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# Population and comparative genetics of thermotolerance divergence between yeast species

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#### Abstract

Many familiar traits in the natural world—from lions' manes to the longevity of bristlecone pine trees—arose in the distant past, and have long since fixed in their respective species. A key challenge in evolutionary genetics is to figure out how and why species-defining traits have come to be. We used the thermotolerance growth advantage of the yeast *Saccharomyces cerevisiae* over its sister species *Saccharomyces paradoxus* as a model for addressing these questions. Analyzing loci at which the *S. cerevisiae* allele promotes thermotolerance, we detected robust evidence for positive selection, including amino acid divergence between the species and conservation within *S. cerevisiae* populations. Because such signatures were particularly strong at the chromosome segregation gene *ESP1*, we used this locus as a case study for focused mechanistic follow-up. Experiments revealed that, in culture at high temperature, the *S. paradoxus ESP1* allele conferred a qualitative defect in biomass accumulation and cell division relative to the *S. cerevisiae* allele. Only genetic divergence in the *ESP1* coding region mattered phenotypically, with no functional impact detectable from the promoter. Our data support a model in which an ancient ancestor of *S. cerevisiae*, under selection to boost viability at high temperature, acquired amino acid variants at *ESP1* and many other loci, which have been constrained since then. Complex adaptations of this type hold promise as a paradigm for interspecies genetics, especially in deeply diverged traits that may have taken millions of years to evolve.

Keywords: evolution; genetics; Saccharomyces; thermotolerance; adaptation; ancient

### Introduction

A central goal of research in evolutionary genetics is to understand how new traits are built. Much of the literature to date focuses on adaptive trait innovation within a species, in the wild (Chan et al. 2010; Will et al. 2010; Linnen et al. 2013; Cleves et al. 2014; Field et al. 2016; Asgari et al. 2020) and in the lab (Blount et al. 2012; Tenaillon et al. 2016; Good et al. 2017; Castro et al. 2019). These systems have enabled studies of short-term adaptation, its genetics (Barroso-Batista et al. 2014; Garud et al. 2015; Good et al. 2017; Harris et al. 2018; Castro et al. 2019; Xie et al. 2019), and its dynamics (Toprak et al. 2011; Blount et al. 2012). Such work on recent adaptations serves as a backdrop for the study of evolution over longer timescales. Many familiar traits from the natural world have been acquired over millions of generations. In the modern-day, such characters manifest as differences between deeply diverged, reproductively isolated lineages. They can represent the abiding fitness strategies of their respective species, and are thus of particular interest in the field. But their evolutionary mechanisms pose a key challenge, given that the relevant events happened so long ago. For these ancient traits, candidate-gene studies have implicated individual loci (Baldwin *et al.* 2014; Anderson *et al.* 2016; Massey and Wittkopp 2016; Sulak *et al.* 2016; Li and Fay 2017; Liu *et al.* 2018; Sackton *et al.* 2019; Tian *et al.* 2019) and reconstructed the mutational path by which a given determinant evolved (Bridgham *et al.* 2009; Finnigan *et al.* 2012; Anderson *et al.* 2015; Liu *et al.* 2018; Pillai *et al.* 2020). Even in such landmark cases, the tempo and mode of evolution of deep trait divergences have remained largely out of reach. To meet the latter challenge, one would need to trace the rise of causal alleles in the respective species and the selective forces that drove it, and pinpoint the timing of these events.

In prior work, our group mapped multiple housekeeping genes underlying the difference in thermotolerance between *Saccharomyces cerevisiae* and other species in its clade, and found that *S. cerevisiae* harbored derived alleles at these loci (Weiss *et al.* 2018). Here, we set out to investigate when and how *S. cerevisiae* 

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acquired the putatively adaptive determinants of thermotolerance, using a population-genomic approach. We then used the results as a jumping-off point for additional analyses of the molecular mechanisms by which variants at thermotolerance genes confer their effects.

