Barcoded reciprocal hemizygosity analysis via sequencing illuminates the complex genetic basis of yeast thermotolerance

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37 ABSTRACT

Decades of successes in statistical genetics have revealed the molecular underpinnings of traits as they vary across individuals of a given species. But standard methods in the field can't be applied to divergences between reproductively isolated taxa. Genome-wide reciprocal hemizygosity mapping (RH-seq), a mutagenesis screen in an inter-species hybrid background, holds promise as a method to accelerate the progress of interspecies genetics research. Here we describe an improvement to RH-seq in which mutants harbor barcodes for cheap and straightforward sequencing after selection in a condition of interest. As a proof of concept for the new tool, we carried out genetic dissection of the difference in thermotolerance between two reproductively isolated budding yeast species. Experimental screening identified dozens of candidate loci at which variation between the species contributed to the thermotolerance trait. Hits were enriched for mitosis genes and other housekeeping factors, and among them were multiple loci with robust sequence signatures of positive selection. Together, these results shed new light on the mechanisms by which evolution solved the problems of cell survival and division at high temperature in the yeast clade, and they illustrate the power of the barcoded RH-seq approach.

INTRODUCTION

Understanding how and why organisms from the wild exhibit different traits is a central goal of modern genetics. Linkage and association mapping have driven decades of success in dissecting trait variation across individuals of a given species (Ott et al. 2015; Tam et al. 2019). But since these methods can't be applied to reproductively isolated taxa, progress in the field of interspecies genetics has lagged behind. However, newer statistical-genetic methods appropriate to comparisons between species have been proposed in the recent literature (Weiss and Brem 2019), which hold promise for elucidating the genetics of ancient traits. For most such methods, limitations accruing from throughput and/or coverage issues remain to be refined.

The budding yeast Saccharomyces cerevisiae grows better at high temperature than any other species in its clade (Sweeney et al. 2004; Gonçalves et al. 2011; Salvadó et al. 2011; Hittinger 2013; Weiss et al. 2018), in keeping with its likely ecological origin in hot, East Asian locales (Peter et al. 2018). This derived and putatively adaptive trait serves as a model for the genetic study of deep evolutionary divergences. Thermosensitivity, the ancestral phenotype in the clade, is borne out in S. paradoxus, a close sister species to S. cerevisiae, making the former a useful point of comparison. Our group previously used this system as a testbed to develop RHseq (Weiss et al. 2018), a genomic version of the reciprocal hemizygosity test (Stern 2014) that is well-suited to the mapping of natural trait variation between sister species. This technique starts with the generation of large numbers of random transposon mutant clones of a viable but sterile interspecies hybrid. In a given clone, loss of function from a transposon insertion in one species' allele of a gene reveals the function of the uncovered allele from the other species. These hemizygotes are competed en masse in a condition of interest; the abundance of each hemizygote in turn in the selected pool is quantified by bulk sequencing, and used in a test for allelic impact on the focal trait. In previous work, we identified eight genes through this approach at which species divergence contributed to thermotolerance (Weiss et al. 2018).

Against a backdrop of successful biological and evolutionary inference from our yeast RH-seq pilot (Weiss et al. 2018; Abrams et al. 2021), we noted that the combination of *S. cerevisiae* alleles of all eight genes mapped to thermotolerance recapitulated only <20% of the difference between the species (AlZaben et al. 2021). Thus, many of the determinants of yeast thermotolerance likely remain undetected. If so, boosting the replication and throughput of

genetic mapping, to enable higher statistical power, could help meet the challenge. In our initial implementation of RH-seq, we had quantified the abundance of hemizygotes in a sample by sequencing across the transposon junction with the genome, using one universal primer that recognized the transposon and another recognizing a ligated adapter at DNA fragment ends (Weiss et al. 2018). This protocol, though rigorous, is labor-intensive and expensive, limiting the potential for throughput and coverage. A higher-throughput alternative starts with the tagging of transposon sequences by random short DNA barcodes (Wetmore et al. 2015). After mutagenesis of a genotype of interest by these barcoded transposons, and then selection of the mutants in bulk in a challenging condition, mutant abundance can be quantified from sequencing of DNA straight from the pool with a simple PCR. We set out to adapt this barcoding strategy to enable highly replicated RH-seq, with application to yeast thermotolerance as a test case to achieve a deeper exploration of the complex genetics of the trait.

MATERIALS AND METHODS

Construction of a randomly barcoded piggyBac transposase pool

For barcoded RH-seq, we constructed a pool of plasmids, each harboring the piggyBac transposase and a randomly barcoded copy of the piggyBac transposon, via Golden Gate cloning of random 20bp barcodes flanked by universal priming sites into a plasmid backbone containing the piggyBac machinery, modified from pJR487 (Weiss et al. 2018) as follows (Figure S1).

Preparation of the backbone vector

To allow the use of BbsI as the Type IIS restriction enzyme for Golden Gate cloning of barcodes into pJR487 (see below), we first removed all three BbsI cut sites from pJR487 by introducing silent mutations that disrupted the restriction enzyme's recognition pattern. The resulting plasmid was called pCW328. We next modified pCW328 to make a Golden-Gate-ready vector, with the final identifier pJC31, by replacing transposon nucleotides with those of a stuffer at a location 70 nucleotides from the end of the right arm of the transposon (Table S1); see Supplementary Note and Figure S2 for a description of this choice. The stuffer contained two BbsI cut sites with custom Type IIS overhang sequences from (Lee et al. 2015), and a NotI cut site in between the two BbsI cut sites. All cloning steps were carried out by GenScript, Inc.

Preparation of barcode oligonucleotides

To make barcodes, we acquired an oligonucleotide pool from IDT that contained random 20 bp sequences (from hand-mixed random nucleotides) flanked by universal priming regions, U1 and U2 (Wetmore et al. 2015, Coradetti et al. 2018). These custom oligos were produced and PAGE purified by IDT. Additionally, we designed forward (FW_Bbsl_JC) and reverse (REV_Bbsl_JC) primers which each contained a Bbsl cut site, Bbsl overhang sequences complementary to the backbone vector, and either universal priming sequence (Table S2) (Coradetti et al. 2018). We set up 50 μ L amplification PCR reactions with 1 μ L of random 20 bp barcodes as template, from a 2.5 μ M stock, and 0.25 μ L of each of the forward and reverse primers from a 100 μ M stock. Amplification used Phusion High Fidelity polymerase (NEB) and the following cycling protocol: 98°C for 30 seconds, (98°C for 10 seconds, 58°C for 30 seconds, 72°C for 60 seconds) × 6, 72°C for five minutes. PCR products were purified (Zymo DNA Clean & Concentrator kit) and then combined. This yielded the final donor barcodes: random 20bp barcodes flanked by universal priming regions, with Bbsl cut sites at the extreme edges.

Cloning barcodes into plasmids

To clone barcodes into pJC31, we proceeded in two barcoding reactions.

The first reaction contained 2:1 molar ratio of vector to barcodes (4 μ g of pJC31 and 128 ng of donor barcodes), 5 μ L of 10X T4 Ligase Buffer (ThermoFisher), 2.5 μ L of T4 Ligase (ThermoFisher), 2.5 μ L FastDigest Bpil (ThermoFisher), and sterile water up to 50 μ L. The cycling program was: 37°C for five minutes, (37°C for two minutes, 16°C for five minutes) x 25, 65°C for 10 minutes. Then a mixture containing 5 μ L 10X FastDigest Buffer (ThermoFisher), 3.13 μ L BSA 2 mg/mL (NEB), 12.5 μ L FastDigest Notl (ThermoFisher), and 12.5 μ L FastDigest Bpil (ThermoFisher) was spiked into the reaction and incubated at 37°C for 16 hours to digest unbarcoded backbone vectors. Ten of these reactions were combined, purified, and eluted in H₂O (Zymo DNA Clean & Concentrator). To spot-check this cloning, 5 μ L of this product was transformed into 25 μ L of *E. coli* 10beta electrocompetent cells (NEB). Sanger sequences across the barcode regions of 20 individually miniprepped *E. coli* colonies showed 95% barcoding efficiency.

The second reaction contained 2:1 molar ratio of vector to donor barcodes (4µg of pJC31 and 128 ng of donor barcodes), 5 µL of 10X T4 Buffer (ThermoFisher), 2.5 µL T4 Ligase (ThermoFisher), 2.5 µL Bpil (ThermoFisher), and sterile water up to 50 µL. The cycling program was: 37°C for five minutes, (37°C for two minutes, 16°C for five minutes) x 25, 65°C for 10 minutes. Then a mixture containing 2.5 µL 10X FastDigest Buffer (ThermoFisher), 2.5 µL G Buffer, (ThermoFisher), 3.13 µL BSA 2 mg/mL (NEB), 12.5 µL FastDigest Notl (ThermoFisher), and 12.5 µL Bpil (ThermoFisher) was spiked in the reaction and incubated at 37°C for 16 hours to digest remaining unbarcoded backbone vectors. Six of these reactions were combined, purified, and eluted in H_2O (Zymo DNA Clean & Concentrator). Then every 5 µL of cleaned eluted product was redigested with 5µL of Notl-HF (NEB), 5 µL 10X CutSmart buffer (NEB), and 35 µL H_2O at 37°C for 16 hours then 80°C for 20 minutes. The reactions were purified again (Zymo DNA Clean & Concentrator) and pooled. Spot checks of this cloning reaction proceeded as above, and Sanger sequences across the barcode regions of 20 individually miniprepped *E. coli* colonies showed 95% barcoding efficiency.

Purified plasmids from the two reactions were combined in a master tube of DNA before transforming into electrocompetent *E. coli* cells (NEB) to generate the final barcoded piggyBac pool (final identifier P58). Each electroporation cuvette (BTX) contained 25 μ L of 10beta electrocompetent cells (NEB) and 5 μ L of cleaned master tube DNA from the previous golden gate barcoding step. We performed 21 electroporation reactions in total using the Bio-Rad GenePulser Xcell machine set to 2.0 kV, 200 Ohms, 25 μ F. After electroporation, each culture was recovered in provided outgrowth media (NEB) by shaking at 37°C at 250 rpm for 1.5 hours. After recovery, all independent 21 electroporation reactions were combined.

