Mat Formation in *Saccharomyces cerevisiae* Requires Nutrient and pH Gradients

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The ability of *Saccharomyces cerevisiae* to form morphologically complex colony-like structures called mats requires expression of the cell surface glycoprotein Flo11p and growth on a semisolid surface. As the mat grows, it forms two visually distinct populations called the rim (edge of the mat) and the hub (interior of the mat), which can be physically separated from one another based on their agar adherence properties. Here, we show that growth of the mat on a semisolid agar surface creates concentric glucose and pH gradients in the medium that are required for the differentiation of the hub and rim. Disruption of the pathways that respond to changing levels of glucose block mat formation by decreasing *FLO11* expression. However, in wild-type cells, Flo11p is expressed in both portions of the structure. The difference in adherence between the rim and hub appears to be a consequence of the reduced adherence of Flo11p at the elevated pH of the rim.

Microbes exhibit “multicellular” behaviors such as swarming and the formation of colonies, fruiting bodies, and biofilms (1, 12, 26, 27, 28). All of these behaviors depend on cells that interact with one another and the local environment. For example, fruiting body formation in *Myxococcus xanthus* occurs in response to starvation conditions (12). Biofilm formation is regulated by more diverse stimuli, depending on the microbe, but can be divided into two basic categories, surface conditions and nutrient conditions (15, 16, 20). Many of these multicellular behaviors depend upon a solid support and are not manifest in liquid cultures.

Mat formation in the baker’s yeast *Saccharomyces cerevisiae* has many of the features of other microbial multicellular behaviors. The formation of mats is dependent upon the nature of the surface, the concentration of glucose, and the genetic background of the strain (24). Mats are formed on semisolid agar surfaces (0.3%) and not in liquid, and experiments have shown that the surface is required for the initial formation of this structure (19).

As *S. cerevisiae* grows on the wet surface of a semisolid agar petri plate, it forms a mat that spreads over the agar. The mat is a structure with an interior region called the hub, which is distinguished by channels and wrinkles, and a smooth periphery called the rim (Fig. 1C). Mat formation has been shown to be dependent on the *FLO11* gene: a flo11Δ null mutant fails to form a mat but, instead, grows as a poorly spreading mass of cells (Fig. 1A) (24). *FLO11* is a member of a superfamily of genes encoding cell-surface adherence proteins found in *S. cerevisiae* and other yeasts including *Candida albicans* and *Candida glabrata* (31).

In this report, we designed experiments to identify aspects of the semisolid surface that create differences between the rim and hub. We found that the mat forms glucose and pH gradients in the agar, and these gradients lead to alterations in mat formation. These results suggest that a higher pH in the rim may alter Flo11p function so that the mat is less adherent, thereby permitting the vanguard of the structure to continue spreading radially across the surface. The expression of *FLO11* is maintained in both the rim and hub.

**MATERIALS AND METHODS**

**Media.** The medium used in this study consisted of YPD made with yeast extract (VWR or Fisher), Bacto peptone (VWR), and dextrose (Fisher) (29). Plates were made with 2% agar for standard cell growth and maintenance. Low-agar plates were made with YPD medium and 0.5% agar (24). YPD plates containing 200 μg/ml G418 (Sigma or Fisher) were used to select transformants. YPD plates were buffered to pH 4.9, 5.4, or 5.8 by the addition of citrate buffer at pH 4.0, 4.5, or 5.0, respectively, to the medium to a final concentration of 20 mM.

**Yeast strains.** All of the strains used in this study are from the yeast genetic background *S. cerevisiae* strains. The strains and their genotypes are shown in Table 1. Mutants were made by PCR-based gene disruptions using deletion mutants from the Yeast Knock Out Collection (Open Biosystems/Research Genetics) as templates (33). Strains in the Yeast Knock Out Collection carry gene disruptions in which the open reading frame (ORF) is replaced by the G418 resistance cassette *kanMX4* (32). Primers (Table 2) were used to amplify DNA encompassing each deleted ORF plus ~300 bp flanking either side of the ORF. The resulting PCR product was then transformed into TBR1 by standard lithium acetate transformation protocols (8), and transformants were selected on YPD plates containing 200 μg/ml G418 (24). PCR with primers (Table 2) that annealed outside of the disruption construct and inside the *TEF* promoter of the *kanMX* cassette (32) were used to confirm each disruption. Strains that were created by this approach were ykl1Δ and sos2Δ. The sos2Δ mutant was made using the *kanMX6* cassette from plasmid pFA6a-kanMX6 by a method described previously (17). The yeast strain L6906 (9, 10) carrying a hemagglutinin (HA)-tagged form of *FLO11* (*FLO11-HA*) was used for the immunofluorescence analyses. The Flo11-HA protein carries a triple HA tag inserted between amino acids 30 and 31 in the N terminus of the protein.