### Materials and methods Identifying thermotolerance genes

To identify genes at which variation between S. cerevisiae and S. paradoxus impacts thermotolerance, we re-analyzed data from a reciprocal hemizygosity screen of transposon mutants in the interspecies hybrid background (Weiss et al. 2018) as described, with the following differences. Call a<sub>39, i</sub> the average, across technical replicates, of sequencing-based abundances of a hemizygote mutant measured after ~7 generations in biological replicate i of growth at 39°C, and  $a_{\rm 28,\ i}$  the analogous quantity for growth at  $28^{\circ}$ C, for i = [1, 3]. We calculated the mean of the latter across biological replicates, a<sub>28, mean</sub>, and then used it to tabulate three replicate estimates of the temperature effect on growth of the mutant as log<sub>2</sub>(a<sub>39, i</sub>/a<sub>28, mean</sub>). If the coefficient of variation across these biological replicates was greater than 20, we eliminated the mutant from further consideration. Otherwise, for a given gene, we concatenated these vectors of length three across all hemizygote mutants in the S. paradoxus allele for which we had abundance data, yielding the set of temperature effects  $s_{\mbox{\scriptsize Spar}}.$  We did likewise for the S. cerevisiae allele, yielding s<sub>Scer</sub>. We retained for further analysis only genes at which we had at least two mutants' worth of data for each allele. For each such gene, we compared  $s_{Scer}$  and s<sub>Spar</sub> with a Wilcoxon test, and corrected for multiple testing across genes, as described in Weiss et al. (2018).

### Sequence data, alignments, and interspecies diversity

For  $D_{xy}$  analyses in Supplementary Table S2, for a given gene, open reading frame sequences for the strains of each *S. cerevisiae* population from Peter *et al.* (2018) were aligned against the European *S. paradoxus* population from Bergström *et al.* (2014) and, separately, against the North American *S. paradoxus* subpopulation B from Durand *et al.* (2019). For  $D_{xy}$  analysis across species in Table 1, alignments were generated using all *S. cerevisiae* and *S. paradoxus* strains. Alignments used MUSCLE (Edgar 2004) with the default settings for DNA and –maxiters set to 2. Any gene for which, in the alignment, >10% of sites were denoted as gaps or unknown nucleotides (Ns), or sequences from <75% of strains in the population were available, was eliminated from analysis, leaving 4110 to 4781 genes suitable for testing in the respective strain set.

We calculated pairwise nucleotide diversity  $\left( D_{xy} \right)$  for each gene as

$$D_{xy} = \frac{1}{n_x n_y} \sum_{i=1}^{n_x} \sum_{j=1}^{n_y} d_i$$

where  $n_x$  is the number of *S. cerevisiae* strains,  $n_y$  is the number of *S. paradoxus* strains, and *d* is the number of sites with nucleotide differences at the same position for each pairwise sequence comparison. Sites with gaps or unknown nucleotides (Ns) were ignored.

To test for enriched  $D_{xy}$  among thermotolerance genes in a given *S. cerevisiae* and *S. paradoxus* population pair or across the species, we first tabulated  $D_{\text{therm}}$ , the median  $D_{xy}$  across the thermotolerance gene cohort from the appropriate genomes. We next

sampled 10,000 random cohorts of genes from the genome with the same number of essential and nonessential genes as in the thermotolerance cohort (Winzeler *et al.* 1999), and tabulated the median  $D_{xy}$  in each  $D_{rand}$  from the appropriate genomes. We used as an empirical *P*-value the proportion of random cohorts with  $D_{rand} \ge D_{therm}$ .

### Codon alignment and McDonald-Kreitman statistics

Open reading frame sequences for each *S. cerevisiae* strain from Peter *et al.* (2018), the European *S. paradoxus* strains (Bergström *et al.* 2014), and North American *S. paradoxus* strains (Durand *et al.* 2019) were translated to amino acid sequences using Biopython (Cock *et al.* 2009) and aligned using MUSCLE (Edgar 2004) with default settings for amino acids and –maxiters set to 2. The amino acid sequence alignments and unaligned nucleotide sequences were used as input to PAL2NAL (Suyama *et al.* 2006) to create codon alignments for each gene. Sequences with stop codons within the open reading frame or where >10% of sites were denoted as gaps or unknown Ns were discarded. Genes with valid sequences in <75% of strains from each species were removed from the analysis, leaving 3814 genes suitable for testing.

The codon alignments were input into the CodonAlignment module of Biopython 1.78 (Cock *et al.* 2009) and the mktest function reported the number of divergent nonsynonymous ( $D_n$ ), divergent synonymous ( $D_s$ ), polymorphic nonsynonymous ( $P_n$ ), and polymorphic synonymous changes ( $P_s$ ) in each gene. We calculated the Neutrality Index (NI) for each gene as

$$\mathrm{NI} = \frac{P_n/P_s}{D_{n/D_s}}$$

(McDonald and Kreitman 1991). We then used these measures as input into a resampling test for enrichment of low NI, analogous to that used for  $D_{xy}$  (see above).

