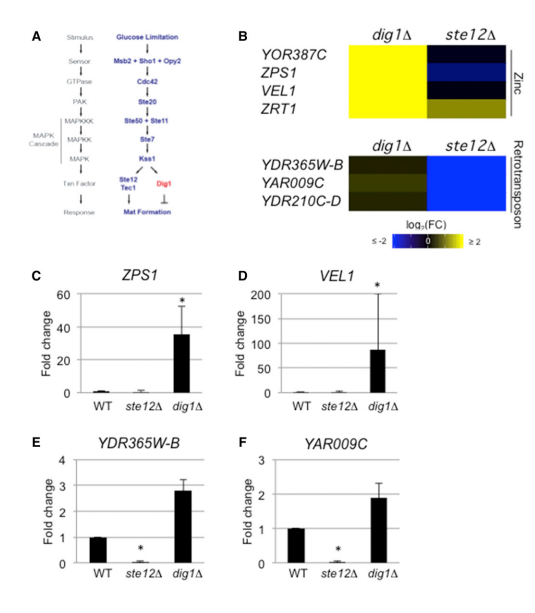


Figure 4 Common and unique genes regulated by Ste12p and Dig1p. (A) Model of the fMAPK pathway showing Ste12p and Dig1p. (B) Heat map of indicated targets regulated in the $dig1\Delta$ or $ste12\Delta$ mutants. (C–F) Bar graphs showing fold change in target mRNA levels for (C) ZPS1, (D) VEL1, (E) YDR365W-B, and (F) YAR009C. See Figure 2C for details. fMAPK, fungal MAPK; $log_2(FC)$, $log_2(Fold Change)$; WT, wild-type.

expression in the $dig1\Delta$ and $ste12\Delta$ sets ($P < 10^{-12}$) (Figure S8), 20 targets were found to be in common and only 50% (10/20) were reciprocally regulated (up in $dig1\Delta$ and down in $ste12\Delta$) (Figure S8). As expected, Ste12p and Dig1p showed reciprocal regulation of a subset of target genes (Figure S8; FLO11/MUC1, PGU1, YLRO42C, and BAR1). However, more genes were independently regulated only by Dig1p or only by Ste12p (Figure S8).

For example, the target with the largest change in expression identified in the $dig1\Delta$ volcano plot was YOR387C (> 111-fold increase) (Figure 2A and Table S1). Although YOR387C lacks associated GO terms, a literature search indicated that YOR387C is induced in zinc-limiting conditions (Higgins et~al.~2003; Wu et~al.~2008). Although YOR387C and its paralog VEL1 (Higgins et~al.~2003) were significantly regulated in the $dig1\Delta$ mutant, neither gene was found to be regulated in the $ste12\Delta$ mutant (Figure 4B and Table S1). Additionally, only the $dig1\Delta$ mutant showed a change in expression of ZAP1 (Table S1), a major transcription factor in-

duced by limiting zinc (Zhao and Eide 1997). Interestingly, the ability of cells to take up trace zinc was determined to be critical for flocculation (Yuan 2000), which is a closely related response to filamentous growth and mat formation. A subset of other upregulated targets in the $dig1\Delta$ mutant were genes induced under low-zinc conditions [Figure 4B, ZPS1 (Lyons et~al.~2000)] as well as a zinc transporter [ZRT1 (Zhao and Eide 1996)]. These genes were not strongly downregulated in the $ste12\Delta$ mutant. The expression profiles of ZPS1 and VEL1 were confirmed by qPCR analysis (Figure 4, C and D).

Likewise, the $ste12\Delta$ mutant showed downregulation of some target genes that were not altered in the $dig1\Delta$ mutant. One of the targets identified in the $ste12\Delta$ volcano plot was the retrotransposon YDR365W-B (Kim et~al.~1998). Other retrotransposons—YDR365W-B, YOL103W-B, YAR009C, and YDR210C-D (Kim et~al.~1998)—were also regulated by Ste12p but less so by Dig1p (Figure 4B). The expression profiles of YDR365W-B and YAR009C were confirmed by qPCR analysis (Figure 4, E and F). Thus, key transcriptional

