Pooled Segregant Sequencing Reveals Genetic Determinants of Yeast Pseudohyphal Growth

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Abstract

The pseudohyphal growth response is a dramatic morphological transition and presumed foraging mechanism wherein yeast cells form invasive and surface-spread multicellular filaments. Pseudohyphal growth has been studied extensively as a model of conserved signaling pathways controlling stress responses, cell morphogenesis, and fungal virulence in pathogenic fungi. The genetic contribution to pseudohyphal growth is extensive, with at least 500 genes required for filamentation; as such, pseudohyphal growth is a complex trait, and linkage analysis is a classical means to dissect the genetic basis of a complex phenotype. Here, we implemented linkage analysis by crossing each of two filamentous strains of Saccharomyces cerevisiae (Σ1278b and SK1) with an S288C-derived non-filamentous strain. We then assayed meiotic progeny for filamentation and mapped allelic linkage in pooled segregants by whole-genome sequencing. This analysis identified linkage in a cohort of genes, including the negative regulator SFL1, which we find contains a premature stop codon in the invasive SK1 background. The S288C allele of the polarity gene PEA2, encoding Leu409 rather than Met, is linked with non-invasion. In Σ1278b, the pea2-M409L mutation results in decreased invasive filamentation and elongation, diminished activity of a Kss1p MAPK pathway reporter, decreased unipolar budding, and diminished binding of the polarisome protein Spa2p. Variation between SK1 and S288C in the mitochondrial inner membrane protein Mdm32p at residues 182 and 262 impacts invasive growth and mitochondrial network structure. Collectively, this work identifies new determinants of pseudohyphal growth, while highlighting the coevolution of protein complexes and organelle structures within a given genome in specifying complex phenotypes.

Introduction

The budding yeast Saccharomyces cerevisiae undergoes a pronounced growth transition in response to nitrogen limitation or glucose limitation, forming multicellular pseudohyphal filaments that can spread outward from a colony and/or invade the surface of a solid growth substrate [1,2]. Yeast pseudohyphal filament formation is a presumed foraging mechanism, accomplished through underlying changes in cell adhesion, cell cycle progression, and budding [1,3,4]. During pseudohyphal growth, yeast cells remain physically connected after cytokinesis via mechanisms encompassing the actin cytoskeleton, regulation of polarity proteins, and delayed G2/M progression [8–12].

The molecular basis of yeast pseudohyphal growth has been studied extensively as a model of conserved signaling pathways controlling cell morphogenesis and polarity. Furthermore, related processes of filamentous development in the principal opportunistic human fungal pathogen Candida albicans are required for virulence, and signaling pathways between the related yeasts are conserved [13]. Classic studies of pseudohyphal growth in S. cerevisiae have resulted most prominently in the identification of core pseudohyphal growth signaling modules encompassing the Kss1p mitogen-activated protein kinase (MAPK) cascade, the cAMP-dependent protein kinase A (PKA) pathway, and the AMP-activated protein kinase ortholog Snf1p [14–20]. The pseudohyphal growth MAPK cascade encompasses Ste11p, Ste7p, and the MAPK Kss1p [10,14]. Kss1p phosphorylates the Ste12p transcription factor, resulting in dissociation of the negative regulatory Dig1p and Dig2p interactors and binding of a Ste12p-Tec1p heterodimer to target promoters, such as the FLO11 promoter [21–23]. Tpk2p, a catalytic subunit of PKA, phosphorylates the Flo8p transcription factor, promoting Flo8p binding and transcriptional activation at the FLO11 promoter and other regulatory sites [17,24–26]. In response to glucose limitation, FLO11 transcription is regulated by Suf1p; the Suf1p-Gal83p isoform promotes cell adhesion during invasive filamentation by antagonizing Nrg1p- and Nrg2p-mediated repression of FLO11 [19,27].

While the central components of these signaling pathways have been identified, the scope of the yeast pseudohyphal stress response is broad [28–33], and the mechanisms enabling these genes and gene products to drive pseudohyphal filamentation are
incompletely defined, as are the genetic determinants within this gene set that underlie filamentation. To further dissect pseudohyphal growth pathways, we implemented a linkage study, coupling whole genome sequencing with pooled segregant analysis. The results present previously unidentified genetic determinants of yeast invasive growth and indicate the coevolution of proteins within complexes in driving phenotype.