# Multi-locus genotype and allele-sharing inference in S. cerevisiae populations

We calculated expected genotype homozygosity, G1 (Harris *et al.* 2018), as follows.

For the allele-sharing inference across all S. cerevisiae in Table 1, we used unphased VCF genotypes for all strains from 1011 Yeast Genomes (Peter *et al.* 2018) as input into SelectionHapStats (Harris *et al.* 2018) with the following parameters: -w (window size, SNPs) = 1200, and -j (jump size, SNPs) = 25. We tabulated G1 in each window whose center fell within the open reading frame, and we calculated the average across the windows. We then used these measures as input into a resampling test for enrichment of high G1 analogous to that used for

**Table 1** Thermotolerance loci are enriched for positive selection

 between species and allele-sharing in S. cerevisiae

	Thermotolerance genes	Genome	Р
D <sub>xy</sub> <sup>a</sup> NI <sup>b</sup>	0.1215 2.0513	0.1016 2.8076	0.0053* 0.0217*
G1 <sup>c</sup>	0.01025	0.0077	0.0491*

 $^{\rm a}$  The columns report the median value of the absolute diversity statistic  $D_{xy}$  in thermotolerance genes and random sets of genes from the genome, respectively, and empirical significance from a resampling test.

<sup>b</sup> Data are as in *a* except that the metric analyzed was the neutrality index NI according to the McDonald-Kreitman test.

 $^{\rm c}$   $\,$  Data are as in a except that the metric analyzed was the allele-sharing statistic genotype homozygosity, G1.  $^{\rm *P}$  < 0.05.

 $D_{xy}$  (see above). For allele-sharing inference in individual populations of *S. cerevisiae* in Supplementary Table S3, we proceeded as above except that we used unphased VCF genotypes for each of the five largest *S. cerevisiae* populations from 1011 Yeast Genomes (Peter *et al.* 2018).

We also evaluated allele-sharing at thermotolerance genes in S. *paradoxus*, as a complement to the above analyses in S. *cerevisiae*. For this purpose, we used unphased VCF genotypes for all S. *paradoxus* genomes from Bergström *et al.* (2014) according to the methods above and found no significant enrichment for G1 at thermotolerance loci (resampling P = 0.19).

#### Polymorphism in Wine/European S. cerevisiae

For polymorphism calculations in Supplementary Figure S1, from the Wine/European S. *cerevisiae* population from Peter *et al.* (2018), we used genotype data as a VCF as input into VCFtools (Danecek *et al.* 2011) with the command –site-pi. This output a polymorphism ( $\pi$ ) value for each SNP. We tabulated the average  $\pi$  across a 1200 SNP window centered at each SNP, with invariant sites contributing  $\pi = 0$ .

#### ESP1 phylogenetic analysis

We used the alignment of the open reading frame of ESP1 from the type strains of S. cerevisiae, S. paradoxus, S. mikatae, S. bayanus, S. uvarum, and S. kudriavzevii from saccharomycessensustricto.org as input into the codeml module of PAML4.9 (Yang 2007). The branch-site model (model = 2, NSsites = 2) was used, and two models, null and alternative, were fitted. In the null model, the dN/dS for the S. cerevisiae branch was fixed at 1.0 and all other branches were described by the same dN/dS ratio ( $\omega$ ). In the alternative model, the branch leading to S. cerevisiae was fitted with one  $\omega$ , and all other branches were fitted with a separate  $\omega$ . A test statistic, calculated by comparing the likelihood ratios of the alternative and null models, was used to calculate a P-value by comparing it to a chi-squared distribution with one degree of freedom, equal to the difference in the number of parameters in the two models. No codons exhibited a posterior probability of positive selection, on the branch leading to S. cerevisiae, higher than 0.9.

### Analysis of cis-regulatory expression divergence between S. cerevisiae and S. paradoxus

For Supplementary Table S5, we analyzed temperature-dependent allele-specific expression measurements in interspecific hybrids as follows. For each gene in turn, from Tirosh et al. (2009) we tabulated the log<sub>2</sub>-ratio of allele-specific expression between alleles of an S. cerevisiae x S. paradoxus hybrid cultured at 35°C, as a difference from the analogous quantity from cultures at 30°C, which we refer to as  $D_{\mbox{\scriptsize ASE}}.$  To test for an enrichment of high-magnitude allele-specific expression differences between species at thermotolerance loci, we took the absolute value of  $D_{ASE}$  for each gene and then tabulated the median of this value across the thermotolerance gene set. We next sampled 10,000 random cohorts of genes from the genome with the same number of essential and nonessential genes as in the thermotolerance cohort (Winzeler et al. 1999), and for each, we calculated the median absolute value D<sub>ASE</sub> . We then used as an empirical P-value the proportion of random cohorts for which the median was equal to or exceeded that seen in the thermotolerance genes. To test for directional cis-regulatory change between species at thermotolerance genes, we repeated the above analysis except that we took the median across signed D<sub>ASE</sub> values for a gene set of interest.