**Overlay adhesion assay.** Strains were grown on low-agar plates at ~23°C for 5 days, and then a piece of either Saron or Reynolds brand commercial plastic wrap was laid over the top of the strain and allowed to adhere to the surface by capillary action. Enough plastic was left on either side so that it could be gripped without touching the agar, and it was then removed by pulling up evenly on both sides. The plastic was inverted and laid on a copy stand and photographed in
order to document the non-agar-adherent cells (rim). The cells remaining on the agar were also photographed to document agar-adherent cells (hub).

Northern blotting. Northern blotting was performed as described by Sambrook and Russell (25) using Church buffer for prehybridization and hybridization steps (6). A Techne Hybrigene oven set at 65°C was used for all incubation and wash steps. The cells were grown in liquid YPD medium to logarithmic phase (optical density at 600 nm [OD600] of between 0.5 and 1.0) or post-logarithmic phase (diauxy; OD 600 of 4.0; cells began growing very slowly and doubled only once more over the next 24 h). The cells were collected by centrifugation, aspirated, flash-frozen in a bath of dry ice and methanol, and stored at −80°C. Total RNA was collected by acid-phenol extraction (13), and 10 μg of total RNA was subjected to Northern blotting analysis. A PCR product (using primer TRO367, ATGCAAAGACCATTTCTACTCG, and primer TRO368, TGCCAGGAGCTTGCATATTGAG) corresponding to the first 484 bp of the FLO11 ORF was used to probe the total RNA. The probe was labeled with [α-32P]dATP using a Stratagene Prime-It II random primer kit, and following

FIG. 1. The mat adheres to the agar plate in a Flo11p-dependent manner and forms a glucose gradient. (A) Wild-type (WT) and flo11Δ strains were inoculated on the center of YPD plates containing 0.3% agar and grown at 23°C for 5 days (Mat). The strains were then overlaid with plastic wrap. This was removed by gripping both sides of the wrap and pulling up. The cells remaining adhered to the agar where photographed (Hub). The plastic wrap and cells that were removed with it were inverted and photographed (Rim). (B) Urinalysis strips (Bayer) were used to measure the glucose levels in different sections of the mat as indicated. The percentage (weight/volume) of glucose in the agar is indicated to the side of the strip. The lines indicate the sampling points. (C) The hub (H) and rim (R) of a wild-type mat are shown in greater detail. Bar, 0.5 cm.

| Strain | Genotype | Reference |
|--------|----------|-----------|
| TBR1   | MATα ura3–52 leu2:hisG his3–hisG flo11∥kanMX6 | 24 |
| TRB5   | MATα ura3–52 leu2:hisG his3–hisG flo11∥kanMX6 | 24 |
| YTR140 | MATα ura3–52 leu2:hisG his3–hisG yak1∥kanMX6 | This study |
| YTR141 | MATα ura3–52 leu2:hisG his3–hisG snf1∥kanMX6 | This study |
| YTR142 | MATα ura3–52 leu2:hisG his3–hisG ras2∥kanMX6 | This study |
| L6906  | MATα ura3–52 his3–hisG FLO11∥HA | 10 |