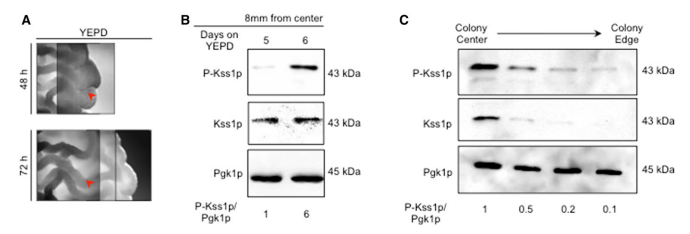


Figure 5 Different ruffling patterns in colonies correspond to different levels of fMAPK pathway activity. (A) Bright-field microscopy of a wild-type colony grown in YEPD for 48 and 72 hr. Red arrow indicates same location on a mat at 48 hr when the region is smooth and 72 hr when the region has become ruffled. Different focal planes of the same colony are shown. (B) Immunoblot analysis using p42/p44 antibodies to detect P-Kss1p levels from extracts prepared from cells before and after ruffling. Immunoblots were also performed using antibodies to Kss1p and Pgk1p as loading controls. (C) Immunoblot analysis using p42/p44 antibodies to detect phosphorylated Kss1p at different parts of a colony. Immunoblots were also performed using antibodies to Kss1p and Pgk1p as loading controls. fMAPK, filamentous growth MAPK; YEPD, yeast extract, peptone, and dextrose.

regulators of the fMAPK pathway operate through mechanisms that do not only involve reciprocal regulation between the two proteins.

Changes in colony patterning correspond to changes in MAPK pathway activity

We also examined the development of colony pattern formation under different conditions and over time. Colonies showed different ruffling patterns under different conditions. For example, growth in preferred carbon sources such as glucose media (YEPD) resulted in colony perimeters that were smooth compared to a ruffled colony interior (Figure 5A, adjacent photographs of a colony are shown at different focal planes on the z-axis). Such differences were not observed in more uniformly ruffled colonies grown on a nonpreferred carbon source (YEP-Gal) (Video S1). Over time, the smooth regions of the colony became ruffled (Figure 5A, red arrow marks the same location in the colony).

One explanation for differences in colony pattern formation might be that different parts of the colony are exposed to different environments. The fMAPK pathway, which is sensitive to nutrient levels, may induce different responses in different parts of the colony. To test this possibility, samples from regions of the colony were collected and examined for fMAPK activity by phosphorylation of the MAP kinase Kss1p (Cullen 2015a). Samples collected from smooth colony perimeters showed low P~Kss1p levels (day 5, smooth). Samples collected from the same physical location that had become ruffled (day 6, ruffled) showed high P~Kss1p levels (Figure 5B). This result indicates that as cells in colonies consume nutrients and subsequently starve, the fMAPK pathway is activated to promote ruffle formation. In line with this possibility, colony interiors showed higher levels of P~Kss1p compared with colony perimeters (Figure 5C). We note that in this experiment, the levels of total Kss1p protein were also

lower at colony perimeters. *KSS1* is a target of the fMAPK pathway, and the regulation of total Kss1p may be a way that fMAPK regulates its activity through positive feedback (Roberts *et al.* 2000). Therefore, signaling pathways may exhibit different activities, and induce target genes to different levels, in a manner that is influenced by nutrient levels (and time) to control adhesion-dependent surface growth.

Genes that respond to oxygen, desiccation, and temperature stress were induced during colony surface growth

Growth on surfaces presents unique challenges compared to the uniform growth in liquid cultures. To further explore the response to colonial growth on surfaces, expression profiling data sets were compared between cells grown in liquid (Adhikari and Cullen 2014) and cells grown on an agar surface (this study). Growth on surfaces caused the differential expression of 3267 genes when compared to growth in liquid (Table S1, wild-type liquid-solid, $|\log_2FC| > 0.585$ and FDR < 0.05). This represents \sim 54% of the yeast genome [6091 genes (Lin et al. 2013)]. Functional classification of the top 15 induced targets (Figure 6A, >25-fold increase in expression) during surface growth included genes induced during anaerobic stress [DAN1 (Sertil et al. 1997) and TIR1 (Cohen et al. 2001), Figure 6A], desiccation/dehydration [SIP18 (Miralles and Serrano 1995) and GRE1 (Garay-Arroyo and Covarrubias 1999), Figure 6A], and elevated temperature [HSP33 (Wilson et al. 2004) and SNO4 (Samanta and Liang 2003), Figure 6A]. An expanded list of the top 50 (>15-fold increase in expression) induced targets included genes that regulate metal homeostasis, RNA processing, and the metabolism of lipids and nonfermentable carbon sources, as well as genes of unknown function (Figure S9).