Results

Pooled Segregant Analysis of Yeast Invasive Filamentation with Deep Sequencing

For linkage analysis, we selected as parents the non-filamentous S288C-derived strain BY4741 and the filamentation-competent strains Σ1278b and SK1 [34,35]. Filamentous-form growth in haploid strains is classically assessed using the plate-washing assay of Gimeno et al. [1] to identify pseudohyphal cells that have invaded the agar substrate. The invasive phenotype of each parent strain in this assay is indicated in Fig. 1A. The experimental design of the linkage study is presented in Fig. 1B. The non-invasive S288C-derived strain was mated with each of the filamentous Σ1278b and SK1 strains, and the resulting diploid strain from each cross was sporulated. Meiotic progeny from dissected tetrads were assayed for agar invasion by plate-washing, and spores indicating strongly non-invasive or invasive phenotypes were pooled for subsequent linkage analysis. Only spores resulting from complete meiosis were included in these phenotypic pools, and intermediate filamentation phenotypes were excluded from subsequent analysis to provide the greatest likelihood of identifying allelic variation with a strong effect on filament formation. Genomic DNA was extracted from each segregant pool and subjected to high-throughput sequencing that yielded greater than 100-fold coverage per pool.

Identification of Allelic Variants Linked with Invasive Phenotype between a Non-Filamentous S288C Derivative and Σ1278b

From the BY4741-by-Σ1278b cross, 31 complete tetrads (124 spores) were screened for agar invasion, identifying 57 strongly invasive spores and 63 non-invasive spores (Fig 2A). The segregant pools were sequenced, and candidate determinants of the invasive phenotype were identified using a linkage LOD score of greater than 3 as an arbitrarily defined cut-off. Table S1 provides a listing of these alleles, encompassing only variants that are in protein-coding sequence and that are non-synonymous with respect to the encoded amino acid sequence. This allelic set affects 50 genes in eleven linkage blocks physically located on seven yeast chromosomes. Figure S1 summarizes the available functional information for this gene set.

Representative plots of non-synonymous allelic variation with respective LOD scores are graphed in Fig. 2B for chromosomes V and IX, highlighting the pseudohyphal growth transcription factor gene FLO8 and the flocculin effector gene FLO11. FLO8 is a pseudogene in S288C-derived strains [36], and in this analysis, the BY1741 allelic variant containing a premature translational stop at codon 142 of the FLO8 sequence yielded a LOD score greater than 17 (Fig. 2B and C). The FLO11 locus exhibits fifteen allelic changes linked with invasive growth phenotypes (Fig. 2B and C). Previous studies identified allelic variation in FLO11 sequence encoding amino- and carboxy-terminal regions linked with the ability to form biofilms on the surface of wine [37]. We recovered these as well as additional sites of DNA sequence variation in FLO11, with the Σ1278b-encoded alleles indicating linkage with strong invasive growth. The FLO11 sequence contains an internal repeat region that is a source of allelic variation between some strains and colonies [38,39]; however, we did not observe a change in the number of these repeats between BY4741 and Σ1278b. Collectively, the identification of these known pseudohyphal growth genes demonstrates the relevance of results obtained from our pooled segregant analysis.

PEA2 Allelic Variation Impacts Cell Morphology and MAPK Pathway Activity

To further identify important determinants of invasion, we screened candidates from Table S1 as follows: 1) we generated gene deletions and assayed for invasive growth phenotypes (Table S2), and 2) for genes yielding deletion phenotypes, we generated mutants with swapped alleles to identify genetic variants required for invasive growth in Σ1278b. In particular, we focused on alleles of genes that contributed to cell polarity, cell cycle progression, cell morphology, and cell responses to nitrogen/carbon limitation, as these are hallmark characteristics of filamentation.

By this approach, we identified variation in PEA2 as an important part of the genetics underpinning invasive growth. Pca2p localizes to sites of polarized growth as a component of a protein complex, termed the polarisome [40,41]. PEA2 is required for wild-type invasive growth, mating projection formation, and bipolar bud site selection in diploids [42,43]. In the filamentous Σ1278b strain, PEA2 codon 409 specifies methionine rather than the leucine residue encoded in the S288C-derived reference genome. The pca2-M409L allele was linked with invasive growth, and generation of an integrated site-specific mutation ( pca2-M409L) reconstituting the S288C-encoded PEA2 allele in Σ1278b resulted in decreased invasive growth (Fig 3A). Relative to wild type Σ1278b, the cell morphology of the pca2-M409L mutant is altered, exhibiting decreased elongation (Fig 3B); over a population of 200 cells, the percentage of pca2-M409L cells with a length/width ratio of less than 1.5 was nearly four-fold the corresponding percentage in a wild type strain. In addition, the pca2-M409L mutant is impaired in Kss1p MAPK signaling activity. The Kss1p kinase activates the Ste12p/Tec1p transcription factor complex, which recognizes a regulatory element (FRE) in the FLO11 promoter. The plasmid-based P<sub>B</sub>α<sub>111-9/10</sub>-lacZ
construct contains the Ste12p/Tec1p-responsive region of the FLO11 promoter fused to lacZ [6], and, by this reporter, the pea2-M409L mutant yields significantly decreased Ste12p/Tec1p-dependent transcriptional activation of FLO11 relative to wild-type Σ1278b (Fig. 3C). In contrast, the pea2-M409L mutation results in wild-type levels of a similarly designed FLO11 promoter fusion responsive to the PKA pathway effector Flo8p (Fig. 3C) [6].