The combined recovered transformation E. coli culture was used to inoculate two 1L fresh LB cultures containing carbenicillin at 100 μ g/mL to select for E. coli cells containing barcoded piggyBac plasmids. Each culture was incubated for 15.5 hours at 37°C, 250 rpm (overnight) to expand the barcoded piggyBac E. coli pool. Then the two cultures were combined yielding the final barcoded transposon plasmid pool, P58. This was aliquoted into 1 mL volumes with 15% glycerol and stored at -80°C.

Sequencing verification of barcoded piggyBac pool plasmid DNA for barcode diversity To verify barcode diversity in the barcoded piggyBac plasmid pool (P58), we sequenced barcodes as follows. One frozen aliquot of P58 was inoculated into 1.25 L of LB containing

190 carbenicillin 100 µg/mL and grown for 16 hours 37°C, 250 rpm or until it reached an OD₆₀₀ of 2.1. This culture was gigaprepped on using a column kit (Invitrogen) to generate 5 mg of 191 plasmid. We used this as input into a PCR with primers (Table S2) annealing to the universal 192 193 priming regions flanking the barcode. These primers were dual-indexed, although in this work we only carried out sequencing of the resulting amplicon from one end (see below), such that 194 195 only one index was used. The generic form of the forward primer was 196 AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCT(N1-4)xxxxxxGTCGACCTGCAGCGTACG, where the N1-4 represent variable amounts of random 197 198 bases from 1-4 to help samples cluster on the Illumina lane and the (x6) represent a unique 6-199 bp index sequence for multiplexing samples. The generic reverse primer was CAAGCAGAAGACGCATACGAGATxxxxxxxGTGACTGGAGTTCAGACGTGTGCTCTTCCGAT 200 201 CTGATGTCCACGAGGTCTCT . Four PCR reactions used 50 ng of prepped P58 plasmid 202 template each. Amplification used Q5 High Fidelity Polymerase (NEB) and a cycling program 98°C for four minutes, (98°C for 30 seconds, 55°C for 30 seconds, 72°C for 30 seconds) x 25, 203 72°C for five minutes. Each PCR product was purified on a column (Zymo DNA Clean & 204 Concentrator-5 Kit) and eluted in 10 µL prewarmed 65°C provided elution buffer (Zymo). Six µL 205 of each were then combined and sequenced off the U2 region via Illumina amplicon 206 207 sequencing, on one lane of HiSeq4000 SR50 at the Genomics Sequencing Laboratory at UC 208 Berkeley. Reads sequenced per library are reported in Table S3. Sequencing of the E. coli 209 vector pool p58 revealed 27,538,142 barcodes with an estimated sequencing error rate of 210 1.38% analyzed as described (Coradetti et al. 2018).

Yeast hemizygote pool construction via barcoded transposon mutagenesis

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We constructed our yeast hemizygote pool essentially as described (Weiss et al. 2018) but with modifications as follows.

To prepare plasmid DNA for mutagenesis, one frozen aliquot of P58 was inoculated into 1.25L of LB containing carbenicillin $100 \mu g ml^{-1}$ and grown for 16 hours at $37^{\circ}C$, 250 rpm or until it reached an OD₆₀₀/mL of 2.1. This culture was gigaprepped on using a column kit (Invitrogen) to generate 5 mg of plasmid.

Next, we transformed yeast in several, smaller subpools which we combined to form a final pool as follows. We carried out mutagenesis of CW27, an F1 hybrid from the mating of *S. cerevisiae* DBVPG1373 with *S. paradoxus* Z1 (Weiss et al. 2018) across the first two days. The first day, we generated one subpool in a single 50 mL culture and one subpool in five 50 mL cultures at $OD_{600}/mL \sim 0.9$ (~45 OD_{600} units of cells each). The second day, we generated two subpools in five 50 mL cultures each at $OD_{600}/mL \sim 0.9$ (~45 OD_{600} units of cells).

To generate subpools consisting of a single 50 mL culture, one colony of CW27 was inoculated into 5 mL of YPD and incubated at 28°C 200 rpm. 24 hours later, the OD₆₀₀/mL of the overnight culture was 3.86. It was backdiluted to an OD₆₀₀/mL of 0.1 in 50 mL of YPD in a 250 mL Erlenmeyer flask and grown with shaking at 28°C , 200 rpm for 5.5 hours. After 5.5 hours, it had reached OD₆₀₀/mL ~0.9 and cells were at mid-log phase. This 50 mL culture was gently pelleted at 1000xg for three minutes. The pellet was washed with 25 mL sterile water and then 5 mL of 0.1 M lithium acetate (Sigma) mixed with 1X Tris-EDTA buffer (10 mM Tris-HCl and 1.0 mM EDTA); after spin-down, to the tube was added a solution of 0.269 mg of P58 mixed 5:1 by volume with salmon sperm DNA (Invitrogen), followed by 3 mL of 39.52% polyethylene glycol, 0.12 M lithium acetate and 1.2X Tris-EDTA buffer (12 mM Tris-HCl and 1.2 mM EDTA). The tube was rested for 10 minutes at room temperature, then heat-shocked in a water bath at 37°C for 26 minutes. The tube was gently spun at 1000q for three minutes after which supernatant

was removed. We transferred the cells to a flask and added YPD to attain an OD₆₀₀/mL of ~0.35–4 in ~70 mL. Each such culture was recovered by shaking at 28°C and 200 rpm for two hours. G418 (Geneticin; Gibco) was added to each at a concentration of 300 µg/mL to select for those cells that had taken up the plasmid, and cultures were incubated with 200 rpm shaking at 28°C for two days until each reached an OD₆₀₀/mL of ~2.5. We transferred cells from this culture, and YPD + G418 (300 µg/mL), to new 250 mL flasks at the volumes required to attain an OD600/mL of 0.2 in 50 mL each. We cultured each flask with 200 rpm. shaking at 28°C overnight until each reached an OD₆₀₀/mL of 3.43. To cure transformants of the P58 URA3+ plasmid, we spun down 10% of this master culture, and resuspended in water with the volume required to attain a cell density of 1.85 OD₆₀₀/mL. Four mL of this resuspension were plated (1 mL per 24.1 cm x 24.1 cm plate) onto plates containing complete synthetic media with 5fluorooritic acid (0.2% dropout amino acid mix without uracil or yeast nitrogen base (US Biological), 0.005% uracil (Sigma), 2% D-glucose (Sigma), 0.67% yeast nitrogen base without amino acids (Difco), 0.075% 5-fluorooritic acid (Zymo Research)). After incubation at 28°C to enable colony growth, colonies were scraped off all four plates and combined into water at the volume required to attain 44 OD₆₀₀/mL, yielding the transposon mutant hemizygote subpool. This was aliquoted into 1 mL volumes with 10% dimethylsulfoxide and frozen at −80°C.

To generate subpools consisting of five 50 mL cultures, one colony of CW27 was inoculated to 100 mL of YPD in a 250 mL Erlenmeyer flask and incubated shaking at 28°C, 200 rpm. Twenty-four hours later, the OD $_{600}$ /mL of the overnight culture was OD $_{600}$ /mL 3.89. The overnight culture was backdiluted to OD $_{600}$ /mL 0.1 in 250 mL of YPD and incubated for 5.5 hours at 28°C, 200 rpm. After 5.5 hours, the OD $_{600}$ /mL reached 0.9 and cells were split into five 50 mL conical tubes, and subjected each to heat shock as above. We then transferred all cells from this post-transformation culture to one 1L flask and added fresh YPD to attain OD $_{600}$ /mL 0.4 in ~750 mL YPD. The transformed culture was recovered by shaking at 28°C, 200rpm, for two hours. G418 (300mg/ul) was added to select for the transposed cells. The culture continued shaking for 48 hours or until the OD $_{600}$ /mL reached 2.1. This culture was then backdiluted to create a new culture at OD $_{600}$ /mL 0.2 in 500 mL of YPD with 300mg/µL G418 shaking for 24 hours at 28°C, 200 rpm until it reached OD $_{600}$ /mL ~3.4 The curing, scraping, and freezing steps were the same as above.

To combine the four subpools to yield the final 160X hemizygote pool (final identifier P75), three 1 mL aliquots of each subpool were thawed on ice for one hour. They were transferred to each of four 1L flasks with 500 mL YPD to OD_{600}/mL 0.2, cultured at 28°C, 200 rpm for 17 hours upon which the OD_{600}/mL was 3.5-4. They were gently pelleted, combined, and resuspended in two ways to reach OD_{600}/mL of 44: YPD with 15% glycerol and YPD with 7% DMSO, aliquoted to 1 mL volumes, and frozen at -80°C.

Tn-seq mapping of yeast hemizygote pool

Tn-seq library preparation

To associate barcoded transposon insertions to genomic location in the hemizygote pool, which we refer to as Tn-seq, we first sequenced barcoded transposon insertions according to the methods of (Weiss et al. 2018) as follows. Each 44 OD_{600} /mL aliquot of each subpool or final pool was thawed on ice, and its genomic DNA (gDNA) was harvested with the ZR Fungal/Bacterial DNA MiniPrep Kit (Zymo Research). gDNA was resuspended in DNA elution buffer (Zymo Research) prewarmed to 65°C, and its concentration was quantified using a Qubit 4.0 fluorometer. Illumina transposon sequencing (Tn-seq) library construction was as described previously. Briefly, gDNA was sonicated and ligated with common adapters, and for each

fragment deriving from a barcoded transposon insertion in the genome, a sequence containing a barcode, a portion of the transposon, and a portion of its genomic context (the barcoded transposon–genome junction) was amplified using one primer homologous to the U1 region immediately upstream of barcode and another primer homologous to a region in the adapter. See Table S2 for the transposon-specific primer ("forward primer"), where Ns represent random nucleotides, and the indexed adapter-specific primer ("reverse primer"). Amplification used Jumpstart polymerase (Sigma) and the following cycling protocol: 94°C for two minutes, (94°C for 30 seconds, 65°C for 20 seconds, 72°C for 30 seconds) × 25, 72°C for 10 minutes. Sequencing of paired-end reads of 150 bp was done over two lanes on a HiSeq4000 at Novogene Corporation (Sacramento, CA) and one lane on a NovaSeq SP at the Genomics Sequencing Laboratory at UC Berkeley (Berkeley, CA). Reads sequenced per library are reported in Table S4.