The most differentially induced target during surface growth was the gene that encodes the anaerobic-responsive

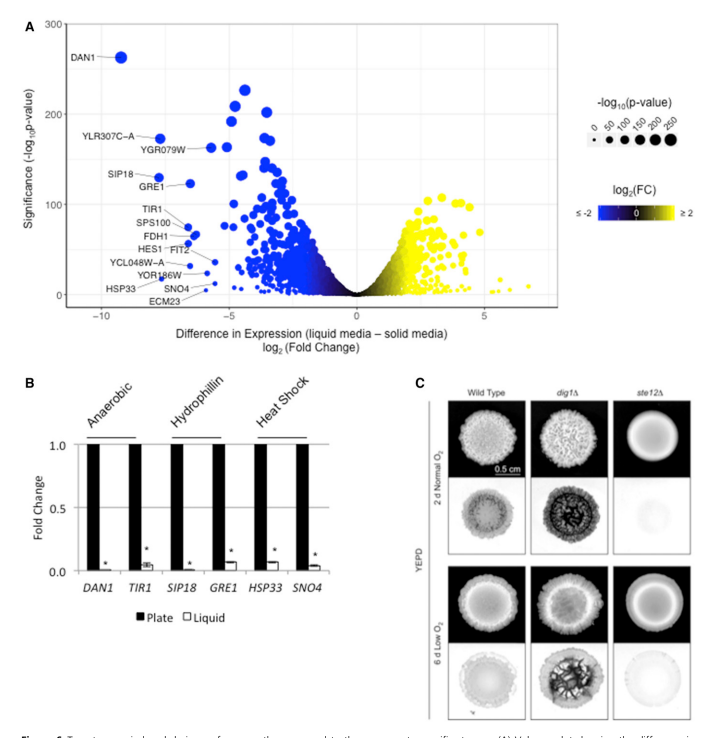


Figure 6 Target genes induced during surface growth correspond to the response to specific stresses. (A) Volcano plot showing the difference in expression and significance of all targets between growth in liquid media vs. solid media. x-axis is $log_2(Fold Change)$; y-axis is $-log_{10}(P$ -value). Labeled targets are the top 15 induced targets during surface growth. (B) Relative fold change of the indicated gene was determined by qPCR analysis. Expression was normalized to ACT1 in the indicated mutants, as determined by qPCR analysis. The experiment was performed in triplicate and error bars represent the standard difference between experiments. * P < 0.05 for differences compared to wild-type. (C) Wild-type cells and the indicated mutants were spotted on YEPD 2% agar media in normal and low-oxygen conditions. Colonies were grown for the indicated times to compare colonies of comparable sizes before photographing (top panels), washing, and rephotographing colonies (bottom panels) to determine colony patterns and invasive growth. Inverted images of the invasive scars are shown. $log_2(FC)$, $log_2(Fold Change)$; qPCR, quantitative PCR; YEPD, yeast extract, peptone, and dextrose.

cell wall mannoprotein Dan1p (Mrsa et al. 1999) (Figure 6A). qPCR analysis showed that DAN1 expression was induced during surface growth (Figure 6B). Examination of the $dan1\Delta$ mutant from an ordered gene deletion collection (Ryan et al. 2012) showed a defect in colony ruffling (Figure S10, $dan1\Delta$). Thus, Dan1p may impact colony patterning in response to stresses associated with surface growth. In addition to DAN1, the four members of the Tir family of anaerobic-responsive cell wall mannoproteins (Cohen et al. 2001) also showed transcriptional induction during surface growth (Figure 6A and Table S1, wild-type liquid–solid), which was verified in a study on the regulation of the cell wall (Chow et al. 2018). Therefore, anaerobic-specific changes to the cell wall also occur during surface growth in yeast.

Because targets that encode anaerobic-responsive proteins were among the top induced targets during colony surface growth [DAN1 and the TIR genes (Figure 6A)], the impact of oxygen in colony ruffling and invasive growth was examined. S. cerevisiae undergoes aerobic alcohol fermentation by what is known as the Crabtree effect (Crabtree 1929). In yeast, this includes the repression of genes involved in aerobic respiration in the presence of glucose during exponential growth (De Deken 1966). Studies that explore this phenomenon in yeast have done so in liquid culture (De Deken 1966; Hagman et al. 2014), but the role of oxygen in regulating aspects of colonial growth has not been explored. To determine the role of oxygen on colony pattern formation, wildtype and $dig1\Delta$ colonies grown in normal atmospheric oxygen (20% oxygen) were compared to colonies grown in low oxygen (5–15% oxygen). Comparing the colonies at similar sizes (2-day normal oxygen and 6-day low oxygen) showed that both were less ruffled in low oxygen (Figure 6C). Wildtype and $dig1\Delta$ colonies showed reduced invasive growth in low oxygen (Figure 6C). Additionally, wild-type, $dig1\Delta$, and $ste12\Delta$ colonies grown in limiting oxygen (5–15% oxygen or 0% oxygen) grew more slowly than in normal oxygen (20% oxygen) (Figure S11). Slower growth was not bypassed with excess glucose (Figure S11, 8% Glu) as might be predicted by the Crabtree effect in yeast. Therefore, oxygen levels impact colony patterning and invasive growth.

