

## Generation of stable, non-aggregating *Saccharomyces cerevisiae* wild isolates\*

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**Cellular aggregates observed during growth of *Saccharomyces cerevisiae* strains derived from various natural environments makes most laboratory techniques optimized for non-aggregating laboratory strains inappropriate. We describe a method to reduce the size and percentage of the aggregates. This is achieved by replacing the native allele of the *AMN1* gene with an allele found in the W303 laboratory strain. The reduction in aggregates is consistent across various environments and generations, with no change in maximum population density or strain viability, and only minor changes in maximum growth rate and colony morphology.**

**Key words:** *AMN1*, aggregation, *Saccharomyces cerevisiae*

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### INTRODUCTION

With its long history in laboratory research, budding yeast has been one of the most intensively used model organisms, and there it is hope that it will be the pre-eminent model organism of the 21st century (Botstein & Fink, 2011). Since 1996, when the genome sequence of *Saccharomyces cerevisiae* was published (Goffeau *et al.*, 1996; Mewes *et al.*, 1997), the amount of accumulated data on its molecular genetics and evolution has increased tremendously. However, knowledge about this species' natural environment, life history, population genetics, and ecology is still incomplete. Because such research cannot be done using the few available laboratory yeast strains, there is interest in strains isolated from various geographical and ecological environments (Diezmann & Dietrich, 2009; Liti *et al.*, 2009; Schacherer *et al.*, 2009) and in the pathogenic effects of clinical *S. cerevisiae* strains.

Two reasons why yeast laboratory strains are so widely used are that they are heterothallic and that they are able to form a homogeneous, non-aggregating and non-adhesive planktonic population in liquid laboratory media (Bruckner & Mosch, 2012; Mortimer & Johnston, 1986). This has allowed for optimization of many basic and indispensable techniques used in the yeast laboratories, such as: assessment of cell number based on optical density measurements; determination of colony forming units (CFU) on agar media (e.g., for viability measurements); "streaking" to obtain colonies derived from single cells; fluorescence-activated cell sorting (FACS); Coulter Counter measurements; efficient staining due to equal availability of the dye for all of the cells; fractionation in various density gradients; and many others. Unfortunately, many wild isolates form large robust aggregates of cells during growth in liquid media ("clumpy"

phenotype), rendering these techniques inaccurate, very laborious, or even impossible to perform with such strains.

It was shown that the clumpy phenotype of the wild isolate RM11 depends on variation in *AMN1*, a gene involved in daughter cell separation (Yvert *et al.*, 2003). Nucleotide sequencing of *AMN1* cDNAs from laboratory strains (BY series and related: W303 and S288C) identified the missense mutation: an A to T change at nucleotide position 1104, which results in an amino acid substitution of Val368 for aspartic acid (D368V). In contrast, most wild isolates (RM11) have aspartic acid (D) at the corresponding position of wt*AMN1* protein (www.yeastgenome.org). Deletion of the *amn1* gene gives the aggregating RM11 strain a non-aggregating phenotype (Yvert *et al.*, 2003). This deletion has been shown to result in decreased vegetative growth rate, competitive fitness, in filamentous growth, and a deficiency in invasive growth (www.yeastgenome.org). When the *AMN1* allele from laboratory yeast (W303), was replaced by *AMN1*-RM11, (an allele from the wild yeast strain RM11), the lab strains acquired the clumpy phenotype of wild yeast (Koschwanez *et al.*, 2011).

We report a new genetic technique for obtaining a non-aggregating population from a previously aggregating *S. cerevisiae* strain.

### MATERIALS AND METHODS

We used seven starting strains (listed in Table 1); all are haploid and come from an existing collection of wild isolates in which *HO* and *URA3* genes were replaced by antibiotic markers, *HygMX4* and *KanMX4* cassettes, respectively (Cubillos *et al.*, 2009). Plasmid pJHK110, containing the laboratory allele of *AMN1*-W303 (kindly provided by John Koschwanez, FAS Center for Systems Biology, Harvard University), was cut with *Xho1* enzyme to target integration at the *AMN1* locus, and then transformed into the strains (Gietz & Woods, 2002), selecting for URA prototrophy *via* growth on synthetic complete media without uracil (SC-ura) plates. After 2 to 3 days of incubation, at least eight of the obtained transformants were streaked out onto new SC-ura plates. Individual colonies from these plates were then transferred into liq-