Tn-seq data analysis

Tn-seq data of the hemizygote pool was analyzed, to infer transposon insertions on the basis of barcodes detected in reads as junctions with genomic sequence, essentially as described (Coradetti et al. 2018) (https://github.com/stcoradetti/RBseq/tree/master/Old_Versions/1.1.4), with the following modifications. For each barcode, instead of scanning positions for the end of the insertion from a sequence specified by a model file, we searched for the final 22 base pairs of the right arm of the piggyBac transposon allowing for two mismatches. For annotation, we converted the annotation file from https://github.com/weiss19/rh-seq for the *S.* cerevisiae D1373 x *S. paradoxus* Z1 hybrid to a compliant GFF3 file using Another GFF Analysis Toolkit (AGAT) - Version: v0.4.0 (https://github.com/NBISweden/AGAT). Then, we used a custom Jupyter notebook to annotate the file generated by the RBseq mapping software.

Quality control for Tn-seq, to eliminate barcodes whose junction genomic sequence mapped to multiple insertion locations in the hybrid genome, and to minimize the proportion of sequencing errors included in final tallies, was as described (Coradetti et al. 2018). Briefly, we eliminated from further consideration any case where a barcode observed in Tn-seq sequencing data differed from another, much more abundant, barcode by a single base (a total of 2,024,812 offby-one barcodes in 2,888,129 reads). We also filtered out off-by-two barcodes (280,949 barcodes in total). Separately, we eliminated barcodes that were detected in sequencing data as a junction with more than one genomic context, suggesting the respective transposon had inserted into multiple locations in one or many clones (98,669 barcodes where this inference was based on multiple strong mapping matches, and an additional 46,583 barcodes where this inference was ambiguous, with one strong mapping match with reads outnumbered by those assigned to weaker mapping matches). The final filtered barcode set comprised 548.129 uniquely barcoded and mapped inferred transposon insertions in the P75 hemizygote pool, at an average read depth of 308.6 reads, and a median read depth of 47 reads; 166,834 of these insertions were mapped as genic. The annotation script, GFF3 file, and modified mapping script are available at https://github.com/melanieabrams-pub/RH-seq with barcoding.

Competition cultures

For the thermotolerance competition at 37° C (Table S5), one aliquot of the yeast hemizygote pool was thawed and inoculated into 150 mL of YPD in a 250 mL unbaffled Erlenmeyer flask and grown for six hours at 28° C, 200 rpm. This pre-culture (T_0 , at T_0 and T_0 at T_0 at

YPD at the same optical density as the starting culture, for a total of 10-15 generations. Dilutions for the 28°C competition cultures were performed after 8.5, 18.5, and 25.5 hours after the T_0 timepoint, and dilutions for the 37°C competition cultures were performed after 8.5, 18.5, 25.5 hours and 32.5 hours after the T_0 timepoint. The entire cell culture was harvested from each of these biological replicate tubes for sequencing as biological replicates.

For thermotolerance competition at 36° C (Table S6), competition cultures were grown as above with the following differences. The high temperature was 36° C, instead of 37° C. The pre-culture (T₀, at OD₆₀₀/mL of 0.693 after 5.5 hours at 28° C, 200 rpm) was backdiluted to a starting OD₆₀₀/mL of 0.02 for competition cultures at 36° C. Dilutions for both the 28° C and 36° C competition cultures were performed after 8.5, 18.5 and 25.25 hours after the T₀ timepoint. Eleven instead of 12 replicates were carried out at 28° C.

Barcode quantification from competition cultures

Bar-seq library preparation

 To determine the abundance of barcoded transposon mutant hemizygote clones after selection, we sequenced barcodes insertions as follows. Each cell pellet from a selection sample was thawed on ice, and its genomic DNA (gDNA) was harvested with the Zymo QuickDNA Kit (Zymo#D6005). gDNA was resuspending in DNA elution buffer (Zymo Research) prewarmed to 65°C, and its concentration was quantified using a Qubit 4.0 fluorometer. The barcode insertion was amplified as above (see *Sequencing verification of barcoded piggyBac pool plasmid DNA for barcode diversity*). Each PCR product was purified on a column (Zymo DNA Clean & Concentrator) and eluted in 10 µL prewarmed 65°C provided elution buffer (Zymo). Six µL of each were then combined and sequenced off the U2 region by Illumina sequencing on one lane of HiSeq4000 SR50 at the QB3 Genomics Sequencing Laboratory at UC Berkeley.

Bar-seq data analysis

Bar-seg mapping and quantification were as described (Coradetti et al. 2018) (https://github.com/stcoradetti/RBseq/tree/master/Old Versions/1.1.4), wherein only barcodes that passed quality control in Tn-seq (see Tn-seq data analysis above) were analyzed for quantitative measures of abundance via Bar-seq. Thus we did not use in our screen any barcode that was detected in Bar-seq sequence data but not Tn-seq data (the product of e.g. sequencing errors in Bar-seq, or a failure to observe in Tn-seq a barcode associated with a bona fide transposon insertion that could be detected in Bar-seq). A total of 301,349 barcodes conformed to these criteria from across all replicates of Bar-seg in competitions for the dissection of determinants of growth at 37°C relative to 28°C, with an average read depth of 305.3 reads and a median of 12 reads; 89,772 of these Bar-seq detected barcodes corresponded to inferred transposon insertions in genes and were analyzed as input to the reciprocal hemizygosity testing pipeline described below. In a given replicate competition culture we detected a median 1 x 10⁵ barcodes. The latter represented a fifth of the size of the total pool of hemizygotes detectable after quality control by Tn-seq (5.5 x 10⁵; see *Tn-seq data* analysis above). Thus, the extent of bottlenecking in any given competition experiment was modest, with diversity retained at the order of magnitude of the mutant pool size.

Competitions for the dissection of growth at 36°C relative to 28°C (Table S6) used the same procedures as above, mapping a total of 230,469 barcodes, 68,523 of which corresponded to inserts in genes and were analyzed as input to the reciprocal hemizygosity testing pipeline

described below. In a given replicate competition culture, we detected a median 5 x 10⁴ barcodes.

Reciprocal hemizygosity testing

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The tabulated counts of abundance from Bar-seq for each barcode in each replicate were used as input into reciprocal hemizygosity tests essentially as in (Abrams et al. 2021), with slight changes as follows. We had in hand each barcode which had been sequenced as a junction with a unique genomic location in the Tn-seq step and had passed quality control there (see Tnseg data analysis above), and which was now detected in competition cultures. We interpreted each such barcode as reporting a hemizygote clone bearing a transposon insertion at the respective position of the respective species' allele (S. cerevisiae or S. paradoxus), with the other species' allele retained as wild-type at that locus. In what follows, we refer to each such barcode as reporting an inferred hemizygote clone, with respect to its growth behavior in competition cultures. As in (Abrams et al. 2021), for a given biological replicate we normalized the abundances attributed to each inferred hemizygote genotype to the total number of sequencing reads in the respective sample, and we eliminated from further analysis insertions which had been annotated as intergenic, or as corresponding to the plasmid used to generate this library. For reciprocal hemizygosity tests, we excluded from consideration any gene with fewer than three inferred hemizygote genotypes per allele. Of the retained genes, for each inferred hemizygote genotype, we tabulated the quantity aexperimental in the sequencing-based abundance measured after the competition culture in biological replicate i of growth at the experimental temperature (36 or 37°C), and, separately, we calculated a_{control,i}, the analogous quantity from growth at the control temperature (28°C), for i = [1,12]. We then took the mean of the latter and used it to tabulate the temperature effect on the inferred hemizygote genotype in replicate i, $t_i = log_2(a_{\text{experimental},i}/a_{\text{control,mean}})$. As in (Abrams et al. 2021), we eliminated an inferred hemizygote genotype if the coefficient of variation of this quantity exceeded 2.0, or there were fewer than 1.1 normalized reads. With the data for the remaining inferred hemizygote genotypes (Tables S5 and S6), for a given gene, we compiled the vector of the t measurements across replicates and all inferred hemizygote genotypes with each species' allele of the hybrid disrupted in turn, and discarded genes where the coefficient of variation of the t measurements across hemizygote inserts for one or both alleles exceeded 10. For the remainder, we used the Mann-Whitney test to compare these two vectors, with Benjamini-Hochberg correction for multiple testing (Tables S7 and S8). For a given gene, we calculated the effect size as the difference between two values: the log₂(abundance at the experimental temperature/abundance at 28°C) of the average inferred hemizygote genotype representing a transposon insertion in the S. cerevisiae allele, and the analogous quantity among inferred hemizygote genotypes representing insertions in the S. paradoxus allele of the gene. Scripts for this modified RH-seg analysis pipeline are available at https://github.com/melanieabrams-pub/RHseq with barcoding. We earmarked top candidate genes for factors contributing to the thermotolerance of S. cerevisiae as those with corrected Mann-Whitney p < 0.05 in the reciprocal hemizygosity test, and an effect size < -0.5, i.e. disrupting the S. cerevisiae allele was associated with a strong defect in thermotolerance relative to disruption of the S. paradoxus allele; we refer to this gene set as our top barcoded RH-seq hit gene list.

Analysis of inferred interactions between top hit genes from barcoded RH-seq

For the circos plot reporting inferred interactions between top hit genes from barcoded RH-seq, we used the STRING database (Szklarczyk et al. 2021), accessed September 30, 2021, which incorporates experimental/biochemical data from DIP, BioGRID, HPRD, IntAct, MINT, and PDB, and curated data from Biocarta, BioCyc, Gene Ontology, KEGG, and Reactome. Widths of

edges between nodes in the circos plot represent STRING confidence scores, each the probability of a true positive interaction between a given two genes (Szklarczyk et al. 2021).

To test the encoded proteins of top barcoded RH-seq hit genes for enrichment of physical interactions with each other, we used curated known interactions from BioGRID (Oughtred et al. 2021) as housed in the Saccharomyces Gene Database, downloaded February 19, 2021. We tabulated the number of physical interactions between the proteins encoded by RH-seq hit genes, and we divided that by the total number of interactions involving one RH-seq hit gene and any other gene in the genome; call this ratio r_{true} . Then, we drew a random sample of genes from the genome, as described above for GO term resampling. We tabulated, in this random gene set, the number of physical interactions between genes in that sample, and we divided that by the total number of interactions involving one gene in the random sample and any other gene in the genome, to yield $r_{resample}$. We repeated this procedure 10,000 times, and we used the proportion of resampled groups where $r_{resample}$ was greater than or equal to r_{true} as a one-sided p value assessing the significance of enrichment of interactions between our genes of interest.