Genetic Variation in PEA2 Modulates Bud Site Selection and Spa2p Binding

Under conditions of vegetative growth haploid yeast cells bud in an axial pattern, with new buds emerging adjacent to the preceding bud site [44]. Haploid cells undergoing pseudohyphal growth, however, adopt a predominantly unipolar budding pattern wherein the first bud forms distal to the original cell division site, and subsequent buds cluster in the distal pole [1,10]. Here, we find that in the Σ1278b background the pea2-M409L mutant, corresponding to the S288C-encoded PEA2 allele, exhibited a decrease in unipolar budding and an increase in axial budding relative to wild-type (p < 0.001), with levels intermediate between an otherwise isogenic wild-type strain and a pea2Δ mutant (Fig. 3D). For this analysis, we examined a population of invasive cells exhibiting three or more bud scars, such that patterns of axial, unipolar, bipolar, and random budding could be reliably distinguished [44,45]. This budding phenotype was evident in invasive cells, but not in an equally sized population of cells scraped from the surface of an agar plate.

Previous studies have indicated that the majority of bud sites are distal in a pea2Δ mutant [12]; results here also indicate that the majority of bud sites are distal in pea2 mutants, but budding pattern analysis does indicate that Pea2p residue 409 impacts unipolar budding in invasive haploid cells.

In the polarisome complex, Pea2p binds the scaffolding protein Spa2p, a large coiled-coil domain-containing protein required for polarisome function [41,46]. Here, we assessed the possibility that allelic variation at the PEA2 locus impacts Spa2p binding, using Protein A (ProA)-tagged Pea2p variants to recover by co-immunoprecipitation Spa2p tagged at its amino terminus with the hemagglutinin (HA) epitope. In the Σ1278b strain, the Pea2p-M409-ProA variant recovered more HA-Spa2p than the Pea2p-L409-ProA variant, representing the S288C-encoded PEA2 allele (Fig. 3E).

Analysis of Genetic Variation Linked with Invasion between BY4741 and SK1

The BY4741-by-SK1 cross was implemented as described in Experimental Procedures, and phenotypic analysis of meiotic progeny identified 51 and 24 strongly invasive and non-invasive spores, respectively (Fig. 4A). Subsequent deep-sequencing of the phenotypic pools identified allelic variation linked with invasive growth phenotypes within eleven separated locus blocks encompassing 88 genes exhibiting non-synonymous changes and a LOD score of greater than 4 (Table S3). A functional breakdown of these genes is indicated in Figure S2. In this analysis, we used a higher LOD score relative to the Σ1278b-by-BY4741 cross in order to limit the number of selected allelic variants to a manageable size for further study, as the SK1 and BY4741 genomes are more divergent (99.5% sequence identity) than the Σ1278b and BY4741 genomes (99.7% identity) [47]. Very few allelic variants linked with invasive growth in Σ1278b were also identified in SK1, aside from a few sequences near FLO8 that are unlikely to be causative. The set of identified alleles was primarily distinct between the two linkage studies, and deletion phenotypes for tested genes in SK1 are indicated in Table S4.

In haploid spores from this cross, genetic variation is most strongly linked with the invasive growth phenotype over a region of roughly 80,000 bp on chromosome XV, encompassing SFL1 (Fig. 4B and C). SFL1 encodes a transcriptional repressor of pseudohyphal growth that functions by binding to the FLO11 promoter, thereby blocking transcriptional activation [24,25,48–51]. Consistent with its function in repressing FLO gene expression, deletion of SFL1 results in exaggerated invasive growth [28,32,49]. Interestingly, the SK1 strain contains an allelic variant of SFL1 with respect to S288C-derived strains, resulting in the conversion of codon 477 (CAA encoding glutamine) to a TAA stop codon (Fig. 4D). This premature stop codon truncates SFL1 prior to the sequence encoding a domain (AA 571–658) that is strongly similar to a conserved region in Muc oncoproteins [48]. Previous studies have identified hyperactive filamentation in a mutant of the CEN.PK 113-7D background upon introduction of a premature translational stop at SFL1 codon 320 (Q320-stop) [51]. Here, we found that allelic variation in SFL1, encompassing a premature stop codon (C1430T, Q477-stop) in the SK1 background, is linked to the aggressively invasive phenotype of SK1 relative to BY4741.