Separately, we repeated the above analysis using measurements from 37 to  $33^{\circ}$ C cultures of an S. cerevisiae  $\times$  S. uvarum hybrid from Li

and Fay (2017), which are reported as allele-specific expression  $a_{T, s}$  for the allele from species s at temperature T. We tabulated  $D_{ASE}$  for each gene as  $log_2(a_{37, Scer}/a_{37, Suv}) - log_2(a_{33, Scer}/a_{33, Suv})$  and tested for enrichment of high-magnitude and directional cis-regulatory variation across thermotolerance genes as above.

# Interspecies swap strain construction at ESP1 promoter and coding region

To swap the allele of the ESP1 promoter from *S. paradoxus* Z1 into *S. cerevisiae* DBVPG1373, and likewise for the coding region, we designed allele-specific Cas9 guide RNAs for the *S. cerevisiae* background, generated donor DNA from *S. paradoxus*, transformed, and screened for successful transgenesis by Sanger sequencing



**Figure 1** A peak of high-allele frequency in *S. cerevisiae* populations at the 5' end of ESP1. Each panel shows results of analysis of allele frequency at the thermotolerance gene ESP1 in a population of *S. cerevisiae* from Peter et al. (2018). In each panel, the y-axis reports genotype homozygosity, G1, in a 1200-SNP window around the position shown on the x. The ESP1 open reading frame is demarcated with a dark black arrow (direction of transcription is right to left). (A) Wine/European population. (B) Mosaic Region 3 population. (C) Brazilian Bioethanol population. (D) Sake population. (E) Mixed Origin population.

as in Weiss *et al.* (2018). Strains are listed in Supplementary Table S6.

#### Large-format growth assay

For growth measurements in Figure 2 and Supplementary Figure S2, we assayed S. paradoxus Z1, S. cerevisiae DBVPG1373, the full ESP1 swap in the S. cerevisiae background (harboring the promoter and open reading frame from S. paradoxus) from Weiss et al. (2018), and the ESP1 promoter and coding swaps in the S. cerevisiae background (see above) as follows. Each strain was streaked from a  $-80^{\circ}$ C freezer stock onto a yeast peptone dextrose (YPD) agar plate and incubated at room temperature for 3 days. For each biological replicate, a single colony was inoculated into 5 mL liquid YPD and grown for 24 hours at 28°C with shaking at 200 rpm to generate pre-cultures. Each pre-culture was backdiluted into YPD at an  $OD_{600}$  of 0.05 and grown for an additional 5.5-6 hours at 28°C, shaking at 200 rpm, until reaching logarithmic phase. Each pre-culture was again back-diluted into 10 mL YPD in 1-inch diameter glass tubes with a target OD<sub>600</sub> of 0.05; the actual  $OD_{600}$  of each was measured, after which it was grown at either 28 or 39°C with shaking at 200 rpm for 24 hours, and OD<sub>600</sub> was measured again. The growth efficiency for each replicate was calculated as the difference between these final and initial OD<sub>600</sub> values. The pipeline from inoculation off of solid plates through preculture, two back-dilutions, and growth at 28 or 39°C we refer to as a day's growth experiment. For each day's

experiments, we calculated the average efficiency  $\langle e_{Scer} \rangle$  across the replicates of wild-type S. *cerevisiae*, and we used this quantity to normalize the efficiency  $e_s$  measured for each replicate assayed on that day of a genotype of interest s. Thus, the final measurement used for analysis for each replicate on a given day was  $e_s/\langle e_{Scer} \rangle$ . We carried out a total of 2–5 days' worth of replicate growth experiments for each genotype, with three separate transformant strains analyzed by this workflow in the case of the coding swap. For a given genotype we used the complete cohort of measurements of  $e_s/\langle e_{Scer} \rangle$  from all days and strains as input into a one-sample, one-tailed Wilcoxon test to evaluate whether  $e_s/\langle e_{Scer} \rangle$  was less than 1 (i.e., that the strain grew worse in the condition of interest than wild-type S. *cerevisiae*).