Another class of genes induced during surface growth has been described as regulated during desiccation. These included the gene that encodes the hydrophilin Sip18p (Miralles and Serrano 1995; Dang and Hincha 2011) (Figure 6A). A second hydrophilin, Gre1p, (Garay-Arroyo and Covarrubias 1999) was also among the top 10 induced targets (Figure 6A). Four of six hydrophilins and putative hydrophilins were induced during surface growth (Table S1, wild-type liquid-solid). Hydrophilins such as Sip18p allow for survival under desiccation stress and during the dehydration/rehydration process (Rodríguez-Porrata et al. 2012). The induction of hydrophilin-encoding genes SIP18 and GRE1 during surface growth was confirmed by qPCR analysis (Figure 6B). By testing mutants from an ordered deletion collection (Ryan et al. 2012), we found that the loss of either hydrophilin, SIP18 or GRE1 (Figure S10A; $gre1\Delta$ and $sip18\Delta$), increased mat ruffling and hyper-invasive growth, indicating that intracellular hydration may be a trigger for both invasive growth and colony ruffling (Figure S10B; $gre1\Delta$ and $sip18\Delta$). Therefore, cells in mats may experience desiccation stress, and genes induced under this condition can impact colony patterning.

Genes encoding the heat-shock proteins Hsp33p (fourth most-induced target), Sno4p, Hsp26p, and Hsp12p were also induced during mat growth. Additionally, as shown below, a ruffled colony showed a temperature differential of 0.8° between its warmest and coolest regions (see Figure 7D, below). Modest differences in temperature have previously been shown to effect metabolism in *S. cerevisiae* (Jones and Hough 1970). Spg4p, described as being essential for growth at high temperature (Martinez *et al.* 2004), was also among the top targets. Using an ordered deletion collection, we explored the roles of SNO4 and HSP26 (Bentley *et al.* 1992) (Figure S10A; $sno4\Delta$ and $hsp26\Delta$), as the $hsp33\Delta$ mutant was not present in the deletion collection. Deletion of either of these genes resulted in smoother colonies.

None of the mutants tested were necessary for growth in low-oxygen (5–15% $\rm O_2$), high-temperature (37°), or desicated conditions (Figure S11). This may be a result of functional redundancy among the major induced targets. Taken together, oxygen, temperature, and desiccation stresses impact colonial growth in yeast. Genes induced during surface growth impact colony patterning that may aid in the response to these stresses.

Flo11p-dependent adhesion protects cells in colonies from a nematode predator

Flo11p is an established regulator of filamentous growth, and its role in maintaining contacts between cells to form filaments has obvious benefits during surface penetration (e.g., invasive growth) into new environments. However, the broader functional implications of Flo11p-dependent adhesion between cells on the colony surface are not clear. The interactions among yeast cells mediated by a related flocculin, Flo1p, can provide protection from toxins and antibiotics (Smukalla et al. 2008), and cell adhesion has more generally been postulated to provide protection to individual cells of a colony (Granek and Magwene 2010). In bacterial communities, the growth of cells in mats can deter predation by other microbes and multicellular predators, like nematodes (Darby et al. 2002). To explore whether cell adhesion contacts might similarly protect fungal cells from predation, we developed a mock predator-prey assay between S. cerevisiae and the nematode C. elegans.

C. elegans is a free-living nematode that is commonly used in research laboratories (Brenner 1974), and has been used to study the effects of *C. albicans* and *S. cerevisiae* accumulation in the gut (Jain *et al.* 2009; Bois *et al.* 2013). *C. elegans* feed on bacteria and other microbes including yeasts (Félix and Braendle 2010). The pharynx, a neuromuscular pump, contracts and relaxes rhythmically to draw in liquid and suspended particles to grind them up before transporting them