TABLE 2. Primers used in this study

| Primer | Purpose | Sequence |
|--------|---------|----------|
| TRO369 | Confirm disruptions | GCACGGTCAGGCTGTCAGGG |
| TRO158 | Disrupt YAK1 | TACAAAATAGCCCGATGTCG |
| TRO159 | Disrupt YAK1 | CTGAGCTTCTTATCTTIGTTTACCT |
| TRO160 | Confirm yak1Δ | AGAATGACATATACACTATATGTCG |
| TRO309 | Disrupt SNF1 | ATAGAAAGTTTTTTTGGATAACAA |
| TRO310 | Disrupt SNF1 | AGAACCTCCATATCATTCTT |
| TRO94  | Confirm snf1Δ | CCAAAGCATAGGCTTTGGGCTT |
| TRO331 | Disrupt RAS2 | CAAGGTTACATACGAACACAAC |
| TRO312 | Disrupt RAS2 | ATTCCAAGTGGAACACTCCTT |
| TRO313 | Confirm rasΔ | TGACATTTAGGCCGTAGAGC |
| FLO11 fw | RT primer FLO11 | CAAG |
| FLO11 rev | RT primer FLO11 | CTTGCAATTTAGGCCGCACTAC |
| ACT1 fw | RT primer ACT1 | CTTCCACCATTGTAGAAGAAGA |
| ACT1 rev | RT primer ACT1 | CAAAGGCGACGTAACATAGTTT |
| SNR190 fw | RT primer SNR190 | TCTTTCTCTCGTCCGATTCG |
| SNR190 rev | RT primer SNR190 | TACCTTGGACATGAGAAGAG |

TRO369 was used as a reverse primer in combination with the primers listed as confirming primers to confirm disruptions on the chromosome by PCR. fw, forward; rev, reverse.
probing, the blots were visualized on a Storm Phosphorimager. The data were quantified using ImageQuant software, and *FLO11* expression for each strain was normalized to bands from 28S rRNA quantified on a Bio-Rad Chemidoc XRS photodocumentation system using QuantityOne software from Bio-Rad.

Real-time RT-PCR. Seven-day-old mats were utilized for real-time reverse transcription-PCR (RT-PCR). A Pipetman was used to pipette up cells from the rim, the hub, or what was referred to as the “middle,” an intermediate area in between the rim and the hub. The isolated cells were added to ~25 ml of ice-cold water, pelleted by centrifugation at 4°C, and washed with 1 ml of ice-cold water (during which time they were transferred to a microcentrifuge tube), and the washed pellet was frozen at −80°C. This process was carried out on ice for each sample within 10 min to ensure optimal preservation of the mRNA. Total RNA was isolated as described previously (23). Total RNA was further purified and treated with DNase twice using an RNeasy Mini Kit (Qiagen 74104). Reverse transcription was performed using TaqMan reverse transcription reagents (catalog no. N808-0234; Applied Biosystems) according to the manufacturer’s standard protocol, and real-time PCR was performed with Sybr Green PCR Master Mix (catalogue no. 4309155; Applied Biosystems) or TaqMan Universal PCR Master Mix without AmpErase uracil-N-glycosylase (catalog no. 4326614; Applied Biosystems) on an Applied Biosystems 7500 RT-PCR system using a two-step program (15 s melting at 95°C and 60 s annealing-extension at 60°C).

Immunofluorescence of Flo11 on the cell surface of cells from the rim and hub. Mats were grown on low-agar plates for 5 days, and then a pipette tip, cut off at the end to create a wider bore, was used to collect cells from the rim or hub of the mat. The cells were resuspended in phosphate-buffered saline (PBS) with 4% formaldehyde and fixed for 1 h at room temperature. The cells were washed with PBS, blocked with PBS–2% bovine serum albumin (BSA), and incubated with anti-HA antibody (HA.11; Covance) for 1 h. The cells were washed three times with PBS–2% BSA, incubated with goat anti-mouse secondary antibody conjugated to Cy3 (Jackson Immunobiology), washed three times with PBS–2% BSA again, resuspended in PBS–2% BSA, and viewed on an Olympus BX50 microscope.

RESULTS

Fractionation of cells from the rim and hub of the mat. Growth on low-agar plates produced a clear visual differentiation between the cells at the rim and those at the hub of a wild-type mat (Fig. 1C). In order to analyze the two populations of cells, we overlaid the mat with plastic wrap. When the plastic wrap was withdrawn, the nonadherent cells at the rim were drawn up with the wrap (Fig. 1A, rim), whereas the agar-adherent cells in the hub remained behind (Fig. 1A, hub).