#### Temperature dose-response growth assay

To evaluate temperature dose-responses in Figure 3, we assayed S. *paradoxus* Z1, S. *cerevisiae* DBVPG1373, and the full ESP1 swap in the S. *cerevisiae* background (harboring the promoter and open reading frame from S. *paradoxus*) from Weiss *et al.* (2018) as follows. Each strain was streaked from a  $-80^{\circ}$ C freezer stock onto a YPD agar plate and incubated at room temperature for 3 days. For each biological replicate, a single colony was inoculated into 5 mL liquid YPD and grown for 48 hours at  $28^{\circ}$ C with shaking at 200 rpm to create a stationary phase pre-culture. From each preculture we made eight back-dilution experimental cultures in a standard PCR strip tube, each in  $200 \,\mu$ l YPD, and we incubated



**Figure 2** The *S. cerevisiae* ESP1 coding region, but not the promoter, is required for thermotolerance. Each column represents results from biomass accumulation assays of a wild-type or transgenic yeast strain cultured at high temperature. The y-axis reports the optical density of a culture of the indicated strain after 24 h at 39°C, normalized to the analogous quantity from wild-type *S. cerevisiae* (dashed line). Each point reports results from one biological replicate, and each bar height reports the average across replicates (n = 6-18). The first two columns report results from wild-type (WT) strains of *S. paradoxus* Z1 (Sp) and *S. cerevisiae* DBVPG17373 (Sc). The last three columns report strains with the indicated region of ESP1 from *S. paradoxus* swapped into *S. cerevisiae* at the endogenous location; ESP1 full swap denotes transgenesis of both the promoter and the coding region. \*Wilcoxon test P < 0.004 in a comparison against wild-type *S. cerevisiae*. Culture data at 28°C are given in Supplementary Figure S2.

these in a thermocycler using a gradient protocol from 37.0 to ( $40.8^{\circ}$ C. After 24 hours, 150 µl from each culture was removed and D<sub>600</sub> was measured. The pipeline from inoculation off of solid plates through pre-culture, back-dilution, and growth we refer to as a day's growth experiment for the dose-response of a strain. If For each day's experiments, at a given temperature we calculated the average efficiency < $e_{scer, 37}$ > across the replicates of wild-type *S. cerevisiae* at 37°C, and used it to normalize the efficiency  $e_{s, T}$  measured for each replicate assayed on that day of a strain of interest s at temperature T. Thus, the final measurement used to ranalysis for each replicate and temperature on a given day was  $e_{s, T}/<e_{scer, 37}$ >. We carried out 2 days' worth of replicate growth experiments, and used the complete cohort of measure-

### Microscopy

Microscopy was performed as described in Weiss *et al.* (2018). Images were scored, blinded, for the size of dyads, omitting all clumps of >2 cells. Two replicates per strain and condition were imaged, and 7–16 images per replicate were scored. Significance was evaluated using a two-factor ANOVA test to evaluate strain by temperature effects. The range and mean number of dyads scored per image and per strain are reported in Supplementary Table S7.

ments of  $e_{s, T}/\langle e_{Scer, 37} \rangle$  from all days and all temperatures as

input into a two-factor type 2 ANOVA test for a temperature-by-

strain effect comparing s with S. cerevisiae.

### Data availability

Strains and plasmids are available upon request. The authors affirm that all data necessary for confirming the conclusions of the article are present within the article, figures, and tables. Custom scripts for sequence preparation and population genetics statistics for absolute sequence divergence, absolute nucleotide divergence, and the McDonald-Kreitman neutrality index are available at https://github.com/clairedubin/thermotolerance. Custom scripts for RH-seq reanalysis, multi-locus genotype, and allelesharing inference, and allele-specific expression analysis are available at https://github.com/melanieabrams-pub/thermotoler ance-loci-across-yeasts. Supplemental Material is available at figshare: https://doi.org/10.25387/g3.14461965.

#### Results

# Signatures of adaptation and constraint at S. *cerevisiae* thermotolerance loci

With the goal of investigating evolutionary mechanisms of thermotolerance divergence between yeasts, we started by addressing the genetics of the trait. Our earlier study used genome-scale screens with the reciprocal hemizygosity test (Steinmetz *et al.* 2002; Stern 2014) to identify eight genes at which *S. cerevisiae* harbored pro-thermotolerance alleles relative to those of *S. paradoxus* (Weiss *et al.* 2018). We re-processed these screen data with an improved statistical workflow, to boost power and genome coverage (see Materials and Methods). The results recapitulated seven loci that we had reported and validated, plus an additional seven that had not risen to significance in our original analysis (Supplementary Table S1). We considered the expanded set of loci as a more complete model of the genetic architecture of the trait, which would be well-suited to population and evolutionary analyses.