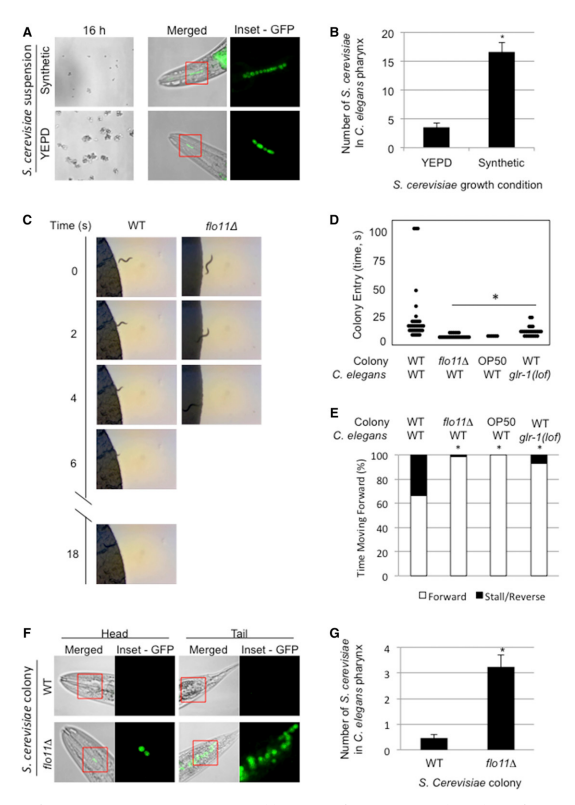


Figure 7 Impact of Flo11p on predator deterrence in yeast colonies. (A) Suspensions of GFP-labeled WT cells (PC6733) grown for 16 hr in synthetic or YEPD media (16 hr) were examined. Merged fluorescent and brightfield images of heads of mounted *C. elegans* showing fluorescent *S. cerevisiae* inside the animal pharynx after 45 min feeding (merged). Fluorescent images of individual *S. cerevisiae* inside the animal pharynx (inset = GFP). (B) Bar graph showing difference in average number of *S. cerevisiae* cells in *C. elegans* pharynx between YEPD- and synthetic-grown suspensions. Error bars are SE. * P < 0.05. (C) Selected images from videos showing WT *C. elegans* entering WT or $flo11\Delta$ yeast colonies. A set of representative videos are available in the supplement (Videos S8–S15). (D) Dot plot showing the time required for worms to enter the indicated colonies. WT (N2) or glr-1(lof) *C. elegans* worms were placed on plates containing WT or $flo11\Delta$ yeast, or the *E. coli* food strain OP50. (E) Stacked bar graphs comparing *C. elegans* forward movement to stalls and/or reversals in the indicated yeast or *E. coli* colonies. (F) Three-day-old colonies of WT (PC538) and $flo11\Delta$ (PC1029) cells

to the intestine. Although the pharynx pumps continuously, previous studies have shown that microbe size is a determining factor in C. elegans feeding; large food is often excluded during this grazing behavior (Avery and Shtonda 2003; Shtonda and Avery 2006; Fang-Yen et al. 2009). One function of cell adhesion molecules is to allow cells to adhere to each other, leading to large colonies. To observe the impact of such adhesion, wild-type S. cerevisiae cells were grown in conditions that either permit cell-cell adherence (Figure 7A, YEPD) or suppress adherence (Figure 7A, synthetic). C. elegans were transferred into suspensions of S. cerevisiae as per established methods (Bois et al. 2013). We found that lessadherent S. cerevisiae (Figure 7A-B, synthetic) were taken into the pharynx of C. elegans in greater numbers than S. cerevisiae that adhered to each other (Figure 7A-B, YEPD). This finding is consistent with the idea that cell-cell adhesion protects cells from ingestion by C. elegans.

We next assessed how yeast cells growing in adhesive colonies might impact the feeding behavior of C. elegans. To test this possibility, the time required for worms to penetrate yeast colonies from nose to tail was measured for ≤ 100 sec (Figure 7, C and D and Videos S8–S15). Worm stalling and reversing out of the colony was also examined (Figure 7E). After 3 days of colony growth, a wild-type yeast strain that formed normal, ruffled colonies slowed *C. elegans* as it attempted to penetrate the colony (Figure 7, C-E, and Videos S8 and S9). In some cases, worms abandoned entering the colony entirely. In contrast, a $flo11\Delta$ colony, which fails to form ruffled colonies, did not deter worms from entering (Figure 7, C–E, and Videos S10 and S11). flo11 Δ colonies were equally vulnerable to nematode entry as a colony of OP50, an E. coli laboratory food strain for C. elegans (Brenner 1974) (Figure 7, D and E, and Videos S12 and S13). Additionally, C. elegans transferred directly into flo11\Delta S. cerevisiae colonies ingested more cells than C. elegans transferred into wild-type S. cerevisiae colonies (Figure 7, F and G). It is important to note here that our results outline an association between a lack of physical barrier and a higher rate of being consumed by C. elegans. As such, we expect any mutant that lacks the physical barrier conferred by adhesive colonies would be similarly vulnerable.