Allelic Variation in MDM32 Affects Mitochondrial Morphology and Invasive Growth

In the BY4741-by-SK1 cross, allelic variation in MDM32 was linked to invasive growth more strongly than any other identified locus, with a LOD score of 9. As indicated in Fig. 4B-D, MDM32 is found on chromosome XV, and relative to BY4741, the SK1 allele of MDM32 encodes Ser182 and Phe262 rather than Cys and Leu, respectively. MDM32 encodes a protein complex...
subunit of the mitochondrial inner membrane required for membrane organization, the maintenance of elongated mitochondrial morphology, and mitochondrial DNA nucleoid stabilization [52]. Mitochondrial function is required for pseudohyphal growth, as filamentation-competent strains of *S. cerevisiae* containing a deleted version of the mitochondrial genome are unable to form pseudohyphae [53,54]; however, a role for Mdm32p in enabling invasive growth remains to be identified.

To determine the effect of *MDM32* allelic variation on pseudohyphal growth, we replaced the SK1-encoded *MDM32*-C546/T787 allele, specifying Mdm32p-S182/F262, with BY4741-encoded *MDM32*-G546/A787, specifying Mdm32p-C182/L262, in the SK1 genetic background. This allelic swap decreased invasive growth in SK1, and agar invasion was rescued upon reintroduction of the native SK1-encoded *MDM32* allele (Fig. 5A). The SK1 mutant containing the BY4741-encoded allele of *MDM32* exhibited a more rounded cell morphology, with the percentage of cells displaying a cell length:width ratio of less than 1.5 increasing from 17% in wild-type SK1 cells to 68% in the SK1 mutant (Fig. 5B). Reintroduction of SK1-encoded *MDM32* recovered levels of cell elongation similar to wild type.
To assess the impact of this allelic variation on mitochondrial function, we grew the SK1 mutant with the BY4741 allele of MDM32 on medium containing non-fermentable glycerol as the sole carbon source. As shown in Fig. 5C, the allele-swapped SK1 mutant grows poorly in glycerol-containing media, indicating that oxidative phosphorylation is impaired. The structure of the mitochondrial network is also perturbed upon introduction of the BY4741 allele of MDM32 in the SK1 background. Using the mitochondrion-specific MitoTracker fluorescent dye, which diffuses passively across the plasma membrane and concentrates in active mitochondria by membrane potential, we can visualize a compact and collapsed mitochondrial network in SK1 cells containing the BY4741-encoded Mdm32p-C182/L262 variant, similar to that observed in mdm32Δ (Fig. 5D). MDM32 is a paralog of MDM31, and the encoded proteins have been found to interact, albeit transiently and weakly, as components of protein complexes at the mitochondrial inner membrane [52]. We, therefore, assessed the effect of allelic variation at MDM32 on Mdm31p binding; however, we observed no difference in the recovery of Mdm31p by co-immunoprecipitation between the respective Mdm32p variants (Figure S3). In sum, MDM32 is a determinant of invasive growth, and replacement of the native SK1 allele of MDM32 with the BY4741-encoded allele yields a mutant filamentous growth phenotype.