Thermotolerance is a defining and putatively adaptive character of *S. cerevisiae*, shared among isolates within the species and distinguishing it from the rest of the *Saccharomyces* clade (Sweeney et al. 2004; Gonçalves et al. 2011; Salvadó et al. 2011). We hypothesized that the loci underlying thermotolerance had evolved under positive selection before the radiation of modern S. cerevisiae populations. To test this, we made use of a broad population survey of S. cerevisiae (Peter et al. 2018), and the deepestsampled S. paradoxus populations available [from vineyards and European collection locales (Bergström et al. 2014) and from North America (Durand et al. 2019)]. With these genomes, we first sought to quantify sequence diversity between the species, at thermotolerance genes. The absolute diversity statistic  $D_{xy}$ reaches high levels in a lineage after selection when compared to a representative of the ancestral state (Nei 1987), and is preferred over relative-divergence metrics as a suggestive statistic of adaptation (Noor and Bennett 2009). Using the entire set of population genomes from S. cerevisiae and S. paradoxus, we found enrichment for high D<sub>xy</sub> among our thermotolerance genes (Table 1 and Supplementary Table S2), as expected from previous smallerscale analysis (Weiss et al. 2018). The latter result was mirrored by analyses of individual S. cerevisiae populations (Supplementary Table S3), ruling out demographic artifacts as the source of signal in our species-wide test. Thus, divergence from S. paradoxus at thermotolerance loci is a trend that pervades >30 S. cerevisiae populations, collected in Europe, Asia, Africa, and the Americas, supporting a model of a selective event in the ancestor of modern S. cerevisiae.

We next reasoned that, if evolution had used predominantly amino acid variants in building the thermotolerance trait, the underlying loci would exhibit striking coding variation between species, relative to within-species polymorphism and relative to synonymous changes, as analyzed in the family of methods derived from the McDonald-Kreitman test (McDonald and Kreitman 1991). For enrichment analyses, we used the neutrality index (Stoletzki and Eyre-Walker 2011), which reaches low values in cases of adaptive amino-acid evolution between species. The results revealed a 1.37-fold reduced neutrality index among thermotolerance genes relative to the genome as a whole (Table 1 and Supplementary Table S2). Such a signal strongly supports a history of adaptation at thermotolerance loci, with a mechanism involving changes to protein structure and/or function.

Under our model of thermotolerance evolution, after an ancestral S. cerevisiae population gained the trait long ago, it was maintained by purifying selection throughout the species. To assess signatures of constraint within S. cerevisiae on thermotolerance loci, we reasoned that haplotype-level analyses would have greater power than site-by-site tests. The SelectionHapStats suite (Garud et al. 2015; Harris et al. 2018) can be used for this purpose to detect very recent soft selective sweeps in young populations or, for a putatively ancient adaptation like ours, to report conservation more generally. In analyses using our complete set of genomes from S. cerevisiae, thermotolerance loci were enriched for high genotype homozygosity (Table 1 and Supplementary Table S2), as seen at any given selected site after a sweep as a product of strong allele-sharing (Garud et al. 2015; Harris et al. 2018). As a control for potential demographic effects in this whole-species analysis, we repeated the test paradigm on individual well-sampled S. cerevisiae populations, and again detected elevated genotype homozygosity at thermotolerance genes (Supplementary Tables S2 and S4). Together, our sequence-based analyses establish hallmarks of directional selection at these loci: sequence divergence between S. cerevisiae and S. paradoxus, particularly at amino-acid coding sites, and tight constraint within S. cerevisiae.