Gene Ontology analyses of top hit genes from barcoded RH-seq

 To test top barcoded RH-seq hit genes for enrichment for overrepresentation of a particular Gene Ontology (GO) term, we mapped each gene to its Gene Ontology groups based on data from geneontology.com (Ashburner et al. 2000). We filtered out GO terms with fewer than five or with more than 200 gene members. We also filtered out GO terms with identical membership in the genome. We took the subset of the remaining GO terms with at least one member among our top barcoded RH-seq hit genes. Then, we randomly sampled genes from the genome, ensuring the same proportion of essential genes as in our set of top barcoded RH-seq hit genes based on the essentiality annotations of (Winzeler 1999). We tabulated whether our random sample had greater or fewer genes with that GO term than our candidate set. We repeated this procedure 10,000 times and used the proportion of these resampled groups that had more genes in the given GO term as the initial *p* value assessing the significance of the enrichment of that GO term. Then, we applied Benjamini-Hochberg correction for multiple hypothesis testing to generate final, adjusted *p*-values for the enrichment of the given GO term among top barcoded RH-seq hit genes.

To test Biological Process ontologies for enrichment for large magnitudes of the effect of allelic variation on thermotolerance, we used the latter as tabulated in **Reciprocal hemizygosity testing** above. We filtered GO terms as above, and then excluded all genes absent in our barcoded RH seq analysis, which would have no associated quantity for the effect of allelic variation to resample. For each retained term in turn, we first tabulated the median absolute value of the effect size of the gene members for which we had data, e_{true} . Then, we tabulated the analogous quantity for a random sample of the same number of genes from the genome, $e_{resample}$, ensuring the same proportion of essential genes as above. We repeated this procedure 100 times, and used the proportion of the resampled groups for which $e_{resample}$ was greater than or equal to e_{true} as an initial p value assessing the enrichment of large effects of allelic variation in the genes the term. For all GO terms with an initial p value < 0.1, we repeated this procedure 10,000 times to calculate a more precise p value. Then, we applied the Benjamini-Hochberg correction for multiple hypothesis testing to generate final, adjusted p-values for the enrichment of the given GO term for large effects of allelic variation on thermotolerance.

Molecular evolution analysis of RH-seq hit genes

Branch length PAML analysis with codeML was performed as in (Dubin et al. 2020). Hits were manually inspected for the quality of the alignment, and one, *YAL026C*, was discarded for poor alignment quality leading to an artifactually high branch length. We used the inferred branch lengths as input into a resampling test as in **Gene Ontology analyses of top hit genes from barcoded RH-seq** above, and we performed a one-sided significance test for long branch lengths along the *S. cerevisiae* lineage. Branch-site PAML analysis with codeML was performed as in (Abrams et al. 2021). Jalview version 2 was used to visualize the percent identity of amino acid sequence alignments (Waterhouse et al. 2009). McDonald-Kreitman analysis statistics were calculated as in (Abrams et al. 2021). Fisher's exact test was used to compute *p*-values for individual loci, and these were adjusted using the Benjamini-Hochberg correction for multiple hypothesis testing.

Data availability statement

Sequencing data are deposited in the Sequence Read Archive under the accession PRJNA735401. Strains and plasmids are available upon request. Custom scripts for the barcoded RH-seq analysis are available at https://github.com/melanieabrams-pub/RH-seq with barcoding. The authors affirm that all data necessary for confirming the conclusions of the article are present within the article, figures, and tables.

RESULTS

Dissecting thermotolerance divergence between species by barcoded transposon mutagenesis

With the goals of boosting RH-seq throughput and power, and achieving new insights into the genetics and evolution of yeast thermotolerance, we set out to generate an RH-seg reagent for yeast incorporating barcoded transposons (Wetmore et al. 2015). For this purpose, we first generated a pool of plasmids, each encoding a barcoded copy of the piggyBac transposon and its transposase (Figure S1A-C). To use these in RH-seq, we revisited our previously characterized model system: a comparison between DVBPG1373, a thermotolerant Dutch soil strain of S. cerevisiae, and Z1, an S. paradoxus isolate from the UK (Weiss et al. 2018; AlZaben et al. 2021; Abrams et al. 2021). The F1 hybrid formed from the mating of these strains exhibits a thermotolerance phenotype intermediate between those of the two species parents, and thus is well-suited to mapping of allelic effects on the trait (Weiss et al. 2018). We transformed this F1 hybrid with barcoded plasmids, yielding a pool of hemizygote mutants, which we expanded and then banked (Figure S1D). Next, to catalog the genomic locations of transposon insertions, we used the DNA from a culture of the pool in standard conditions as input into a first round of sequencing library construction, whose primers recognized a common site on the transposon and a common DNA adapter ligated to DNA fragment ends ("Tn-seq"; Figure 1A). Sequencing and data analysis, with quality controls to eliminate barcodes that could not be uniquely associated with a single transposon insertion location (see Methods), yielded a catalog of 548,129 barcoded hemizygotes in the pool whose genomic insertion locations were tabulated. At this point we could harness the pool for highly replicated screens, each of which could quantify hemizygote abundance in a condition of interest via relatively cheap and straightforward barcode sequencing ("Bar-seq"; Figure 1B).

Thus, with our barcoded hemizygote pool, we implemented an RH-seq screen to search for genes at which *S. cerevisiae* and *S. paradoxus* alleles drove differences in strain abundance at high temperature. For this, we subjected the pool to growth assays with 12 biological replicate cultures at 37°C, alongside controls at 28°C. We used DNA from each culture as input into

barcode seguencing (Figure 1B). The resulting data revealed a total of 301,349 cases where a barcode, representing a hemizygote clone with a transposon insertion catalogued by Tn-seq (Figure 1A), was detectable in our growth assays. Transposon insertion positions corresponding to these informative barcodes were evenly split between S. cerevisiae and S. paradoxus alleles of genes throughout the F1 hybrid genome (Figure S3). We took the normalized count of a given barcode in a sequencing data set as a report of the fitness of the respective hemizygote, i.e. its relative abundance after growth in the pool in the respective condition. We then used the complete set of such counts as the input into reciprocal hemizygosity tests to compare, for a given gene, the temperature-dependent abundance of strains harboring a disruption in the S. cerevisiae allele, relative to that of strains with the S. paradoxus allele disrupted. A pipeline for these tests, including filters for coverage and reproducibility and multiple testing correction (see Methods), revealed 83 genes at a 5% false discovery rate (Figure 2 and Table S7). This contrasted with the much smaller set of eight genes at which species' alleles drove differences in high-temperature growth, in our original non-barcoded RH-seq approach (Weiss et al. 2018), which had involved only three biological replicates. The 10-fold increase in the number of significant hits in our barcoded RH-seq screen reflects the statistical power afforded by our highly-replicated method to detect even quite small effects.

In our barcoded RH-seq screen hits, as a positive control we first examined the set of genes known to contribute to thermotolerance divergence from our earlier study (*AFG2, APC1, CEP3, DYN1, ESP1, MYO1, SCC2*, and *DYN1*) (Weiss et al. 2018). Several did not meet the experiment-wide statistical thresholds of our barcoded RH-seq pipeline (Figure S4A), suggesting an appreciable false negative rate of the latter overall. However, manual inspection made clear that hemizygosity effects at all gold-standard thermotolerance loci were borne out: in each case, in barcoded RH-seq data, strains with disruptions in the *S. cerevisiae* allele, and a wild-type copy of the *S. paradoxus* allele, had worse thermotolerance than did strains with only the *S. cerevisiae* allele intact (Figure S4A-B), as we had previously reported (Weiss et al. 2018). Furthermore, the list of gene hits from barcoded RH-seq also included *HFA1* (Figure 2B and Tables S7 and S9) which was reported and validated separately as a determinant of thermotolerance differences between yeast species (Li et al. 2019). On the strength of these controls, we considered our deep sampling of thermotolerance loci to serve as a useful proof of concept for the barcoded RH-seq method.

Functional-genomic analysis of thermotolerance genes

We next aimed to pursue deeper analyses of the novel gene hits from barcoded RH-seq in our yeast thermotolerance application. We considered that a focus on the strongest and most evolutionarily relevant sources of mapping signal would likely yield the most informative results. As such, in light of our interest in explaining the exceptional thermotolerance of purebred *S. cerevisiae*, we earmarked the 44 genes from our larger candidate set at which the *S. cerevisiae* allele boosted the trait most dramatically relative to that of *S. paradoxus* (Figure 2 and Table S9). In what follows, we refer to these genes as our top RH-seq hits, and we analyze them as our highest-confidence predictions for factors that nature would have used in evolving the *S. cerevisiae* phenotype.

We sought to use our mapped loci to explore potential functional mechanisms underlying the thermotolerance trait. We hypothesized that *S. cerevisiae* thermotolerance genes could participate in an interacting network, jointly shoring up particular aspects of cell machinery that were critical for growth at high temperature (AlZaben et al. 2021). Consistent with this notion, the STRING database, which collates experimentally detected protein-protein interactions, genetic interactions, and pathway membership (Szklarczyk et al. 2021), inferred multiple

interactions among our top genes from barcoded RH-seq, with salient signal involving cell cycle factors (Figure 3). A more focused analysis revealed an enrichment, among our top barcoded RH-seq hits, for protein-protein interactions with one another as tabulated in BioGRID (Oughtred et al. 2021), to an extent beyond the null expectation (resampling p = 0.014). We also implemented qualitative gene set enrichment tests, which revealed that chromosome segregation and mitosis factors, although relatively few in number among our top barcoded RHseg hit loci, were significantly enriched relative to the genomic null (Table 1). And we developed a complementary, quantitative test to screen Gene Ontology terms for large allelic effect size (the impact on thermotolerance when the S. cerevisiae allele of a given gene was disrupted in the hybrid, as a difference from the analogous quantity for the S. paradoxus allele; see Methods). Top-scoring in this test was a mitosis gene group, encoding components of the septin ring (GO:0000921; resampling p < 0.0001). Together, these results suggest that our top thermotolerance gene hits share commonalities in function, most notably involving cell cycle factors. This dovetails with previous phenotypic and genetic characterization of yeast thermotolerance, including the breakdown of cell division in heat-treated S. paradoxus (Weiss et al. 2018), and supports a model in which S. cerevisiae acquired thermotolerance in part by resolving the latter cell cycle defect.