Figure 3. Variation at Pea2p residue 409 impacts invasive growth, budding, and polarisome interactions. A) Invasive growth of the wild-type Σ1278b strain, pea2Δ strain, and a strain containing the integrated pea2-M409L point mutation in the Σ1278b background was assayed by plate washing (++, strong invasive growth; +, invasive growth; −, lack of invasive growth). Invasive growth levels were quantified by determining the ratio of pixel intensity of images of the spotted cultures before and after plate washing; these ratios are indicated to the lower right of each spotted culture image post-washing. B) Cell morphology was assessed by determining the length-to-width ratio of 200 cells each for the indicated strains. Prior to imaging, cells grown on YPD medium were scraped into a suspension in liquid media. For length-width calculations, the longest cell axis was selected for cell length measurements. C) Activity of MAPK and PKA signaling pathways in pea2 mutants. The bar graphs indicate activity of plasmid-based cAMP-dependent PKA and pseudohyphal growth MAPK pathway-responsive P_flo11-lacZ reporters. The PKA pathway-responsive P_flo11 reporter contains a fragment of the FLO11 promoter (P_flo11-6/7) 21.0–21.4 kb from the FLO11 start codon, and the Kss1p MAPK-responsive P_flo11 reporter contains a promoter fragment (P_flo11-9/10) 21.6–22.0 kb upstream of the FLO11 start codon. β-galactosidase activity was measured from cells growing exponentially in SC-Ura medium. Error bars indicate standard deviations of three independent measurements. D) Budding patterns were analyzed for the indicated strains by calcofluor white staining of invasive cells recovered from the agar. Sample images of the observed budding patterns are presented. Budding patterns of the pea2Δ and pea2-M409L mutants are distinct from wild type, with p-values of <0.01 and <0.001, respectively, by Pearson’s chi-squared test. E) Western blots indicate co-immunoprecipitation of Spa2p by Pea2p variants in the Σ1278b background. For this analysis, chromosomal PEA2 and pea2-M409L were tagged at the carboxy terminus by integration of a cassette encoding ProA. Spa2p tagged at its amino terminus with the hemagglutinin (HA) epitope was expressed from a plasmid using a copper-inducible promoter. The binding affinity between Pea2p and Spa2p was revealed by IgG pull-down using Pea2p-ProA as bait. Pea2p and Spa2p levels were normalized against levels observed in total cell lysate. The ratio of recovered Spa2p by the S288C-encoded Pea2p-L409 variant is presented relative to the ratio of Spa2p recovered by Σ1278b-encoded Pea2p-M409; thus, the boxed ratios indicate that Pea2p-L409 recovered 60% the amount of Spa2p co-immunoprecipitated with Pea2p-M409.

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The linkage analysis presented here identifies a broad gene set contributing to pseudohyphal growth (Fig. 6). The pattern of allelic linkage indicates a large number of determinant loci underlying invasive growth, consistent with results from systematic single-gene deletion and overexpression studies [28,31,32]. Within this gene set, components of the polarisome and mitochondria play important roles in enabling invasive growth. We report here that Pea2p residue 409 impacts bud site selection in haploid cells undergoing invasive growth, although the effect is less pronounced in determining initial distal-versus-proximal budding in virgin mother cells. Pea2p residue 409 lies distinct from the Pea2p coiled-coil region between residues 236 and 327 and is important for Spa2p binding. Spa2p interacts with Ste11p and Ste7p from the Kss1p MAPK pathway, providing a mechanism for polarisome-mediated regulation of Kss1p MAPK activity [41]. Our results further indicate that nuclear-encoded Mdm32p is required for invasive growth in SK1, and that residues 182 and 262, located outside of the mitochondrial pre-sequence (AA 1–102) and at the boundary or outside of a transmembrane domain (AA 161–184 and 636–653), are important in enabling invasion, as well as in determining aerobic respiratory function and mitochondrial morphology. Mdm32p is proposed to function cooperatively with other inner membrane proteins and components of the outer mitochondrial membrane in the maintenance of mitochondrial morphology [52], potentially through cytoskeletal interactions that may be affected by variation at these sites.

**Discussion**

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This analysis highlights two additional points. First, the non-filamentous BY4741 background is not uniformly repressive with respect to pseudohyphal growth. From the Σ1278b and SK1 crosses, we identified three and five blocks of allelic variation,
respectively, in BY4741 linked with the invasive growth phenotype; full listings of the encompassed alleles with respect to each cross are presented in Tables S5 and S6. Alleles in S288C-derived strains that promote pseudohyphal growth are antagonized by alleles that repress filament formation, such as the pseudogene form of FLO8; similarly, alleles in SK1 linked with the non-invasive phenotype may be offset by alleles that promote invasion, such as the SFL1 allele containing a premature stop codon.

Second, the identified allelic variation in PEA2 and MDM32 and the allele-swapping experiments performed here indicate that within a given genome, functionally interacting genes coevolve to impact phenotype. The majority of genetic variation linked with invasive phenotype in this study involves site-specific changes that do not create pseudogenes. Alleles of these genes yield functional proteins within the respective genomic contexts; however, a given allele results in a hypomorphic phenotype when introduced into a non-native strain. It should be noted that the BY4741-encoded allele of PEA2 may be viewed as being pseudohyphal competent, as Liu et al. [36] reported that the introduction of S1278b-encoded FLO8 in a S288C-derived strain is sufficient to enable at least some degree of pseudohyphal growth. The data here suggest that partner genes have likely co-evolved with genes such as PEA2 and MDM32, and the resulting protein complexes are, thus, an important determinant of cell phenotype. These findings highlight the utility in studying these complexes as a whole, in supplement to individual proteins, in order to accurately identify the functions and properties that specify phenotype.