This analysis revealed a new aspect of mat formation: the growing mat could be partitioned into two distinct subpopulations based on agar adherence properties. The point on the growing mat at which the appearance changed from smooth (rim) to channel-ridden (hub) (Fig. 1C) is also the point at which the adherence phenotype changed. By contrast, the *flo11* strain, which is uniformly smooth, failed to adhere to the agar and was completely removed from the agar by the plastic wrap (Fig. 1A).

A glucose gradient is present in the mat and influences its development. Since a low glucose concentration promotes adhesion of *Saccharomyces* cells to polystyrene (24), we determined whether there was a gradient of glucose that correlated with the different cell populations revealed by the overlay adherence assay. This analysis revealed that the uniform glucose level present prior to growth of the mat (2%) was replaced during growth by a concentration gradient that decreased from the rim toward the hub (Fig. 1B). Thus, there is a gradient of glucose in the growing mat such that the concentration is higher at the rim and lower at the hub.

Glucose limitation appeared to be an important signal for regulating mat formation, but other nutrients could be depleted in the hub and might play a role as well. If glucose is the key nutrient in regulating mat formation, then medium containing increased levels of glucose, but identical levels of all other components, should slow mat formation. Plates were poured that contained 1% yeast extract, 2% peptone, and 2%, 4%, or 6% glucose. It was found that, in comparison to 2% glucose plates, hub formation was delayed by 1 day or 2 days in plates containing 4% or 6% glucose, respectively (Fig. 2).

Although a glucose gradient is formed in the growing mat (Fig. 1B), it was unclear if the gradient itself was required for mat formation. In order to test this, the gradient was perturbed.
in a mat by the addition of glucose to the medium outside of the growing mat. If a glucose gradient were required for maturation of the mat, then the addition of glucose to one side of the mat should slow maturation of the mat on the side to which glucose was added. After a wild-type mat had grown for 3 days, a small pit was dug in the agar outside of the mat with a pipette tip, and 50 μl of 40% glucose or water was added to the pit (Fig. 3, 3d). After two more days (5d) the plates were examined and photographed. The black circles mark the pit in the agar where the glucose or water was added.

The fact that a gradient of glucose drives mat formation and that the hub is depleted of glucose compared to the rim suggested that cells in the rim were actively growing whereas those in the hub had stopped growing. This was tested by a “healing” assay. The mat was damaged by using a pipette tip to tear a hole in both the rim and the hub on day 5 (Fig. 4). Two days later (day 7), the hole made in the rim was detectable only as a depression in the agar beneath the mat as cells had proceeded to grow over it. Conversely, the hole in the hub looked much like it had at day 5 with only slight changes in appearance (Fig. 4). This result indicated that the cells in the hub exhibited little to no growth, whereas the cells in the rim continued to divide and spread.

Glucose-sensing genes control mat formation and FLO11 expression. Several signaling pathways are associated with sensing glucose levels and regulating filamentous growth in *S. cerevisiae*. These pathways include the Ras/cyclic AMP, Snf1p kinase, and Yak1p kinase pathways. Genes representing these pathways include *SNF1* (3), *YAK1* (18), and *RAS2* (11). *SNF1* has been shown to be required for mat formation on low agar (21), biofilm formation on plastic, *FLO11* expression (14), and filamentous growth on solid 2% agar (7). Isolectric mutants for these genes were generated in the strain TBR1 (24) and tested for mat formation. Disruptions of *SNF1*, *YAK1*, or *RAS2* all caused defects in mat formation (Fig. 5). Although they differed in the extent of their adhesion defects (Fig. 5), all three mutants formed few or no discernible channels and spread poorly.

The three glucose-sensing mutants, *yak1Δ*, *ras2Δ*, and *snf1Δ*, were tested for their effects on *FLO11* expression. It was found that all three mutants caused a decrease in *FLO11* expression compared to the wild-type strain when grown to exponential phase in liquid YPD medium containing 2% glucose (Fig. 6A). The *ras2Δ* mutant did not cause a strong decrease in *FLO11* expression compared to the *yak1Δ* or *snf1Δ* mutants. However,
when the three mutants were compared to wild type in liquid YPD medium during glucose stress (growth to post-exponential phase), the ras2Δ mutant, like the yak1Δ and snf1Δ mutants, showed a significant decrease in FLO11 expression (Fig. 6B).