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**Abbreviations:** *AMN1* gene, antagonist of mitotic exit network gene; FACS, fluorescence-activated cell sorting; cDNA, complementary DNA; *HygMX4*, hygromycin antibiotic marker; *KanMX4* genetic antibiotic marker; SC-ura, synthetic complete medium without uracil; YPD, yeast peptone dextrose medium; 5-FOA, 5-Fluoro-ortho acid medium; OD, optical density; MGR, maximum growth rate

Table 1. a) Strains positively transformed with pJHK\_110\_amm1\_W303 inducing the *AMN1* gene to switch into the laboratory W303 allele.

Parental strains	Source	Average number of cells in clumps		Percentage of the aggregates in the population		MGR (underlined higher MGR);	
		parental	trans formed	parental	trans formed	Parental/t-test results *stat.significant	transformed
*UWOPS83-787 mating type a,	Fruit. <i>Opuntia stricta</i>	6.2	1.0	29.9±6.6	2.3±1.3	<u>0.51±0.03/</u> $t_{10, 0.0004} = 6.86$	0.39±0.02
*UWOPS87-2421 mating type a	Fruit. <i>Opuntia megacantha</i>	10.3	1.7	62.2±4.2	9.7±2.5	<u>0.53±0.04/</u> $t_{10, 0.014} = 2.97$	0.48±0.02
*DBVPG1373 mating type a	Soil	6.4	1.7	73.6±3.5	24.7±1.6	<u>0.42±0.02/</u> $t_{10, 0.000003} = -9.39$	<u>0.51±0.02</u>
UWOPS87-2421 mating type alpha	Fruit. <i>Opuntia megacantha</i>	6.1	1.9	89.4±0.8	8.9±2.5	<u>0.50±0.01/</u> $t_{10, 0.024} = -2.67$	<u>0.53±0.03</u>
*Y55 mating type a	Grapes	9.9	1.7	73.3±11.3	22.5±2.2	<u>0.56±0.01/</u> $t_{10, 0.000003} = 9.26$	<u>0.57±0.01</u>
DBVPG1106.3 mating type a	Grapes	7.5	1.6	64.4±1.4	18.3±1.2	<u>0.56±0.01/</u> $t_{10, 0.2} = -1.39$	0.52±0.01
378604X mating type alpha	Clinical isolate ( <i>Sputum</i> )	3.4	2.0	46.0±5.1	19.1±3.4	<u>0.51±0.01/</u> $t_{10, 0.0005} = -4.99$	<u>0.54±0.01</u>
S288c mating type alpha. reference	Rotting fig; laboratory strain	1.7	1.7	7.3±2.3	7.3±2.3	0.49±0.01	0.49±0.01

uid yeast peptone dextrose (YPD) medium and grown at 30°C overnight to allow the plasmid to “pop out” by homologous recombination between the two copies of the *AMN1* gene sequence. From each culture, ~10<sup>5</sup> cells were plated onto 5-Fluoro-ortic acid medium (5-FOA) plates, which selects for cells lacking a functional *URA3* gene, in order to obtain cells which had lost the plasmid and thus retained only one *AMN1* gene (containing either the endogenous or the W303 allele). Colonies that grew on 5-FOA medium were chosen for microscopic observation. The number of aggregates, and the average number of cells within the aggregates, were counted using a Bürker chamber (hemocytometer). Approximately eight transformant clones showing no aggregate formation were obtained from each of the starting strains, and were then subjected to genomic confirmation to determine whether they carried the expected non-clumping W303 *AMN1* allele. Genomic DNA from each clone was obtained using Genra DNA purification Kit with prior lysis of the cell wall by lyticase treatment. To amplify the *AMN1* gene region, the following primers (designed by J. Koschwanez) were used: F:ACGAACCTCCAGGAGACGACACA; R: AAAGGCCTCCCGCG-GACTTAAAACAGAAC, with the PCR program: 95°C – 3 min; 34 cycles of [95°C – 30 s; 55°C – 30 s; 72°C – 1 min]; 72°C – 10 min. The PCR products were treated with the Exo-Sap clean-up kit (Fermentas) and then Sanger sequenced with the Big Dye Terminator kit 3.1 (ABI), using the F primer, and run on a 3130 x 1 genetic analyzer (ABI). The sequences were analyzed using BioEdit software. Clones verified to have the GTT (Val) at codon 368 of the *AMN1* gene, i.e., the W303 allele, were analyzed using the 5 assays described below. (i) *Clumpiness*. The ancestral and transformed strains were grown in liquid YPD medium at 28°C. Then the populations were diluted, and single cells and aggregates were counted using a microscope (Nikon Eclipse E400) and Bürker chamber. Each of the populations was also photographed using a microscope (Nikon Eclipse 80i) and Lucia software (Quality Capture 2560×1920, ex. time