It is interesting that the studies here indicated very little overlap between alleles linked with invasive growth in the S1278b and SK1 strains with respect to BY4741 (Fig. 6). Studies mapping quantitative trait loci (QTL) in a cross of the laboratory strain BY4716 and the vineyard strain RM-11 identified hotspots impacting gene expression, protein abundance, and small molecule-dependence [55–58]. These hotspots were principally due to alleles in the BY4716 background, leading Ronald and Akey [59] to suggest that the causative polymorphisms may occur at low frequency. The non-overlapping allele sets identified in our analysis are not suggestive of hotspots, but rather highlights the substantial importance of epistatic interactions in determining the sum filamentous phenotype resulting from variant alleles in the haploid segregants. These epistatic interactions likely represent instances of gene coevolution, which has been suggested to occur at an elevated rate for genes encoding proteins of shared biological functions and/or for proteins that have coevolved between species [60,61]. Clark et al. [60] have analyzed the rate of covariation for pairs of proteins over evolutionary time, and by this analysis,
polarisome components as a whole do not exhibit statistically significant evidence of covariation, although many mitochondrial complexes do yield signature indicating evolutionary rate covariation. Further analyses of individual protein pairs from the strains used in our study will be necessary to identify a set of coevolved proteins that drive the filamentous growth phenotype.

In summary, we used pooled segregant whole-genome sequencing to dissect gene networks that determine yeast pseudohyphal growth. This analysis identified allelic variation in the known pseudohyphal growth genes \textit{FLO8} and \textit{FLO11}, while also revealing variation in the negative regulator \textit{SFL1}, the coding sequence of which contains a premature stop codon in the invasive SK1 background. We further found that amino acid 409 in the polarisome protein Pea2p is a site of allelic variation critical for the protein’s ability to signal through the Kss1p MAPK pathway, establish unipolar budding during pseudohyphal growth, and bind the Spa2p polarisome scaffold. Linkage analysis identifies variation in \textit{MDM32} as a determinant of invasive growth between S288C derivatives and the SK1 strain; the 182 and 262 residues are sites of variation and contribute to Mdm32 function in aerobic respiration and invasive growth.

Materials and Methods

Yeast Strains and Growth Conditions

A listing of yeast strains and plasmids used in this study is provided in Tables S7 and S8. Haploid deletion mutants were constructed by PCR-mediated gene disruption using pFA6a-KanMX6 or pUG72 \cite{62,63}. Yeast strains were propagated on rich YPD medium (1% yeast extract, 2% polypeptone and 2% glucose) medium or synthetic medium as described \cite{64}. Yeast invasive growth was assayed on YPD medium.

Figure 6. Chromosomal location of genes exhibiting allelic linkage with invasive growth from the BY4741 cross with Σ1278b and SK1, respectively. Genes within a chromosomal region are listed in the boxes with color-coding to indicate the respective cross from which the linkage was identified. Hash marks indicate 100,000 bases, and the full length of each chromosome is indicated.

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Pooled Segregant Analysis

The statistical modeling used to derive the probabilities of identifying linkage are described by Birkeland et al. [65]. Following mating as indicated in Fig. 1, resulting strains were sporulated and asci were dissected. The dissected spores were grown overnight at 30°C and were individually tested for mating type. Spores resulting from complete meiosis (four viable spores with two each of the α and a mating types) were then used for whole genome sequencing. Each spore was assigned to invasive or non-invasive pools based on its invasive growth phenotype. To ensure equal representation of all segregants in a pooled population, each haploid strain was grown overnight at 30°C in individual 4 ml YPD cultures. The OD600 of the cultures was determined and used to calculate the appropriate volume of each strain so that upon mixing, we would achieve equal numbers of cells. A mate-pair library with 300-bp fragments was prepared for each of the phenotypic pools, and each library sequenced as paired-end reads using the Illumina Genome Analyzer (University of Michigan DNA Sequencing Core). Sequence analysis was performed as described [65]. To obtain an estimate of the recombinant and non-recombinant spore counts in each phenotypic pool for a given observed sequence variant, we multiplied the number of spores in the pool by the fraction of sequence reads from that pool that matched the corresponding allele variant. These values were then used in standard LOD score calculations.