Flo11p is expressed on the surface of cells in the rim and hub. Since the glucose-sensing genes affect the expression of FLO11 and glucose levels are lower in the hub, the morphological differences between the rim and the hub could be a consequence of FLO11 expression. FLO11 mRNA levels were measured in the rim, the intermediate region where the rim and hub meet (middle), and the center of the hub by real-time RT-PCR. If ACT1 is used to normalize the FLO11 levels, there is no significant difference in FLO11 mRNA levels between the rim and hub (Fig. 7A). By contrast, if SNR190 is used for normalization, FLO11 mRNA levels appear highest in the rim and lowest in the hub (Fig. 7B). This difference in expression is also observed in the ACT1 mRNA levels, which appear to be higher in the rim than the hub when SNR190 is used for normalization (Fig. 7B). ACT1 levels are affected by the growth state of cells (5, 22), but SNR190 levels are stable between cells in logarithmic and stationary phase (7a). The indeterminate growth state of cells in the hub makes an accurate measurement of FLO11 mRNA levels difficult, but it is clear that FLO11 mRNA is expressed in both parts of the mat.

The availability of a strain expressing a functional version of Flo11p with an HA tag (10) permitted a direct assessment of the levels of Flo11p on the surface of cells in the rim and hub. This strain, containing the FLO11::HA construct inserted into the chromosome at the FLO11 locus, forms mats like the wild-type strain. The percentage of cells expressing Flo11p on their surfaces was assessed by indirect immunofluorescence and was found to be the same in the rim and hub (38% ± 2% and 38% ± 5%, respectively) (Fig. 8). As has been previously reported (10), there was heterogeneity of FLO11 expression among the cells, but there did not appear to be any difference.
The same RNAs from the rim, hub, and middle that were used to assess the \textit{FLO11} levels were used to ascertain the levels of the other \textit{FLO} genes (\textit{FLO1}, \textit{FLO5}, \textit{FLO9}, and \textit{FLO10}), which are known to affect aggregation and adhesion (30, 31). Although strains from the \textit{S288c} background do not express these other \textit{FLO} genes in liquid, they might be expressed on the low-agar petri dishes. Analysis of these transcript levels by real-time RT-PCR using primers specific for these genes showed no significant expression in any portion of the mat using either \textit{ACT1} or \textit{SNR190} as a normalization standard (Fig. 7A and B, respectively).

**Mat formation is controlled by a pH gradient.** Analysis of the mat revealed that in addition to the glucose gradient (Fig. 1B), the mat also established a pH gradient. The pH of the medium was originally 5.8 but was altered by the yeast, and after growth of the mat, a pH gradient formed between the sites of the hub (4.7) and rim (5.0) (Fig. 9). Moreover, the cells at the edge of the rim were juxtaposed to medium at pH 5.8. To test whether the pH gradient was an important component in the distribution of the cells expressing Flo11p in the two populations.

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exhibited generally decreased spreading compared to mats growing on plates without citrate buffer (Fig. 1A). A slight rim and hub region could be detected in mats grown on plates buffered to pH 5.4, based on appearance (Fig. 10). However, at pH 5.8, the channels that characterize the hub did not form, and the surface of the whole mat was smooth. In the cases of both types of plates (pH 5.4 and 5.8), the surfaces of the whole mats were removed by the overlay adhesion assay (Fig. 10). In contrast, the entire surfaces of mats formed on pH 4.9 plates resembled hubs, and no material could be removed from the surface of the plate using the overlay adhesion assay (Fig. 10). This strong adherence at pH 4.9 is dependent on \(FLO11\) as the flo11\(\Delta\) mutant was completely removed from pH 4.9 plates during the overlay adhesion assay (Fig. 10).