C. elegans might be excluded from wild-type colonies due to the physical barrier encountered by wild-type colonies expressing *FLO11*. Additionally, exclusion might also result from a change in a worm's sensory response to the colony surface. Wild-type *C. elegans* halt their forward locomotion and initiate backward movement in response to a light touch to their anterior-most tip (nose). This avoidance of touch to the worm's nose has been termed the nose-touch response (Kaplan and Horvitz 1993), a behavior involving the ASH polymodal nociceptive sensory neurons (with a small contri-

bution from the FLP and OLQ sensory neurons) (Kaplan and Horvitz 1993). The glutamate-gated ion channel GLR-1 functions in the downstream command interneurons and is required for the nose-touch response (Hart et al. 1995; Maricq et al. 1995). To test the possibility that C. elegans do not effectively enter yeast colonies due to the activation of this mechanosensory avoidance response, experiments were performed with C. elegans glr-1(n2461) loss-of-function (lof) mutant animals, which are defective for nose touch, exposed to a wild-type yeast colony. The glr-1(lof) mutant animals penetrated colonies faster than wild-type worms, though this was still slower than wild-type worms penetrating $flo11\Delta$ yeast colonies (Figure 7D, and Videos S14 and S15). In addition, the glr-1(lof) mutant animals had fewer stalls and reversals upon initial colony entry than wild-type C. elegans (Figure 7E). Taken together, our data reveal a clear association between the formation of adhesive contacts between yeast cells in surface-growing colonies and predation in a laboratory setting, both by becoming too large to eat and also by forming a physical barrier. Further experiments are needed to establish if protection against macroscopic predators is actually an evolutionary driving force for this phenotype.

Flo11p-dependent ruffles impact colonial heat dissipation

Colony ruffling increases the surface-to-volume ratio and may provide benefits such as efficient thermoregulation, or other such adaptations to the environment (Palková and Váchová 2006). We also observed that heat-shock response proteins were among the most induced surface-growth targets (Figure 6A, HSP33 and SNO4). To test whether ruffles made by Flo11p-dependent adhesion impact the temperatures of colonies, ruffled wild-type colonies and smooth flo11 Δ colonies were examined by infrared imaging (Figure 8A). Thermal imaging showed that the average temperature of a ruffled colony was 0.31° Cooler than the average temperature of a smooth colony (Figure 8B, black bars, P < 0.05, n = 6). The average coolest region of a ruffled colony was 0.33° cooler than the average coolest region of a smooth colony (Figure 8B, white bars, P < 0.05, n = 6). Additionally, the coolest regions of the ruffled colonies (Figure 8C, <30°) were the ruffles themselves (Figure 8C, merged). In a study of cell wall stresses, $flo11\Delta$ cells were also shown to have modest temperature sensitivity at 37° (Chow et al. 2018). In summary, Flo11p-dependent ruffling might aid thermoregulation in yeast colonies.

Discussion

Here, we explored the role of signaling pathways that regulate filamentous growth in mediating adhesion-based surface

expressing-plasmid borne GFP (PC2560). *C. elegans* transferred into the colony for 45 min before mounting. Merged fluorescent images of head and tail regions of *C. elegans* showing fluorescent *S. cerevisiae* inside the animal (merged). Fluorescent images of individual *S. cerevisiae* inside the animal (inset = GFP). (G) Bar graph showing difference in average number of *S. cerevisiae* cells in the *C. elegans* pharynx between WT and $flo11\Delta$ colonies. Error bars are SE. * P < 0.05. WT, wild-type; YEPD, yeast extract, peptone, and dextrose.

responses, including mat formation and colony patterning. Fungal cells commonly grow on surfaces, which poses unique challenges due to the heterogeneity of the nutrients within different parts of the colony and direct exposure to environmental conditions. Understanding the responses to growth on surfaces is important because fungal pathogens exhibit mat growth and form invasive filaments (e.g., hyphae) on the surface of the host, and on inert surfaces during early steps in host colonization.

Analysis of signaling pathways that coregulate adhesion-dependent surface responses

Intracellular signaling pathways can operate in functionally interconnected networks (Levchenko 2003). How multiple pathways operate in a coordinated manner to achieve morphogenetic responses with high fidelity remains a mystery. By examining the major signaling pathways that regulate filamentous growth in yeast, we have identified new roles for the filamentation network in regulating colony patterning and mat growth. Thus, the filamentation regulatory pathways may have a general function in regulating the growth of cells in surface communities.

We also have identified the chromatin remodeling complex, SAGA, to be a regulator of filamentous growth. SAGA and the previously characterized Rpd3p pathway (Bernstein *et al.* 2000; Chavel *et al.* 2010) are involved in the epigenetic modification of chromatin. SAGA is an evolutionarily conserved chromatin remodeling complex and a member of the histone acetyltransferase family of proteins (Wang and Dent 2014). SAGA's functions are diverse and include changes in transcription that result in cell differentiation (Wang and Dent 2014; Hirsch *et al.* 2015). In yeast, SAGA controls the expression of a specific set of growth-promoting genes (Bruzzone *et al.* 2018). In fission yeast, TORC1 and TORC2 converge to regulate SAGA in response to nutrient availability (Laboucarié *et al.* 2017). Moreover, components of SAGA have recently been shown to impact the virulence of *Fusarium* (Gao *et al.* 2014).