Analysis of Yeast Budding Pattern

Budding patterns of invasive cells were determined as previously described [44,45]. In brief, equal concentrations of mid-log phase cultures were spotted onto YPD plates and incubated for 7 days at 30°C; surface cells were subsequently washed off under a gentle stream of water. Residual invaded cells were recovered from the agar using a sterile toothpick. Cells were washed twice in sterile water and were stained with 2 μg/ml calcoflour white. Bud scars were visualized by fluorescence microscopy. Cells with more than three bud scars were examined. Budding patterns were determined by criteria previously described [45]. Budding patterns were divided into four subgroups: axial, bipolar, unipolar and random. The axial pattern was defined as a long chain of bud scars on the proximal cell pole. Cells with a cluster of scars exclusively at the distal pole were classified as exhibiting unipolar budding. A pattern of medial bud scars was scored as random budding, whereas cells with bud scars equally distributed on both proximal and distal poles were classified as undergoing bipolar budding. For these analyses, 200–250 cells from each strain were scored.

Plasmid construction. For expression of Σ1278b alleles in null mutants, yeast open reading frames of the Y826 background with 1 kb of upstream sequence and 300 bp of downstream sequence were cloned into a Gateway vector by standard methods of recombination-based Gateway cloning (Invitrogen) [66,67]. For fusing enhanced green fluorescent protein (eGFP) and tandem affinity purification (TAP) cassettes to the carboxy-terminus of the budding yeast Mdm31p and Mdm32p proteins, Gateway plasmids 416-GPD-ccdB-EGFP and 414-GPD-ccdB-TAP were used [68]. Plasmid pCu-Spa2p-3xHA-URA3 was modified from pCu-3xHA-URA3 using standard restriction enzyme digestion and ligation-based techniques. The Sall and HindIII restriction sites of this plasmid was used to integrate the SPA2 open reading frame between the copper-inducible CUP1 promoter and 3xHA/GFP tag.

Integrated point mutations. Allelic variants of PEA2 were generated in the Σ1278b background as integrated point mutations.

A URA3 cassette was amplified from the pUG6 plasmid and then used to replace the PEA2 open reading frame in Y826. Subsequently, 5-FOA-mediated counter selection [69] was applied to replace the integrated URA3 cassette with a 500 bp cassette encompassing the PEA2 allelic change (M409L), recreating the 2080C-derived allele in the Σ1278b background. The integrated point mutation was confirmed by sequencing of an amplified PCR fragment using the University of Michigan Sequencing Core.

Yeast invasive growth assays. Invasive growth of haploid strains was determined by the standard plate washing assay of Gimeno et al. [1]. Mid-log phase cultures were spotted onto YPD plates and incubated for 7 days at 30°C, and surface cells were subsequently washed off under a gentle stream of water. Residual cells from the spotted culture on the plate were imaged using a Nikon Eclipse 80i upright fluorescence microscope.

FRE-lacZ assays. Plasmids pLG669-Z FLO11-6/7 and -9/10 were transformed into designated yeast strains by standard protocols [64]. Cells were grown in SC-Ura media overnight, and then inoculated into fresh media to an OD600 of approximately 0.2. Cultures were subsequently grown for 3–4 hours to an OD600 of approximately 0.8–1.0. β-galactosidase activity was determined with the use of ortho-nitrophenyl-β-galactoside as a substrate (Sigma Aldrich) [70,71].

Assays for respiratory activity. Respiratory ability was assessed using YPG plates with glycerol as a non-fermentable carbon source. Cells were pre-grown overnight and then inoculated into fresh SC medium to an OD600 of approximately 0.2; culture was subsequently incubated an additional 3 hours to an OD600 of approximately 0.8. The same amount of cells were serially diluted and spotted onto YPD and YPG [1% yeast extract, 2% polypeptone and 2% glycerol] plates, and incubated for 2 days at 30°C.

Fluorescence microscopy and image processing. For live-cell imaging, spotted cultures were grown in YPD plates for 7 days at 30°C. After washing the surface of the plates with a gentle stream of water, cells that had invaded the agar were recovered using a steel scalpel, and the extracted cells were washed twice with distilled water. Images were taken using a Nikon Eclipse 80i upright fluorescence microscope. To determine budding patterns, cells were stained with Calcofluor White as previously described [45]. Over 250 cells were counted twice for each strain. Statistics were analyzed using the ANOVA approach. To reveal mitochondrial morphology, living cells were stained with 50 nM MitoTracker Red, and the staining pattern was visualized by fluorescence microscopy [72].