The genetics of yeast thermotolerance likely also involves mechanisms beside mitosis, given the known role of mitochondrial genes (Baker et al. 2019; Li et al. 2019) and those operating during stationary phase (AlZaben et al. 2021). Indeed, functional-genomic tests revealed enrichment for secretion genes and for regulatory factors in our top RH-seq hits, although no such group constituted a large proportion of the total hit list (Table 1). Annotations in transcription and translation, mitochondrial function, and signaling were also apparent in our top thermotolerance loci (Figure 2B). These trends are consistent with a scenario in which evolution built the trait in *S. cerevisiae* by tweaking an array of housekeeping mechanisms, beside those that involve cell cycle machinery.

Evolutionary analysis of thermotolerance genes

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645 646 We anticipated that sequence analyses of the genes we had mapped to thermotolerance by barcoded RH-seg could shed light on the evolutionary history of the trait. To explore this, we used a phylogenetic approach in Saccharomyces sensu stricto. We first inferred speciesspecific branch lengths in the phylogeny of each gene in turn, and focused on the lineage leading to S. cerevisiae. The distribution of branch lengths along this lineage among top thermotolerance gene hits was not detectably different from that of the genome as a whole, with the exception of two rapidly evolving thermotolerance genes, TAF2 and BUL1, encoding a transcription initiation factor and ubiquitin ligase adapter respectively (Figure S5). Separately, we quantified protein evolutionary rates in top hits from barcoded RH-seq. A branch-site phylogenetic modeling approach (Yang 2007) detected striking evidence for positive selection along the S. cerevisiae lineage in the amino acid permease GNP1, the kinetochore DNA binding factor CBF2, and the sister chromatid cohesion factor CTF18 (Figure 4). Interestingly, however, McDonald-Kreitman tests (McDonald and Kreitman 1991) on population-genomic data did not detect an overall excess of amino acid variation relative to synonymous changes, at these three genes or any other barcoded RH-seq hit locus (Table S10). Thus, even at genes harboring individual codons with likely signatures of selection, we could not detect evidence for a scenario where S. cerevisiae stacked up a large number of unique amino acid changes, in the evolution of thermotolerance. Together, however, our analyses do highlight thermotolerance genes with marked signal for derived alleles in S. cerevisiae at single codons or in the overall DNA sequence—cases where species divergence is likely to be of phenotypic and evolutionary importance.

DISCUSSION

RH-seq power and the interpretation of mapped loci

In this work, we established the barcoded RH-seq method for genetic dissection of trait variation between diverged lineages. RH-seq falls into a family of recently-developed methods that can dissect natural trait variation across species barriers (Weiss and Brem 2019). A chief distinction of RH-seq is its low cost and low overhead, and the barcoding feature we add here cuts down labor and cost even further, enabling high replication.

Our application to yeast thermotolerance serves as an informative model for the performance of barcoded RH-seq on highly genetically complex traits. We pinpointed dozens of candidate genes at which species-level variation contributes to growth at high temperature. And yet we also observed evidence for a sizeable false negative rate among our barcoded RH-seq results, since some validated thermotolerance loci from our earlier screen did not appear among the hits here. Likewise, a separate barcoded RH-seq mapping of yeast species' differences in growth under milder heat stress revealed little signal above the noise (Table S6 and Table S8), likely reflecting very weak genetic effects under this condition. We thus expect that, as would be true for a classical linkage or association scan, the statistical power of a barcoded RH-seq experiment is a function of signal to-noise, genetic complexity, and genetic effect size; and that many thermotolerance loci remain to be identified even in our very deep set of screen results from high-temperature growth.

By virtue of our focus on pro-thermotolerance alleles in *S. cerevisiae*, our work has left open the functional and evolutionary genomics of loci at which the allele from *S. cerevisiae* instead conferred worse thermotolerance than that of *S. paradoxus*, when each in turn was uncovered in the hybrid. Our barcoded RH-seq identified a number of such genes at high statistical significance. These loci may well reflect the accumulation of advantageous alleles in *S. paradoxus*, or deleterious alleles in *S. cerevisiae*, by drift, even as the latter was under selection to improve the trait in evolutionary history. Analogously, in linkage mapping results, the effect of an allele in recombinant progeny from a cross often does not conform to that expected from the respective parent's phenotype (Burke and Arnold 2001; Brem and Kruglyak 2005). It is also possible that some such allelic effects are the product of epistatic interactions between a locus of interest and the hybrid background, and would be phenotypically buffered (and thus evolutionarily irrelevant) in the purebred species. Molecular validation will be necessary to confirm the phenotypic impact of variation at our mapped loci, and its potential dependence on genetic background.

That said, we consider genes with pro-thermotolerance *S. cerevisiae* alleles according to barcoded RH-seq to be strong candidates for *bona fide* determinants of the trait from the wild in this species. Indeed, earlier work has shown that for such genes mapped by RH-seq in the hybrid, the advantage of *S. cerevisiae* alleles is borne out in tests in purebred backgrounds (Weiss et al. 2018). Accordingly, we have shown here that as a cohort, barcoded RH-seq hits with advantageous *S. cerevisiae* alleles exhibit functional and sequence-based attributes consistent with a role in thermotolerance evolution in the wild.

Cellular and molecular mechanisms of thermotolerance

Our top RH-seq hits revealed strong evidence for chromosome segregation and other mitosis functions as a linchpin of *S. cerevisiae* thermotolerance. As a complement to earlier

characterization of six such genes (*APC1*, *ESP1*, *DYN1*, *MYO1*, *CEP3*, and *SCC2*) (Weiss et al. 2018; Abrams et al. 2021), we now report seven new thermotolerance determinants that function in cell division (*MEC1*, *MLH1*, *CTF13*, *CTF18*, *MCM21*, *CBF2*, and *MYO2*). The emerging picture is one in which the ancestor of modern-day *S. cerevisiae*, faced with dysfunction of a slew of mitotic factors at high temperature, acquired variants across the genome to shore up their activity under these conditions. Under one model of *S. cerevisiae* evolution, the particular niche to which this species specialized was one of avid fermentation, producing (and resisting) heat and ethanol at levels that eliminated its microbial competitors (Goddard 2008; Salvadó et al. 2011). In such a scenario, the maximum benefit could well accrue to the organism if it were able to undergo rapid cell division under the challenging conditions of its own making. Consistent with this notion, another budding yeast, *Hanseniaspora*, which often dominates in early fermentation prior to takeover by *S. cerevisiae* (Fleet 2003), underwent evolutionary loss of much of the cell-cycle checkpoint machinery, consistent with a strategy of accelerated growth at any cost to outcompete other species at the respective stage (Steenwyk et al. 2019).

However, since our current hit list includes many genes from other housekeeping pathways, from transcription/translation to transport and lipid metabolism, mitosis does not appear to be the whole mechanistic story for the thermotolerance trait in *S. cerevisiae*. Indeed, other housekeeping factors also showed up in our previous screen (Weiss et al. 2018) and in an elegant complementary study of mitochondrial determinants of thermotolerance divergence between yeast species (Baker et al. 2019; Li et al. 2019). The panoply of functions detected among our mapped loci conforms well to current models of the mechanisms of thermotolerance, which invoke many essential genes and housekeeping processes (Leuenberger et al. 2017).

The latter idea emerged largely from a proteomic study which showed that thermotolerant organisms had higher thermostability of essential proteins of many functions, across the tree of life (Leuenberger et al. 2017). Were sequence changes that led to improved protein stability a linchpin of thermotolerance evolution in S. cerevisiae? Our data are consistent with a mechanistic role for properties of the protein sequences of many thermotolerance genes, in that variation in coding regions has come to the fore in our sequence tests here and those of an earlier small-scale analysis (Abrams et al. 2021). And interestingly, an experimental case study of one of our mapped thermotolerance loci revealed no impact on the trait from variation in the promoter, only from that in the coding region (Abrams et al. 2021). We cannot rule out noncoding determinants in some cases, especially given that a few hundred genes exhibit temperature-dependent cis-regulatory programs unique to S. cerevisiae (Tirosh et al. 2009; Li and Fay 2017). But if coding regions do hold the exclusive key to the mechanism of S. cerevisiae thermotolerance, they could well involve variants that improve protein function and regulation alongside folding/structure at high temperature. Overall, then, we envision that nature could have used a variety of molecular mechanisms in building the trait, given the apparent complexity of the problem. Biochemical studies will be necessary to nail down exactly how S. cerevisiae alleles advance thermotolerance.

In summary, our data reveal a newly detailed picture of the highly polygenic architecture for a natural trait divergence between species. It is tempting to speculate that evolution may draw on a vast number of variants across the genome to refine a trait over millions of generations, making effects stronger, weaker, or less pleiotropic, adding regulatory control, and so on (Orr 1998). If so, these architectures may ultimately conform to the omnigenic model (Boyle et al. 2017)—which was originally applied to human disease genetics, but may also prove to be an apt description of ancient adaptations.

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FIGURE CAPTIONS

Figure 1. Barcoded RH-seq mapping of yeast thermotolerance loci. (A) Barcoded RH-seq sequencing analysis steps. Left, in a pool of *S. cerevisiae* x *S. paradoxus* hybrid hemizygotes, each harboring a transposon (grey rectangle) marked with a unique 20-mer barcode (multicolored) flanked by universal primer sites (U1 and U2), each barcode is associated with its insertion location by transposon sequencing (Tn-seq). Genomic DNA from the pool is extracted, sheared, and ligated to universal adapters (pink ovals), followed by PCR amplification with a transposon-specific primer (forward black arrow) and an adapter-specific primer (reverse black arrow) and sequencing. Right, for barcode sequencing (Bar-seq) to quantify hemizygote strain abundance after pool growth in a condition of interest, genomic DNA is used as input to PCR with primers to universal primer sites for sequencing. (B) Thermotolerance RH-seq screen design. An aliquot of the hemizygote pool was thawed and cultured in large format, then split into small replicate cultures, each maintained in logarithmic growth phase at the temperature of interest by back-dilution, followed by quantification by Bar-seq.