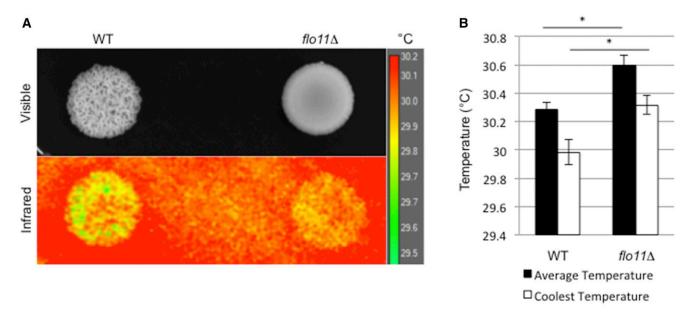
Expression profiling was performed to evaluate the roles for a subset of pathways that regulate adhesion-dependent surface growth. We identified an unexpectedly large number of differentially expressed genes. Many of the genes were regulated by a single pathway. However, as might be expected of a dense network of functionally connected pathways, each pathway impacted every other pathway by regulating overlapping targets as well as genes encoding pathway components. A key example of this overlap, even in the expanded set of pathways, was the gene encoding the adhesion molecule Flo11p. This may be expected given the critical roles that Flo11p plays in regulating filamentous growth (Rupp et al. 1999), mat formation (Reynolds and Fink 2001), and colonial patterning (Granek and Magwene 2010). Moreover, the FLO11 gene is a hub where many signaling pathways and transcription factors converge (Rupp et al. 1999). The other gene was PGU1, which encodes a secreted plant cell walldegrading enzyme. Given that one surface that budding yeast commonly encounter is the surface of plants, especially rotting fruit, it may not be surprising that yeast induce this gene when undergoing surface growth. Transcriptional induction of the *PGU1* gene may also result from the growth of cells on agar, which is a potential substrate, and may be independently regulated. Therefore, one function of the filamentation regulatory network may be to coordinately regulate target genes that are needed to respond to the challenges of growing on surfaces.

We also uncovered new connections between signaling pathways that regulate adhesion-dependent surface growth. One connection was between the RAS and RTG pathways, which coregulate a substantial number of target genes. The coregulated targets support a function for the RAS pathway in mitochondrial control. The RAS pathway is a global nutrientsensing pathway in yeast and other organisms (Zaman et al. 2008). The fact that RAS coregulates targets of the RTG pathway supports the idea that RAS plays a critical role in the response to mitochondrial stress. Our results are consistent with previous observations that connect RAS to the overall regulation of the mitochondria. RAS is required for growth on nonfermentable carbon sources, mitochondrial enzyme content (Dejean et al. 2002), and citrate synthase activity (Swiegers et al. 2006; Chavel et al. 2014). Interestingly, components of the RAS pathway, including the GTPase activating protein Ira1p and adenylate cyclase Cyr1p, associate with mitochondrial membranes, which might impact RAS pathway function or activity at this site (Belotti et al. 2012). Moreover, RAS has been implicated in working with RTG to promote longevity in yeast (Kirchman et al. 1999). Thus, the functional interaction between the two pathways to regulate aspects of mitochondrial health might impact the cell's overall life span. Further experiments will be required to determine how the RAS and RTG pathways coordinate the response to mitochondrial problems. Though the nature of this relationship is unclear, subsequent studies may better characterize the mechanism by which each pathway regulates the other's hallmark targets.

We also show that Dig1p and Ste12p, which are commonly thought to reciprocally regulate the same set of target genes, actually regulate a partially nonoverlapping set of targets. One that stood out in the Dig1p-regulated set were genes that play a role in zinc uptake and metabolism. This observation is consistent with a previous set of findings that show unique expression profiles and binding sites for this activator–repressor pair (Breitkreutz *et al.* 2003; Zeitlinger *et al.* 2003).