1/125s) (phase contrast PlanApo 40×/0.95) (Table 1). A minimum of 3 independent colonies of each transformant and 6 independent colonies of the control (untransformed parent) were checked. The average percentage of single and double cells and aggregates (defined as groups of cells containing more than 2 cells) was calculated (Table 1). To calculate the average number of cells in the clumps, the number of cells in at least 250 aggregates was counted using the microscope. (ii) *Population density*. Populations of parental and transformed strains were grown for 2 days in liquid YPD media. The density was assessed by counting cells from serially diluted populations using hemocytometer. (iii) *Monocolony phenotype*. Using a micromanipulator, four single cells in stationary phase from each parental (non-transformed) strain and from one of its transformed derivatives were placed on growth and stress media (YPD, synthetic dextrose minimal medium with uracil, SD+ura, ZnSO<sub>4</sub>, NaCl). Cells were allowed to form colonies at 28°C for 3 weeks, then scanned (Epson Perfection 2480 Photo), washed from the plate. Washed plates were scanned then to check for the invasive colony growth. (iv) *Population viability*. Four of the parental and transformed strains in triplicate (marked with an asterisk in Table 1) were incubated at 28°C in 10 ml of water for 14 weeks. Every two weeks, a sample of each liquid culture was diluted and plated on YPD, and the viable cells that formed colonies were then counted. (v) *Maximum growth rate*. Microtiter plates with microcultures of 200 µl YPD media were incubated with agitation of 1.250 rpm (Tetramax 1000; Heidolph Instruments) and subjected to a wide-band optical density (OD) measurement every 20–60 minutes using the Infinite m200 Microplate Reader (Tecan). Before each OD measurement, parental strain populations were mixed using a pipette, which does not break small aggregates. The OD read-outs were log-transformed, and used to determine the range of exponential growth. The latter was done by finding lower and upper limits of OD between which the correlation coefficient (Pearson's *r*) was the highest when averaged over all cultures within