Figure 2. Hits from barcoded RH-seq mapping of yeast thermotolerance. (A) Each panel reports barcoded RH-seq results for a gene at which the *S. cerevisiae* allele is associated with better thermotolerance than the *S. paradoxus* allele, when uncovered in the hybrid background. In a given panel, the *x*-axis reports the log₂ of abundance, measured by RH-seq after selection at 37°C, of a clone harboring a barcoded transposon insertion in the indicated species' allele in a given replicate, as a difference from the analogous quantity for that clone after selection at 28°C on average across replicates. The *y*-axis reports the proportion of observations of all clones bearing insertions in the indicated allele that exhibited the abundance ratio on the *x*, as a kernel density estimate. Shown are the top six genes from among all barcoded RH-seq hit loci in terms of allelic effect size; see Table S7 for effect sizes of the complete set of hits. (B) Subcellular localization of RH-seq hit genes, where available from (Pierleoni et al. 2007) and (Huh et al. 2003). Genes at which effects of allelic variation on thermotolerance were reported previously (Weiss et al. 2018; Li et al. 2019) are denoted in bold type.

Figure 3. Interactions between thermotolerance loci. Each node represents a top hit gene from barcoded RH-seq mapping of thermotolerance. Each chord represents an inferred interaction, taking into account physical and genetic interactions as well as pathway membership, from the STRING database (Szklarczyk et al. 2021). Chords are weighted by the confidence of the inference of interactions; nodes are colored by the number of inferred interactions with other top hits, such that genes with higher numbers of interactions among the hits are represented by warmer colored nodes.

Figure 4. Codons under positive selection in thermotolerance loci. Each panel shows the amino acid sequence context of the codon(s) (red bar) inferred to be under positive selection along the *S. cerevisiae* lineage, in a hit gene from RH-seq thermotolerance mapping. Alignments are colored by percent identity, with darker purples indicating a higher percent identity. (A) *YDR508C/GNP1*. (B) *YGR140W/CBF2*. (C) *YMR078C/CTF18*.

904 TABLES

	n _{observed} :	Adjusted		
GO term	n _{expected}	p	Name	Total n
		(Cellular Component	
GO:0000775	5:1	0.0366	chromosome, centromeric region	75
GO:0000778	4:0	0.0256	kinetochore	40
		1	Molecular Function	
GO:0000149	3:0	0.0701	SNARE binding	28
GO:0008081	2:0	0.0256	phosphoric diester hydrolase activity	11
GO:0004843	2:0	0.0998	thiol-dependent deubiquitinase	24
			Biological Process	
GO:0007165	3:0	0.0923	signal transduction	59
GO:0001403	3:0	0.0923	invasive growth in response to glucose limitation	42
			negative regulation of Ras protein signal	
GO:0046580	2:0	0.0256	transduction	6
GO:0001934	2:0	0.0256	positive regulation of protein phosphorylation	5
GO:0016042	2:0	0.0923	lipid catabolic process	26
GO:0034087	2:0	0.0923	establishment of mitotic sister chromatid cohesion	16

Table 1. Functional enrichment among thermotolerance loci. Each row with numerical data reports a Gene Ontology (GO) term enriched for RH-seq hit genes. nobserved, the number of genes from among top hits from thermotolerance RH-seq that were annotated with the term. nexpected, the number of genes annotated with the term in the same number of randomly chosen genes from the genome, as a median across samples. Adjusted *p*, resampling-based significance of the enrichment after Benjamini-Hochberg correction. Total *n*, the total number of genes annotated in the GO term in *S. cerevisiae*.

SUPPLEMENTARY FIGURE CAPTIONS

Supplementary Figure 1. Making barcoded hemizygotes in a yeast hybrid background for RH-seq. (A) A pool of random N20 barcodes (colors), each flanked by universal priming sites (U1 and U2), was used as input into a PCR with primers containing recognition sites for the Bbsl type IIS restriction enzyme. (B) In a plasmid harboring an un-barcoded piggyBac transposon (gray rectangle) (the kanamycin resistance cassette, kan^R, flanked by left and right transposon arms) and transposase (teal rectangle) (Weiss et al. 2018), a 42 nucleotide stuffer sequence, consisting of two Bbsl restriction enzyme sites flanking a Notl restriction enzyme site and custom overhang sequences (Lee et al. 2015), replaced 42 nucleotides in the right arm of the transposon. (C) Bbsl digestion of the barcodes and stuffer-containing plasmid, followed by stuffer loss and ligation of a barcode into each plasmid, yielded a pool of barcoded plasmids. (D) Transformation of the barcoded transposase plasmid into *S. cerevisiae* x *S. paradoxus* hybrids, followed by transposition and plasmid loss, yielded a pool of marked transposon hemizygote insertion genotypes in the hybrid background.

Supplementary Figure 2. Modifying yeast piggyBac to test barcode insertion positions and transposase optimization, and effects of barcode insertion positions and transposase sequence on yeast piggyBac transposition efficiency. (A) Left, a plasmid from (Weiss et al. 2018) containing the unbarcoded piggyBac transposon (gray) and transposase (teal) was modified to eliminate three BbsI restriction enzyme sites, and used as a backbone for further modifications. Right, test plasmids were mutated at transposase sites designed to optimize codons and increase activity (Yusa et al. 2011). Bottom, test plasmids were modified to incorporate into the transposon a single 20 nucleotide barcode flanked by universal priming regions and custom two-nucleotide overhang sequences (blue squares), either by insertion between the 3'-most end of the left arm and 5' end of the TEF promoter of the kanamycin cassette (bottom left) or replacing endogenous nucleotides inside the right arm of the transposon (bottom right). Pink rectangles indicate transposase binding sites from (Morellet et al. 2018). (B) Each pair of boxes reports transposition test results from a plasmid schematized in (A) in the *S. cerevisiae* x *S. paradoxus* F1 hybrid, with transformation at the indicated temperature. For a given box, the thick black line reports the median; the box extent report quartiles; whiskers report outliers.

Supplementary Figure 3. Gene coverage and read depth in thermotolerance Bar-seq. (A) The *x*-axis reports the number of inferred hemizygote clones in a given gene (corresponding to transposon insertion mutants) whose abundance was detectable in Bar-seq (see Figure 1B), and each bar height reports the number of genes with the number of detectable hemizygotes on the *x*, for the indicated species' allele in the diploid hybrid. The dotted red line indicates the cutoff used in our quality control pipeline for tests of allelic impact on thermotolerance, whereby only genes with greater than three inserts for an allele in the Bar-seq counts were considered. (B) The *x*-axis reports the total number of Bar-seq reads, for a given inferred hemizygote clone in the indicated species' allele, in competition cultures grown at 28°C; each bar height reports the number of inferred hemizygote clones with the Bar-seq abundance on the *x*. (C) Data are as in (B) except that competitions at 37°C were analyzed.

Supplementary Figure 4. Impact on high-temperature growth of allelic variation, in barcoded RH-seq, at genes from a previous thermotolerance screen. (A) Each row reports the allelic effect, the thermotolerance conferred by disruption of the *S. cerevisiae* allele, relative to the analogous quantity for the *S. paradoxus* allele, as measured in barcoded RH-seq, of a gene at which allelic variation was previously reported to impact thermotolerance (Weiss et al. 2018). Genes marked with asterisks were significant at p < 0.05, after quality control for noise and number of inserts and multiple-hypothesis correction. (B) The *x*-axis reports allelic effect for a given gene as in (A); the *y*-axis reports the proportion of genes with the allelic effect on the *x*, with the blue trace showing the distribution across all genes with barcoded RH-seq data, as a kernel density estimate. Red dotted vertical lines represent genes from (A).

Supplementary Figure 5. **Accelerated evolution of thermotolerance loci.** Shown are results of analyses of branch length of top hit genes from barcoded RH-seq mapping of thermotolerance, as inferred from gene trees and normalized for gene length. Each vertical bar reports inferred branch length, along the *S. cerevisiae* lineage, of the indicated RH-seq hit gene. Horizontal lines report median branch lengths across the indicated gene sets. A resampling test for long branches on the *S. cerevisiae* lineage

among top RH-seq hits revealed significant evidence for enrichment (p = 0.0465) but not when *TAF2* and *BUL1* were eliminated (p = 0.1574), attesting to the particularly strong inference of accelerated evolution in the latter two genes.

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SUPPLEMENTARY TABLES

Supplementary Table 1. Plasmids used in this study.

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				insertions; homologous to
				adapter (Figure 1A, left)
	CAAGCAGAAGACGGCATACGAGATC ACTGTGTGACTGGAGTTCAGACGTGT	Index5	ACAGTG	Indexed reverse primer to sequence transposon insertions; homologous to adapter (Figure
P7_MOD_TS_index5	GCTCTTCCGATCT	Index6	GCCAAT	1A, left) Indexed reverse primer to sequence transposon insertions;
P7 MOD TS index6	CAAGCAGAAGACGGCATACGAGATAT TGGCGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCT			homologous to adapter (Figure 1A, left)
P7 MOD TS index7	CAAGCAGAAGACGGCATACGAGATG ATCTGGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCT	Index7	CAGATC	Indexed reverse primer to sequence transposon insertions; homologous to adapter (Figure 1A, left)
P7_MOD_TS_index8	CAAGCAGAAGACGGCATACGAGATT CAAGTGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCT	Index8	ACTTGA	Indexed reverse primer to sequence transposon insertions; homologous to adapter (Figure 1A, left)
	Bar-seq			
P1_BS3_IT001	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNGCACTAGTCGACCTGCAG CGTACG	IT001	ATCACG	Indexed forward Bar- seq primer (Figure 1A, right)
P1 BS3 IT002	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNTGTAGCGTCGACCTGCA GCGTACG	IT002	CGATGT	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT003	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNCGGATTGTCGACCTGCA GCGTACG	IT003	TTAGGC	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT004	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC	IT004	TGACCA	Indexed forward Bar- seq primer

			1	T .=.
	CGATCTNNNNACCAGTGTCGACCTG CAGCGTACG			(Figure 1A, right)
P1 BS3 IT005	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNGTGACAGTCGACCTGCAG CGTACG	IT005	ACAGTG	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT006	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNTAACCGGTCGACCTGCA GCGTACG	IT006	GCCAAT	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT007	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNCTAGACGTCGACCTGCA GCGTACG	IT007	CAGATC	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT008	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNNAGTTCAGTCGACCTGC AGCGTACG	IT008	ACTTGA	Indexed forward Bar- seq primer (Figure 1A, right)
P1 BS3 IT009	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNGACTAGGTCGACCTGCAG CGTACG	IT009	GATCAG	Indexed forward Bar- seq primer (Figure 1A, right)
P1 BS3 IT010	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNTTCGATGTCGACCTGCAG CGTACG	IT010	TAGCTT	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT011	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNCATCGGGTCGACCTGC AGCGTACG	IT011	GGCTAC	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT012	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNNATGTTCGTCGACCTGC AGCGTACG	IT012	CTTGTA	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT013	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNAACTGAGTCGACCTGCAGC GTACG	IT013	AGTCAA	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT014	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNCCTTGAGTCGACCTGCAG CGTACG	IT014	AGTTCC	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT015	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC	IT015	ATGTCA	Indexed forward Bar- seq primer