**DISCUSSION**

Gradients of glucose in the petri dish may maintain the expression of \(FLO11\). The semisolid support of the agar plate permits the maintenance of the multiple gradients required for formation of the mat. Previous studies have shown that mat formation is enhanced by low glucose (24). The concentration of glucose is modulated in the petri plate to create a glucose gradient, from the lowest concentration in the hub to the highest in the rim (Fig. 1B). As we have shown, glucose-sensing pathways are required for the upregulation of \(FLO11\) transcription under low

![FIG. 9. Mat formation correlates with a gradient of pH. pH strips were used to measure the pH of medium outside the mat and inside the rim and the hub. The pH values are indicated next to the strips. The lines indicate the sampling points.](image)

![FIG. 10. pH controls mat formation. Wild-type or flo11\(\Delta\) strains were inoculated onto low-agar YPD plates that were buffered to pH 4.9, 5.4, and 5.8 with 20 mM Na citrate buffer; cultures were grown for 5 days at 23°C and photographed. The mat in each case was then subjected to the overlay adhesion assay.](image)
glucose concentrations (Fig. 6B). Loss of glucose sensing results in attenuation of mat formation (Fig. 5).

One possibility is that the glucose gradient helps to maintain Flo11 protein at a relatively constant level throughout the mat. As the cells in the hub are no longer growing (Fig. 4), glucose sensing may provide a compensatory mechanism to ensure that Flo11p is maintained at the cell surface.

**Flo11 protein is expressed throughout the mat.** The finding that the *FLO11* transcript is expressed throughout the mat is consistent with the requirement of *FLO11* for mat formation but raised the question of why cells at the hub were more adherent than those at the rim. One possibility was that Flo11 protein was not properly expressed at the cell surface in rim cells but was in hub cells. In order to function, Flo11p must transist the secretory pathway, arrive at the plasma membrane, and be cross-linked to the cell wall (31). Alterations in any of the steps along this pathway could lead to failure of Flo11p to be properly displayed at the cell surface. However, using a tagged version of *FLO11* (Fig. 8), we showed that there are roughly the same proportion of cells in the rim and the hub with Flo11p on the surface. Moreover, the distribution of the fluorescence was ostensibly the same in the rim cells as it was in the cells within the hub. These methods did not measure protein levels of Flo11p and thus would only detect gross differences in Flo11p on the cell surface. Nonetheless, there is no obvious difference in protein expression or distribution in the populations.

**Other FLO genes are not differentially activated in the mat.** Of the five members of the FLO gene family (*FLO1, FLO5, FLO9, FLO10, and FLO11*), only *FLO11* is transcribed in the ∑1278b strains we used. The other FLO genes are normally transcriptionally silenced (31). However, it was possible that one or more of these silenced genes was transcribed either in the rim or hub under the unique conditions of mat formation. This model would likely require that one of these normally silent FLO genes be coexpressed with *FLO11* as none of the other FLO genes tested (*FLO1* and *FLO10*) can compensate for the loss of *FLO11* function during mat formation in the *flo11Δ* strain. In addition, disruption of *FLO1, FLO5, or FLO10* had no effect on mat formation (T. B. Reynolds and G. R. Fink, unpublished data). This desilencing hypothesis appears to be ruled out by our finding by real-time RT-PCR that there are very low levels of *FLO1, FLO5, FLO9, or FLO10* transcription in the rim and hub (Fig. 7A and B). Of course, there could be other cell surface genes that are required, but they also must cooperate with *FLO11*. There are more than 38 genes encoding glycosylphosphatidylinositol-linked cell wall proteins in yeast (4), leaving a number of other possibilities.

**Gradients of pH in the petri dish affect mat formation.** The fact that the mat establishes a pH gradient within itself may provide an explanation for the differential adherence between the rim and hub (Fig. 9). Flo11p-dependent adhesion has been shown to be dependent on pH, increasing as the pH decreases between values of 5.5 and 3.9 (2). The pH in the hub is ~4.7, but it is 5.0 in the rim and higher adjacent to the rim (Fig. 9).

The importance of the pH gradient was supported by experiments showing that stabilization of the pH in solid medium alters the level of adhesion exhibited by mat cells (Fig. 10). This gradient of decreasing pH appears to contribute to the formation of the mat by increasing Flo11p-dependent adhesion in the hub of the mat compared with the rim.