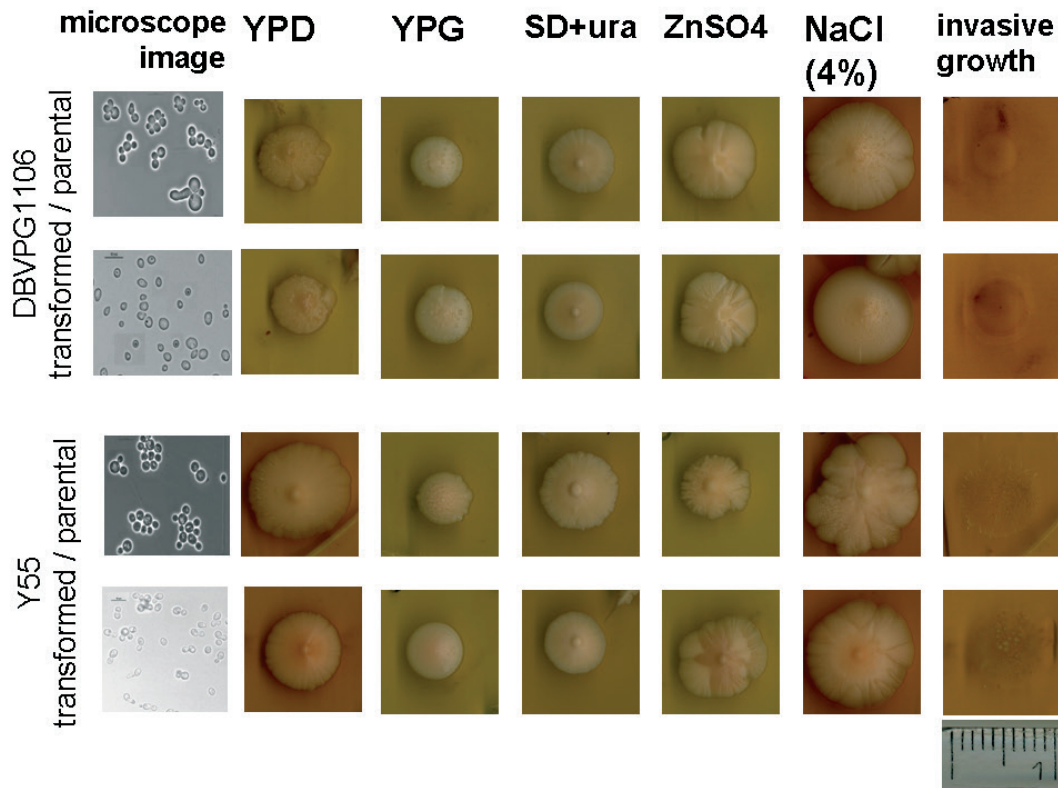


Figure 1. Scans of monoclonies of two strains that were chosen: DBVPG1106 and Y55. Small bars on the microscope images represent 10  $\mu$ m.

one test environment. This usually yielded no less than 5 time points with the mean squared  $r$  at 0.999 and rarely dropping below 0.99. The final estimator of fitness was the coefficient of slope in a linear regression. Every strain was assayed independently 6 times.

## RESULTS AND DISCUSSION

We obtained at least one clone from each strain (Table 1) that was positively verified by double sequencing to have the GTT (Val) at codon 368 of the *AMN1* gene, i.e., the W303 allele. Below are results from the five assays that those strains were subjected to:

(i) **Clumpiness.** We observed substantial reduction in the size and percentage of the aggregates in all of the transformed strains (Table 1). In the transformed strains, the highest number of cells counted in a clump was 5, while some aggregates in the non-transformed strain were composed of more than 24 cells. These results were similar irrespective of the media tested (YPD, SD+ura, ZnSO<sub>4</sub>, 0.5 mg/ml, NaCl, 4%), liquid and agar plates (stable environments) and temperature (22°C, 38°C and 35°C) (data not shown). At the same time, an increase in the number of single and double cells in a population of transformed strains was observed.

(ii) **Population density.** The transformation had no effect on the maximum average population density (ANOVA  $F_{1,46} = 3.55$ ;  $p = 0.07001$ ) which was measured using hemocytometer.

(iii) **Monocolony phenotype.** Most strains demonstrated no noticeable differences in the colony morphology. However, in minimal media (SD+ura) and under oxidative stress (ZnSO<sub>4</sub>; 0.5 mg/ml), parental colonies were more wrinkled and striated compared to the smoother surface of the transformed colonies (Fig. 1).

The “washing test” for invasive growth (Piccirillo & Honigberg, 2010) showed no difference in the phenotype for all strains tested on all the media (Fig. 1).

(iv) **Population viability.** The ratio of log N parental/transformed remained stable, suggesting that transformation did not influence viability (Regression  $r = 0.03$ ;  $F_{1,91} = 0.63$ ;  $p = 0.81$ ).

(v) **Maximum growth rate.** There was a significant difference between all but one parental and transformed strains (Table 1), with the largest difference of 1.2% (strain UWOPS83-787). The effect of transformation on growth rate varied for different strains.

In summary, we describe a method to eliminate or drastically diminish the clumpy aggregating phenotype in a yeast strain by changing a single codon in the *AMN1* gene. Several comparisons of the ancestral and *AMN1*\_W303 transformed strains show that this change has a minor influence on growth rate and colony morphology (in certain media). In our opinion these differences are not drastic. Such modification allows for using a growing number of strains isolated from various ecological origins and of different phylogeny distances for comparative studies, isolation of fraction of cells in a similar physiological state and research on origin of multicellularity. However, there is need for other assays to fully characterise the effect of codon replacement we propose here. We hope that this method and the modified strains could be useful for scientists interested in research using wild isolates of yeast *Saccharomyces cerevisiae*.

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