		1		T
	CGATCTNNNACTGTAGTCGACCTGCA GCGTACG			(Figure 1A, right)
P1_BS3_IT016	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNNCCTGCCGTCGACCTG CAGCGTACG	IT016	CCGTCC	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT017	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNGAGATGGTCGACCTGCAG CGTACG	IT017	GTAGAG	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT018	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNCGCCTGGTCGACCTGCA GCGTACG	IT018	GTCCGC	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT019	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNAAAGTGGTCGACCTGCA GCGTACG	IT019	GTGAAA	Indexed forward Bar- seq primer (Figure 1A, right)
P1 BS3 IT020	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNNCCGGTGGTCGACCTG CAGCGTACG	IT020	GTGGCC	Indexed forward Bar- seq primer (Figure 1A, right)
P1 BS3 IT021	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNGCTTTGGTCGACCTGCAGC GTACG	IT021	GTTTCG	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT022	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNGCATGCGTCGACCTGCA GCGTACG	IT022	CGTACG	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT023	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNGGTGAGGTCGACCTGC AGCGTACG	IT023	GAGTGG	Indexed forward Bar- seq primer (Figure 1A, right)
P1_BS3_IT024	AATGATACGGCGACCACCGAGATCTA CACTCTTTCCCTACACGACGCTCTTC CGATCTNNNNCGATGGGTCGACCTG CAGCGTACG	IT024	GGTAGC	Indexed forward Bar- seq primer (Figure 1A, right)
P2_BS3_IT001	CAAGCAGAAGACGGCATACGAGATC GTGATGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCTGATGTCCACGAGG TCTCT	IT001	ATCACG	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT002	CAAGCAGAAGACGGCATACGAGATA CATCGGTGACTGGAGTTCAGACGTGT	IT002	CGATGT	Indexed reverse Bar- seq primer

			1	1
	GCTCTTCCGATCTGATGTCCACGAGG TCTCT			(Figure 1A, right)
		IT003	TTAGGC	Indexed
	CAAGCAGAAGACGGCATACGAGATG			reverse Bar-
	CCTAAGTGACTGGAGTTCAGACGTGT			seq primer
Do Doo ITaaa	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2_BS3_IT003	TCTCT	ITO04	TO 4 CO 4	right)
	CAAGCAGAAGACGGCATACGAGATT	IT004	TGACCA	Indexed reverse Bar-
	GGTCAGTGACTGGAGTTCAGACGTG			seq primer
	TGCTCTTCCGATCTGATGTCCACGAG			(Figure 1A,
P2 BS3 IT004	GTCTCT			right)
		IT005	ACAGTG	Indexed
	CAAGCAGAAGACGGCATACGAGATC			reverse Bar-
	ACTGTGTGACTGGAGTTCAGACGTGT			seq primer
	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2_BS3_IT005	TCTCT	. 		right)
		IT006	GCCAAT	Indexed
	CAAGCAGAAGACGGCATACGAGATAT TGGCGTGACTGGAGTTCAGACGTGT			reverse Bar- seq primer
	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2 BS3 IT006	TCTCT			right)
		IT007	CAGATC	Indexed
	CAAGCAGAAGACGGCATACGAGATG			reverse Bar-
	ATCTGGTGACTGGAGTTCAGACGTGT			seq primer
	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2_BS3_IT007	TCTCT	ITOOC	A OTTO A	right)
		IT008	ACTTGA	Indexed reverse Bar-
	CAAGCAGAAGACGGCATACGAGATT CAAGTGTGACTGGAGTTCAGACGTGT			seq primer
	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2 BS3 IT008	TCTCT			right)
		IT009	GATCAG	Indexed
	CAAGCAGAAGACGGCATACGAGATC			reverse Bar-
	TGATCGTGACTGGAGTTCAGACGTGT			seq primer
DO DOO ITOOO	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2_BS3_IT009	TCTCT	IT010	TAGCTT	right) Indexed
	CAAGCAGAAGACGGCATACGAGATA	11010	IAGUII	reverse Bar-
	AGCTAGTGACTGGAGTTCAGACGTGT			seq primer
	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2_BS3_IT010	TCTCT			right)
		IT011	GGCTAC	Indexed
	CAAGCAGAAGACGGCATACGAGATG			reverse Bar-
	TAGCCGTGACTGGAGTTCAGACGTGT			seq primer
D2 DC2 IT044	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2_BS3_IT011	TCTCT	IT012	CTTGTA	right) Indexed
	CAAGCAGAAGACGGCATACGAGATTA	11012	CIIGIA	reverse Bar-
	CAAGGTGACTGGAGTTCAGACGTGT			seq primer
	GCTCTTCCGATCTGATGTCCACGAGG			(Figure 1A,
P2_BS3_IT012	TCTCT			right)
		IT013	AGTCAA	Indexed
	CAAGCAGAAGACGGCATACGAGATTT			reverse Bar-
P2_BS3_IT013	GACTGTGACTGGAGTTCAGACGTGT			seq primer

		1		T
	GCTCTTCCGATCTGATGTCCACGAGG TCTCT			(Figure 1A, right)
P2 BS3 IT014	CAAGCAGAAGACGGCATACGAGATG GAACTGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCTGATGTCCACGAGG TCTCT	IT014	AGTTCC	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT015	CAAGCAGAAGACGGCATACGAGATT GACATGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCTGATGTCCACGAGG TCTCT	IT015	ATGTCA	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT016	CAAGCAGAAGACGGCATACGAGATG GACGGGTGACTGGAGTTCAGACGTG TGCTCTTCCGATCTGATGTCCACGAG GTCTCT	IT016	CCGTCC	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT017	CAAGCAGAAGACGGCATACGAGATC TCTACGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCTGATGTCCACGAGG TCTCT	IT017	GTAGAG	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT018	CAAGCAGAAGACGGCATACGAGATG CGGACGTGACTGGAGTTCAGACGTG TGCTCTTCCGATCTGATGTCCACGAG GTCTCT	IT018	GTCCGC	Indexed reverse Bar- seq primer (Figure 1A, right)
P2 BS3 IT019	CAAGCAGAAGACGGCATACGAGATTT TCACGTGACTGGAGTTCAGACGTGTG CTCTTCCGATCTGATGTCCACGAGGT CTCT	IT019	GTGAAA	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT020	CAAGCAGAAGACGGCATACGAGATG GCCACGTGACTGGAGTTCAGACGTG TGCTCTTCCGATCTGATGTCCACGAG GTCTCT	IT020	GTGGCC	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT021	CAAGCAGAAGACGGCATACGAGATC GAAACGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCTGATGTCCACGAGG TCTCT	IT021	GTTTCG	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT022	CAAGCAGAAGACGGCATACGAGATC GTACGGTGACTGGAGTTCAGACGTG TGCTCTTCCGATCTGATGTCCACGAG GTCTCT	IT022	CGTACG	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT023	CAAGCAGAAGACGGCATACGAGATC CACTCGTGACTGGAGTTCAGACGTGT GCTCTTCCGATCTGATGTCCACGAGG TCTCT	IT023	GAGTGG	Indexed reverse Bar- seq primer (Figure 1A, right)
P2_BS3_IT024	CAAGCAGAAGACGGCATACGAGATG CTACCGTGACTGGAGTTCAGACGTGT	IT024	GGTAGC	Indexed reverse Bar- seq primer

	GCTCTTCCGATCTGATGTCCACGAGG		(Figure 1A,
	TCTCT		right)

Supplementary Table 2. Oligonucleotides used in this study.

36°C vs. 28°C barcoded RH-seg in

veast

RBMA039A IT011

7,059,133

RBMA039A_IT012	19,837,690
RBMA039A_IT013	14,393,602
RBMA039A_IT014	19,572,649
RBMA039A_IT015	8,814,383
RBMA039A_IT016	14,587,181
RBMA039A_IT017	8,619,661
RBMA039A_IT018	13,216,843
RBMA039A_IT019	12,499,486
RBMA039A_IT020	11,963,866
RBMA039A_IT021	6,626,989
RBMA039A_IT022	11,649,542
RBMA039A_IT023	5,813,357
RBMA039A_IT024	9,349,172

Supplementary Table 3. Bar-seq sequencing data sets. Each row reports numbers of reads sequenced for the indicated Bar-seq experiment. The first set of rows reports results from a check of barcoded piggyBac transposon plasmids as in Figure S1C; the remaining rows report results from quantification of yeast hemizygote insertion genotypes after competition in the indicated condition, as in Figure 1B of the main text. Experiment identifiers are from BioProject PRJNA735401.

Pool	Library	Reads	Platform	Facility
67	RBJC37	38,713,102	Novaseq SP	UC Berkeley
69	RBJC38	38,875,221	PE150	
69	RBJC39	43,194,450		
69	RBJC40	39,778,862		
69	RBJC41	38,836,065		
67	RBJC42	39,265,466		
67	RBJC43	47,124,575		
67	RBJC44	39,762,187		
67	RBJC48	91,531,071	HiSeq4000 PE150	Novogene
67	RBJC48_reseq	86,892,060		
70	RBCJ51	86,254,426		
70	RBCJ51_reseq	86,130,880		
70	RBJC52	52,108,306		
70	RBJC52_reseq	53,363,169		
71	RBJC54	88,154,532		
71	RBJC54_reseq	86,878,835		
71	RBJC55	90,265,981		
71	RBJC55_reseq	82,130,170		
69	RBJC57	84,296,399		
69	RBJC57_reseq	85,606,080		

Supplementary Table 4. Tn-seq sequencing data sets. Each row reports numbers of reads from the indicated sequencing of insertion positions of barcoded transposons in the *S. cerevisiae* x *S. paradoxus* hybrid, as in Figure 1A, left, of the main text. Experiment identifiers are from PRJNA735401; "reseq" indicates the reads from a technical replicate performed to gather additional reads for the indicated library.