**Conclusion.** Mat formation is dependent on gradients of glucose and pH established by the growing cell population. These gradients correlate with changes in both the expression and the function of Flo11p, which is necessary for this phenotype.

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**REFERENCES**

1. Allison, C., and C. Hughes. 1991. Bacterial swarming: an example of prokaryotic differentiation and multicellular behavior. Sci. Prog. 75:403–422.
2. Bayly, J. C., L. M. Douglas, I. S. Pretorius, F. F. Bauer, and A. M. Dranginis. 2000. Characteristics of Flo11p-dependent flocculation in *Saccharomyces cerevisiae*. FEBS Yeast Res. 5:1151–1156.
3. Carlson, M. 1999. Glucose repression in yeast. Curr. Opin. Microbiol. 2:202–207.
4. Caro, L. H., H. Tettelin, J. H. Vossen, A. F. Ram, H. van den Ende, and F. M. Kils. 1997. In silico identification of glycosyl-phosphatidylinositol-anchored plasma-membrane and cell wall proteins of *Saccharomyces cerevisiae*. Yeast 13:1477–1489.
5. Chrobak, M. J. 1991. A general topoisomerase I-dependent transcriptional repression in the stationary phase in yeast. Genes Dev. 5:2315–2326.
6. Church, G. M., and W. Gilbert. 1984. Genomic sequencing. Proc. Natl. Acad. Sci. USA 81:1991–1995.
7. Cutten, P. J., J. Schulz, J. Horecka, B. J. Stevenson, Y. Jigami, and G. F. Sprague, Jr. 2000. Defects in protein glycosylation cause SHO1-dependent activation of a STE12 signaling pathway in yeast. Genetics 155:1005–1018.
8. Duina, A. A., and F. Winston. 2004. Analysis of a mutant histone H3 that perturbs the association of Swi/Snf with chromatin. Mol. Cell. Biol. 24:561–572.
9. Gietz, R. D., and R. A. Woods. 2002. Transformation of yeast by lithium acetate/single-stranded carrier DNA/polychethylene glycol method. Methods Enzymol. 350:87–96.
10. Gao, B., C. A. Styles, Q. Feng, and G. R. Fink. 2000. A *Saccharomyces* gene family involved in invasive growth, cell-cell adhesion, and mating. Proc. Natl. Acad. Sci. USA 97:12158–12163.
11. Hallme, A., S. Bumgarner, C. Styles, and G. R. Fink. 2004. Genetic and epigenetic regulation of the FLO gene family generates cell-surface variation in yeast. Cell 116:405–415.
12. Howard, S. C., Y. W. Chang, Y. V. Budovskaya, and P. K. Herman. 2001. The Ras/FKase signaling pathway of *Saccharomyces cerevisiae* exhibits a functional interaction with the Sin3p complex of the RNA polymerase II holoenzyme. Genetics 159:77–89.
13. Jelsbak, L., and L. Sogaard-Andersen. 2003. Cell behavior and cell-cell communication during fruiting body morphogenesis in *Mycococcus xanthus*. J. Microbiol. Methods 55:829–839.
14. Kohrer, K., and H. Domdey. 1991. Preparation of high molecular weight RNA. Methods Enzymol. 194:398–405.
15. Kuchin, S., V. K. Vyas, and M. Carlson. 2002. Snf1 protein kinase and the repressors Nrg1 and Nrg2 regulate FLO11, haploid invasive growth, and diploid pseudohyphal differentiation. Mol. Cell. Biol. 22:5994–4000.
16. Kuchma, S. L., and G. A. O’Toole. 2000. Surface-induced and biofilm-induced changes in gene expression. Curr. Opin. Biotechnol. 11:429–433.
17. Lejeune, P. 2003. Contamination of abiotic surfaces: what a colonizing bacterial. Biophys. J. 11:179–184.