# Molecular evolution and functional impact of coding variation at ESP1

We anticipated that inspecting allele-sharing within species, at high resolution across genomic loci, could further help reveal facets of the history of thermotolerance genes. Using the largest well-sampled population of S. cerevisiae (collected from localities across Europe and in vineyards elsewhere), we found that genotype homozygosity was not uniform across a given thermotolerance gene, and for most loci, peaks of allele-sharing could be resolved (Figure 1A and Supplementary Figure S1). As expected, the latter corresponded to troughs of polymorphism across the population as measured by the number of pairwise differences between strains (Supplementary Figure S1). Notably, even at allele-sharing peaks, in absolute terms genotype homozygosity was modest. Across all thermotolerance loci, the top-scoring regions in the wine/European population were at the 5' end of the chromosome segregation gene ESP1 (Figure 1A), where the statistic reached at most a value of 0.15. This is consistent with our inference of an ancient date for positive selection at ESP1 and other loci, since the tight conservation and long haplotypes expected immediately after selection would be eroded over longer timescales (Smith and Haigh 1974; Berry et al. 1991; Weigand and Leese 2018). Using this very highest region of allele-sharing in ESP1 as a test case, we inspected it in other well-sampled S. cerevisiae populations and again found elevated genotype homozygosity (Figure 1, B-E), indicating that all these populations likely have had the same forces at play at the locus.

We next sought to gain deeper molecular insight into thermotolerance genetics and, for this purpose, chose to focus further on ESP1 as a testbed, given its high allele-sharing within S. cerevisiae (Figure 1 and Supplementary Table S2) and dramatic impact on the thermotolerance trait (Weiss et al. 2018). First, using a phylogenetic approach across the Saccharomyces genus, we established a pattern of accelerated protein evolutionary rate along the S. cerevisiae lineage in ESP1 (P = 0.046), consistent with our population-level tests of protein evolution on the larger set of thermotolerance genes (Table 1). Next, to investigate the importance of coding variation at ESP1 experimentally, we turned to an allele-swap design. We introduced the ESP1 coding region and, separately, the ESP1 promoter, from wild S. paradoxus (strain Z1, isolated from an oak tree in the United Kingdom) into a wild S. cerevisiae background (strain DVBPG1373, from soil in the Netherlands). Growth experiments revealed a dramatic, temperature-dependent effect of variation in the ESP1 coding region, with the S. paradoxus allele compromising growth under heat treatment (Figure 2 and Supplementary Figure S2). This transgenic fully recapitulated the impact of a larger, regional swap of the S. paradoxus open reading frame and promoter together into S. cerevisiae (Figure 2 and Supplementary Figure S2). By contrast, the S. paradoxus allele of the ESP1 promoter conferred no defect in thermotolerance when analyzed on its own (Figure 2 and Supplementary Figure S2). As an independent analysis of potential promoter effects, we examined cis-regulatory variation between yeast species in measurements of ESP1 gene expression. We found no overall dramatic tendency for overall cis-regulatory divergence between S. cerevisiae and other species, at ESP1 in particular or across thermotolerance genes as a set (Supplementary Table S5A). Likewise, the latter yielded no signal in tests for directional cis-regulatory divergence (Bullard et al. 2010; Supplementary Table S5B). , hese data highlight the evolutionary and functional importance of amino acid variation between S. cerevisiae and S. paradoxus at ESP1, and raise the possibility that coding divergence may also underlie the thermotolerance effects of other mapped loci.

#### Temperature dependence and cell biology of species divergence effects at ESP1

In further pursuit of the molecular mechanisms of S. cerevisiae thermotolerance, we turned to the potential for clues from temperature-dependent genetics. S. cerevisiae outperforms its sister species at a range of elevated temperatures (Sweeney et al. 2004; Salvadó et al. 2011). Our thermotolerance loci were identified in a screen for effects of interspecies divergence at 39°C (Weiss et al. 2018), and their relevance to growth under other conditions is unknown. Drawing again on ESP1 as a model with which to address this question, we assayed biomass accumulation of wild-type and transgenic strains under a temperature dose-response. In these growth experiments, we observed a gradual decline in wildtype S. cerevisiae and S. paradoxus growth as temperature increased, with the latter more sensitive to heat (Figure 3). Our allele-swap strain in the S. cerevisiae background harboring S. paradoxus ESP1 exhibited a sharp drop in growth at  $\sim$ 38°C; it grew readily below this temperature, phenocopying the wild-type S. cerevisiae progenitor, and at higher temperatures, it exhibited the negligible growth seen in wild-type S. paradoxus (Figure 3). Such a dose-response, resembling the sigmoidal behavior of a cooperative biochemical process, was a synthetic property of the ESP1 interspecies swap, distinguishing it from either wild-type species. These data imply that, at least in the S. cerevisiae background, the function of S. paradoxus Esp1 breaks down with a steep temperature dependence, whose midpoint is close to the conditions under which this gene was originally identified (39°C).