Yeast respond to surface growth by expressing a subset of stress-response genes

We also explored the hypothesis that the activity of signaling pathways may change in different parts of a colony over time. We found this to be the case: the activity of the fMAPK pathway was different in different parts of the colony and in different aged colonies. This finding has broad implications in understanding how target genes might be induced during surface growth. For example, it might be expected that other time points or harvesting from certain regions of the colony might



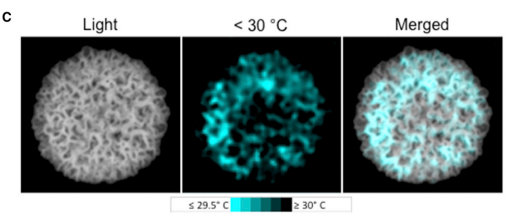


Figure 8 Effect of cell–cell adhesion on thermoregulation. (A) Visible and infrared images of WT and $flo11\Delta$ colonies. Colonies were incubated at 30° and removed from the incubator. The image was taken at < 30 sec after removal of lid. Scale temperature in degrees Celsius. (B) Difference in average and coolest recorded temperatures between WT and $flo11\Delta$. * P < 0.05, n = 6. (C) Visible light (Light) and thermal images of cool colony regions (< 30°) of the WT colony from (A). "Merged" shows the overlapping of cool regions with colony ruffles. WT, wild-type.

result in different expression profiles. The comparative RNAseq analysis performed here may be an oversimplification of the actual gene expression changes that occur within a colony, which may vary in different parts of a colony and over time.

In the study, we also examined genes that were differentially expressed during surface growth as compared to growth in liquid culture. The genes that were differentially expressed tell us about the challenges of growing in the unique environment of surface growth. Several of the differentially expressed genes identified fit with what one expects cells to encounter when growing on surfaces. One might expect that cells experience anaerobic stress within colonies compared to when growing in liquid conditions, where they are uniformly aerated by shaking. Likewise, cells in colony exteriors would be expected to be vulnerable to desiccation. Interestingly, we show that colony growth patterns change in response to these stresses. Specifically, differentially regulated genes impact colony patterning in response to stress.

One prominent class of differentially expressed genes regulates the heat-shock response. This discovery was at first perplexing, given that cells were grown in liquid and on plates at the same temperature. However, thermal imaging verified that colonies do not have a uniform temperature. This may result from the generation of heat due to metabolic activities of cells within the colony, which may lead to temperature differences in different parts of the colony. We also show that ruffles dissipate heat more efficiently than other parts of the colony. The increased surface area of ruffles may promote heat dissipation. Therefore, one function of colony pattern formation might be to efficiently dissipate heat that results from cellular metabolic activities.

Adhesive contacts between yeast cells may offer protection from macroscopic predators

Yeast and other microbes produce colonies that have a ruffled appearance due to contacts between cells. One established

reason for cooperation among individuals in microbial populations is protection (West et al. 2007). For example, cells in the exterior of flocs protect cells in the interior from toxins and antibiotics (Smukalla et al. 2008). Microbial cells are also subject to predation by macroscopic predators. Macroscopic predators, like nematodes, graze on microbial communities. Predator-prey interactions have been studied in bacterial populations, which have developed diverse strategies to evade macroscopic predators. Depending on the species, bacteria have been shown to evade predators by the formation of mats (Darby et al. 2002; DePas et al. 2014; Nandi et al. 2016), by the secretion of proteases (Vaitkevicius et al. 2006) and exopolysaccharides that interfere with the predator's responses (Begun et al. 2007), and by phase variation, in which bacterial cells express different classes of cell surface proteins to evade detection (Dahl et al. 2011).

Although predator-prey studies are less-well characterized in fungal species, yeast and other fungal microorganisms have ecologically relevant interactions with animals in the wild, including grazing by wasps (Stefanini et al. 2016), flies (Fischer et al. 2017), and other invertebrates. Using two wellcharacterized laboratory genetic models, we provide evidence that adhesive contacts between yeast cells may deter efficient predation by nematodes. While individual cells in suspension were vulnerable to being ingested, cells that adhered to each other and formed clumps were too large to be eaten. Once cells settled out of suspension and grew into a colony, adhesive colonies formed a physical barrier to deter penetration by worms and also trigger the "nose-touch" mechanosensory behavioral avoidance response. Although the predator-prey model set up in the laboratory may or may not reflect a true ecological relationship between the two species, experiments using model organisms can provide insights into encounters that may occur in the wild. Our data provide an empirical framework to test the hypothesis that the physical interaction among yeast cells in communities can protect individuals from macroscopic predation in the wild.

In conclusion, we have explored the roles of filamentous growth regulatory pathways in regulating multiple adhesion-dependent surface responses in yeast. Dissection of the regulatory pathways by comparative RNA-seq and mutant phenotype analysis revealed new components and targets, which emphasize the common and unique roles that pathways play in coordinately regulating colonial surface responses in this organism. Stress-responsive genes that respond to challenges associated with surface growth were also characterized. The signaling components, target genes, and stress-responsive genes identified here might be common among other fungi, including pathogens. Our results may be particularly relevant to early steps in fungal virulence, where cells must attach to and penetrate surfaces to colonize the host.

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