18. Longtime, M. S., A. McKenzie, 3rd, D. J. Demarini, N. G. Shah, A. Wach, A. Brachat, P. Philippsen, and J. R. Pringle. 1998. Additional modules for versatile and economical PCR-based gene deletion and modification in *Saccharomyces cerevisiae*. Yeast 14:953–961.
19. Moriya, H., Y. Shimizu-Yoshida, A. Omori, S. Iwashita, M. Katoh, and A. Sakai. 2001. Yak1p, a DHR family kinase, translocates to the nucleus and phosphorlates yeast Posp2p in response to a glucose signal. Genes Dev. 15:1217–1228.
20. Nguyen, B. A., Upadhyaya, A. V. Oudenaarden, and M. P. Brenner, 2004. Elastic instability in growing yeast colonies. Biophys. J. 86:2740–2747.
21. O’Toole, G. A., L. A. Pratt, P. I. Watnick, D. K. Newman, V. B. Weaver, and R. Kolter. 1999. Genetic approaches to study of biofilms. Methods Enzymol. 310:91–109.
22. Palecek, S. P., A. S. Parikh, J. H. Huh, and S. J. Kron. 2002. Depression of
Saccharomyces cerevisiae invasive growth on non-glucose carbon sources requires the Snf1 kinase. Mol. Microbiol. 45:453–469.
22. Paz, I., J. R. Meunier, and M. Choder. 1999. Monitoring dynamics of gene expression in yeast during stationary phase. Gene 236:33–42.
23. Pernambuco, M. B., J. Winderickx, M. Crauwels, G. Griffioen, W. H. Mager, and J. M. Thevelein. 1996. Glucose-triggered signalling in Saccharomyces cerevisiae: different requirements for sugar phosphorylation between cells grown on glucose and those grown on non-fermentable carbon sources. Microbiology 142:1775–1782.
24. Reynolds, T. B., and G. R. Fink. 2001. Bakers’ yeast, a model for fungal biofilm formation. Science 291:878–881.
25. Sambrook, J., and D. W. Russell. 2001. Molecular cloning: a laboratory manual, 3rd ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
26. Shapiro, J. A. 1995. The significances of bacterial colony patterns. Bioessays 17:597–607.
27. Shapiro, J. A. 1998. Thinking about bacterial populations as multicellular organisms. Annu. Rev. Microbiol. 52:81–104.
28. Stoodley, P., K. Sauer, D. G. Davies, and J. W. Costerton. 2002. Biofilms as complex differentiated communities. Annu. Rev. Microbiol. 56:187–209.
29. Styles, C. 2002. How to set up a yeast laboratory. Methods Enzymol. 350:42–71.
30. Verstrepen, K. J., and F. M. Klis. 2006. Flocculation, adhesion and biofilm formation in yeasts. Mol. Microbiol. 60:5–15.
31. Verstrepen, K. J., T. B. Reynolds, and G. R. Fink. 2004. Origins of variation in the fungal cell surface. Nat. Rev. Microbiol. 2:533–540.
32. Wach, A., A. Brachat, R. Pohlmann, and P. Philippsen. 1994. New heterologous modules for classical or PCR-based gene disruptions in Saccharomyces cerevisiae. Yeast 10:1793–1808.
33. Winzeler, E. A., D. D. Shoemaker, A. Astromoff, H. Liang, K. Anderson, B. Andre, R. Bangham, R. Benito, J. D. Boeke, H. Bussey, A. M. Chu, C. Connelly, K. Davis, F. Dietrich, S. W. Dow, M. El Bakkoury, F. Foury, S. H. Friend, E. Gentalen, G. Giaever, J. H. Hegemann, T. Jones, M. Laub, H. Liao, N. Liebundguth, D. J. Lockhart, A. Lucas-Danila, M. Lussier, N. M'Rabet, P. Menard, M. Mittmann, C. Pai, C. Rebischung, J. L. Revuelta, L. Riles, C. J. Roberts, P. Ross-MacDonald, B. Scherens, M. Snyder, S. Sookhai-Mahadeo, R. K. Storms, S. Veronneau, M. Voelkaert, T. R. Ward, R. Wysocki, G. S. Yen, K. Yu, K. Zimmermann, P. Philippsen, M. Johnston, and R. W. Davis. 1999. Functional characterization of the S. cerevisiae genome by gene deletion and parallel analysis. Science 285:901–906.