Immunoprecipitation studies. PEA2 alleles were generated by tagging with ProA on the C-terminus in wild type and pea2- M490L mutants in the Σ1278b strain. The SPA2 open reading frame was cloned into pCu-3HA-URA, such that SPA2 was expressed from a copper-inducible promoter as a fusion protein with three copies of the hemagglutinin epitope at its amino terminus. The resulting plasmids pCu-3HA-ScSPA2 and pCu-3HA-ScSPA2 were introduced by transformation into both wild type and pea2- M490L mutants. The binding affinity between selected proteins was detected by co-immunoprecipitation. For native immunoprecipitation, 2 OD units of cells were lysed in 1 ml lysis buffer (50 mM Tris-HCl [pH 7.5], 150 mM NaCl, 2 mM EDTA, 0.5% Triton X-100, 1 mM PMSF, and Complete EDTA-free protease inhibitor [Roche]) with glass beads. After centrifugation at 15,000 g for 10 min, the resulting supernatant was incubated with protein G-Sepharose 4 Fast Flow (GE Life Tech) for 2 hr at 4°C. After washing the Sepharose with lysis buffer six times, the bound materials were eluted by boiling the Sepharose in SDS-PAGE loading buffer. The resulting eluate was analyzed by Western blotting with designated antibodies (anti-HA, anti-Protein
A and anti-GFP). Blots were developed using the SuperSignal West Dura Extended Duration Substrate (Thermo Scientific).

Supporting Information

Figure S1  Functional characterization of genes exhibiting allelic linkage with invasive growth in the BY4741 cross with Σ1278b. Genes from the analysis that have been previously identified as exhibiting filamentous growth phenotypes upon deletion [32] are listed to the left. The remaining genes are listed to the right, with genes of unknown functions and genes with functions that may be related to pseudohyphal growth indicated separately. Functions are drawn from data deposited in the Saccharomyces Genome Database as of publication. The chromosome in which the gene is located is indicated in parentheses. It should be noted that this list encompasses genes that may only exhibit linkage because of their close proximity to an important allelic determinant.

(TIF)

Figure S2  Functional characterization of genes exhibiting allelic linkage with invasive growth in the BY4741 cross with SK1. Genes from the analysis that were identified as exhibiting filamentous growth phenotypes upon deletion [32] are listed to the left. The remaining genes are listed to the right, with genes of unknown functions and genes with functions that may be related to pseudohyphal growth indicated separately. Functions are drawn from data deposited in the Saccharomyces Genome Database as of publication. The chromosome in which the gene is located is indicated in parentheses. It should be noted that this list encompasses genes that may only exhibit linkage because of their close proximity to an important allelic determinant.

(TIF)

Figure S3  Binding of Mdm31p by BY4741- and SK1-encoded Mdm32p variants. The MDM32 open reading frames from SK1 and BY4741 were cloned into a GPD-cGFP Gateway plasmid, such that the MDM32 sequence was expressed from the GPD1 promoter as an in-frame 3′-fusion to sequence encoding enhanced GFP. MDM31 was cloned into the GPD-TAP plasmid, yielding a fusion of the tandem affinity purification (TAP) tag to the carboxy terminus of Mdm31p upon expression from the GPD1 promoter. The resulting plasmids were transformed into ndm32A mutants. The binding affinity between Mdm32p-GFP and Mdm31p-TAP was revealed by IgG pull-down using Mdm31p-TAP as bait. No significant difference in binding between Mdm31p-TAP and the respective Mdm32p-GFP variants was observed.

(TIF)

Table S1  Genes exhibiting allelic linkage (LOD≥3) from the BY4741-by-Σ1278b cross. Alleles in a single chromosome are demarcated by a double line; groupings of alleles within a possible linkage block are separated by a single line.

(DOCX)

Table S2  Genes assayed for deletion phenotypes with respect to invasive growth in Σ1278b. Deletion phenotypes were assessed using standard plate-washing assays.

(DOCX)

Table S3  Genes exhibiting allelic linkage (LOD>4) from the BY4741-by-SK1 cross. Alleles grouped in a single chromosome are indicated by a double line; groupings of alleles within a single linkage block are indicated with a single line.

(DOCX)

Table S4  Genes assayed for deletion phenotypes with respect to invasive growth in SK1. Deletion phenotypes were assessed for invasive growth by plate-washing assays.

(DOCX)

Table S5  Alleles with variation between BY4741 and SK1 where the BY4741-encoded allele exhibits linkage with the invasive phenotype. Alleles within a given chromosome are separated by a double line.

(DOCX)

Table S6  Alleles with variation between BY4741 and SK1 where the BY4741-encoded allele exhibits linkage with the invasive phenotype. Alleles within a given chromosome are separated by a double line.

(DOCX)

Table S7  Strains used in this study.

(DOCX)

Table S8  Plasmids used in this study.

(DOCX)

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Author Contributions

Conceived and designed the experiments: QS CJ TEW AK. Performed the experiments: QS CJ. Analyzed the data: QS CJ TEW AK. Contributed reagents/materials/analysis tools: QS CJ. Contributed to the writing of the manuscript: QS CJ TEW AK.

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