ESP1 encodes separase, which releases sister chromatids for separation into daughter cells during anaphase, cleaving the cohesin ring that has held them together in metaphase. We reasoned that, if *S. paradoxus* Esp1 failed to function, in actively growing cells harboring this allele we would see hallmarks of arrest late in the cell cycle. Quantitative microscopy bore out this prediction: as in wild-type *S. paradoxus* (Weiss *et al.* 2018), large-budded dyads predominated in cultures of the *S. cerevisiae* transgenic with *S.* 



**Figure 3** Growth function of *S. paradoxus* ESP1 declines sharply with temperature. Each trace reports results from biomass accumulation assays of a wild-type or transgenic yeast strain across temperatures. Strain labels are as in Figure 2. The y-axis reports the optical density of a culture of the indicated strain after 24 h at the temperature on the x, normalized to the optical density of that day's wild-type *S. cerevisiae* at  $37^{\circ}$ C.  $^{\circ}P < 10^{-12}$  for the strain by temperature interaction term of a two-factor ANOVA, in a comparison between the indicated strain and wild-type *S. cerevisiae*.

paradoxus ESP1, when incubated at 39°C (Figure 4). These findings are consistent with a mechanism in which heat treatment compromises separase function of the S. paradoxus allele of Esp1, blocking the progress of the cell cycle and limiting viability and biomass accumulation. Under such a model, evolution in S. cerevisiae would have resolved these defects, introducing genetic changes that foster Esp1 function and boost fitness at high temperature.

### Discussion

In the study of adaptation, a trait that arises in a species, goes to fixation, and is maintained for thousands of generations can be seen as the ultimate evolutionary success story. Here, we have used yeast thermotolerance as a model of this process. We shed light on the forces driving the trait as it has evolved between



**Figure 4** The S. *paradoxus* allele of ESP1 compromises cell division at high temperature. (A) Each panel reports a representative image of a wild-type or transgenic yeast strain after incubation for 24 h at the indicated temperature. Strain labels are as in Figure 2. (B) Each bar reports quantification of replicated imaging data of the indicated strain cultured at the indicated temperature, as in (A). For each bar, the y-axis shows the fraction of dyads in the indicated size category. \*P < 0.015 for the strain by temperature interaction term of a two-factor ANOVA, in a comparison between the indicated strain and wild-type S. *cerevisiae*. Experiment details are given in Supplementary Table S7.

species, and we investigated the molecular genetics and cell biology of divergent alleles at the underlying loci.

Our sequence-based tests of thermotolerance loci—revealing divergence and protein evolutionary rate between species, and conservation within S. *cerevisiae*—strongly suggest that the trait arose under a selective sweep before the radiation of modern S. *cerevisiae* populations. Also consistent with the latter model is the fact that at a given thermotolerance gene, alleles from S. *cerevisiae* isolates from around the world were partially sufficient for the trait, when swapped into a poorly performing S. *paradoxus* background (Weiss *et al.* 2018). Plausibly, the initial rise of thermotolerance early in S. *cerevisiae* history could have been driven by the ecology of hot East Asian niches where the species likely originated (Peter *et al.* 2018).

The scenario of an ancient sweep of thermotolerance, whose effects bear out across modern *S. cerevisiae*, sets up an intriguing contrast with traits that undergo independent, parallel adaptations in distinct lineages of a species (Hoekstra and Nachman 2003; Chan *et al.* 2010; Rosenblum *et al.* 2010; Xie *et al.* 2019). Under one compelling model, thermotolerance alleles acquired by an initially small, specialized *S. cerevisiae* ancestor could have enabled later migrants to colonize other warm niches (Robinson *et al.* 2016). That said, additional lineage-specific adaptations to heighten thermotolerance further could also eventually come to light.

Our data also open a window onto the molecular mechanisms of thermotolerance evolution in the yeast system. The patterns we have seen of nonsynonymous sequence variation in thermotolerance genes, and our molecular-genetic experiments at the ESP1 locus, point to a key role for protein-coding variation. What exactly would such amino acid changes be doing at thermotolerance loci? Given how much our temperature dose-response of ESP1 function looks like a two-state protein unfolding curve, it is tempting to speculate that the S. cerevisiae allele of this gene may act by boosting protein stability. If such a mechanism were to prove the general rule for our loci, it would dovetail with the trend for proteome thermostability seen in heat-tolerant species (Leuenberger et al. 2017). Perhaps most likely, however, given the complexity of the trait, is a picture in which evolution tweaked many protein features (e.g., Sas-Chen et al. 2020), at different times and at different loci, as the S. cerevisiae ancestor gained its unique thermotolerance character.

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### **Conflicts of interest**

None declared.

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