Supplementary Table 5. Abundance of inferred hemizygote insertion genotypes from thermotolerance RH-seq. Each row reports the results of sequencing from one inferred transposon insertion in the *S. cerevisiae* x *S. paradoxus* diploid hybrid after selection of the barcoded transposon pool after competitions comparing growth at 37°C and 28°C, reflecting the abundance in the pool of the respective hemizygote clone harboring the insertion. Chromosome, strand, location, and gene report the fine-scale position of the inferred insertion. Allele, the species parent's homolog in which the transposon insertion lay. Abundance, read counts of the transposon insertion sequenced after selection of the barcoded transposon pool at the indicated temperature, normalized for library size and averaged across the biological replicate cultures. Transposon insertions not detected in any replicate of the indicated selection were assigned an abundance of 1 prior to normalization by library size. CV, coefficient of variation over biological replicates of normalized read counts after selection at the indicated temperature. Barcode, the unique barcode identifier of the transposon insertion.

Supplementary Table 6. Abundance of hemizygote insertion genotypes from RH-seq at 36°C. Data are as in Table S5, except that RH-seq was done using 36°C as the high-temperature condition.

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Supplementary Table 7. Effects of allelic variation in thermotolerance RH-seq. Each row reports the results of reciprocal hemizygote tests on thermotolerance at the indicated gene in the *S. cerevisiae* x *S. paradoxus* diploid hybrid at 37°C. Columns B-G report analyses of abundance upon the aggregation at the gene level of inferred hemizygote genotypes (Table S5) from all biological replicate experiments, filtered for quality control (see Methods). Columns B-D report results of a two-tailed Mann-Whitney statistical test for a difference in the abundance after growth at 37°C, relative to the abundance after growth at 28°C, of hemizygotes harboring transposon insertions in the two species parents' homologs. The Benjamini-Hochberg method was used to correct for multiple testing. Column E reports the log₂(abundance at 37°C/abundance at 28°C) of the average insert in the *S. cerevisiae* allele. Column F reports the analogous quantity among inserts in the *S. paradoxus* allele of the gene. Column G reports the allele-specific effect size, calculated as the difference between the measures of Columns E and F.

Supplementary Table 8. **Effects of allelic variation in RH-seq at 36°C.** Data are as in Table S7, except that the high temperature growth condition was 36°C.

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Hit	Description
VOD400\W\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	Cargo-transport protein involved in endocytosis; interacts with phosphatidylinositol-4-kinase
YGR198W/YPP1	Stt4; GFP-fusion protein localizes to the cytoplasm; YGR198W is an essential gene
VMD0070/LIEA4	Mitochondrial acetyl-coenzyme A carboxylase, catalyzes the production of malonyl-CoA in
YMR207C/HFA1	mitochondrial fatty acid biosynthesis
	Putative protein of unknown function; predicted prenylation/proteolysis target of Afc1p and
VCI 000\A/\AI\/4	Rce1p; green fluorescent protein (GFP)-fusion protein localizes to the cytoplasm and
YGL082W/MIY1	nucleus; YGL082W is not an essential gene
	Component of the Sec23p-Sfb2p heterodimer of the COPII vesicle coat, required for cargo selection during vesicle formation in ER to Golgi transport; homologous to Sec24p and
YNL049C/SFB2	Sfb3p
TNL049C/SFB2	Plasma membrane G protein coupled receptor (GPCR) that interacts with the
	heterotrimeric G protein alpha subunit, Gpa2p, and with Plc1p; sensor that integrates
YDL035C/GPR1	nutritional signals with the modulation of cell fate via PKA and cAMP synthesis
TDE033C/GFTCT	High-affinity glutamine permease, also transports Leu, Ser, Thr, Cys, Met and Asn;
	expression is fully dependent on Grr1p and modulated by the Ssy1p-Ptr3p-Ssy5p (SPS)
YDR508C/GNP1	sensor of extracellular amino acids
121(0000/0141-1	Genome integrity checkpoint protein and PI kinase superfamily member; signal transducer
	required for cell cycle arrest and transcriptional responses prompted by damaged or
YBR136W/MEC1	unreplicated DNA; monitors and participates in meiotic recombination
	Zinc-finger transcription factor of the Zn(2)-Cys(6) binuclear cluster domain type, involved
YML099C/ARG81	in the regulation of arginine-responsive genes; acts with Arg80p and Arg82p
	Adaptor protein required for structural integrity of the SAGA complex, a histone
	acetyltransferase-coactivator complex that is involved in global regulation of gene
YPL254W/HFI1	expression through acetylation and transcription functions
YIL152W/VPR1	Putative protein of unknown function
TIETOZVV VI TCI	Hexameric DNA polymerase alpha-associated DNA helicase A involved in lagging strand
	DNA synthesis; contains single-stranded DNA stimulated ATPase and dATPase activities;
YKL017C/HCS1	replication protein A stimulates helicase and ATPase activities
1112017 0/11001	Essential kinetochore protein, component of the CBF3 multisubunit complex that binds to
	the CDEIII region of the centromere; Cbf2p also binds to the CDEII region possibly forming
YGR140W/CBF2	a different multimeric complex, ubiquitinated in vivo
	Zinc-finger protein involved in transcriptional control of both nuclear and mitochondrial
	genes, many of which specify products required for glycerol-based growth, respiration, and
YJR127C/RSF2	other functions
	Mitochondrial protein of the AAA ATPase family; has ATP-dependent chaperone activity;
	required for assembly of Rip1p and Qcr10p into cytochrome bc(1) complex; mutations in
YDR375C/BCS1	human homolog BCS1L are linked to neonatal mitochondrial diseases
	Protein of unknown function that associates with translating ribosomes; interacts with
YOR091W/TMA46	GTPase Rbg1p
	ATPase of the CDC48/PAS1/SEC18 (AAA) family, forms a hexameric complex; is essential
	for pre-60S maturation and release of several preribosome maturation factors; may be
YLR397C/AFG2	involved in degradation of aberrant mRNAs
	Essential protein, required for biogenesis of the small ribosomal subunit; heterozygous
YNL132W/KRE33	mutant shows haploinsufficiency in K1 killer toxin resistance
	Subunit of a complex with Ctf8p that shares some subunits with Replication Factor C and is
\# 4D0700/077	required for sister chromatid cohesion; may have overlapping functions with Rad24p in the
YMR078C/CTF18	DNA damage replication checkpoint
	Protein of unknown function with similarity to human DOCK proteins (guanine nucleotide
\(\(\mathred{D} \) \(\tag{\tag{\tag{\tag{\tag{\tag{\tag{	exchange factors); interacts with Ino4p; green fluorescent protein (GFP)-fusion protein
YLR422W/DCK1	localizes to the cytoplasm, YLR422W is not an essential protein

	Large subunit of the nuclear mRNA cap-binding protein complex, interacts with Npl3p to
VADAGENI/OTO 4	carry nuclear poly(A)+ mRNA to cytoplasm; also involved in nuclear mRNA degradation
YMR125W/STO1	and telomere maintenance; orthologous to mammalian CBP80
	Multistep regulator of cAMP-PKA signaling; inhibits PKA downstream of Gpa2p and Cyr1p,
YOR371C/GPB1	thereby increasing cAMP dependency; promotes ubiquitin-dependent proteolysis of Ira2p;
TOR3/TC/GPBT	regulated by G-alpha protein Gpa2p; homolog of Gpb2p
VMD004W/CTF12	Subunit of the CBF3 complex, which binds to the CDE III element of centromeres, bending
YMR094W/CTF13	the DNA upon binding, and may be involved in sister chromatid cohesion during mitosis Protein required for mismatch repair in mitosis and meiosis as well as crossing over during
	meiosis; forms a complex with Pms1p and Msh2p-Msh3p during mismatch repair; human
YMR167W/MLH1	homolog is associated with hereditary non-polyposis colon cancer
TIVITY TO 7 VV/IVIETTI	Pheromone-response scaffold protein that controls the mating decision; binds Ste11p,
	Ste7p, and Fus3p kinases, forming a MAPK cascade complex that interacts with the
YDR103W/STE5	plasma membrane and Ste4p-Ste18p; allosteric activator of Fus3p
TERTOSVVISTES	Protein involved in minichromosome maintenance; component of the COMA complex
	(Ctf19p, Okp1p, Mcm21p, Ame1p) that bridges kinetochore subunits that are in contact
YDR318W/MCM21	with centromeric DNA and the subunits bound to microtubules
T D NO TOV V / IVI O IVI Z T	Aminophospholipid translocase (flippase) that maintains membrane lipid asymmetry in
	post-Golgi secretory vesicles; contributes to clathrin-coated vesicle formation and
YAL026C/DRS2	endocytosis; mutations in human homolog ATP8B1 result in liver disease
17(202007)21(02	Subunit of cohesin loading factor (Scc2p-Scc4p), a complex required for loading of cohesin
	complexes onto chromosomes; involved in establishing sister chromatid cohesion during
YDR180W/SCC2	DSB repair via histone H2AX; evolutionarily-conserved adherin
	Non-essential protein of unknown function; involved in signal transduction and the
	genotoxic response; induced rapidly in response to treatment with 8-methoxypsoralen and
YOR092W/ECM3	UVA irradiation
	U1 snRNP protein involved in splicing, required for U1 snRNP biogenesis; contains multiple
YDR235W/PRP42	tetriatricopeptide repeats
	Ubiquitin-specific protease that interacts with Bre5p to co-regulate anterograde and
	retrograde transport between the ER and Golgi; inhibitor of gene silencing; cleaves
YER151C/UBP3	ubiquitin fusions but not polyubiquitin; also has mRNA binding activity
	Ubiquitin-binding component of the Rsp5p E3-ubiquitin ligase complex, functional homolog
	of Bul2p, disruption causes temperature-sensitive growth, overexpression causes
YMR275C/BUL1	missorting of amino acid permeases
	Major apurinic/apyrimidinic endonuclease, 3'-repair diesterase involved in repair of DNA
	damage by oxidation and alkylating agents; also functions as a 3'-5' exonuclease to repair
YKL114C/APN1	7,8-dihydro-8-oxodeoxyguanosine
	GTPase-activating protein that negatively regulates RAS by converting it from the GTP- to
	the GDP-bound inactive form, required for reducing cAMP levels under nutrient limiting
YOL081W/IRA2	conditions, has similarity to Ira1p and human neurofibromin
	Adapter protein for pexophagy and the cytoplasm-to-vacuole targeting (Cvt) pathway;
\/DD0.40.0/4.T0.4.4	directs receptor-bound cargo to the phagophore assembly site (PAS) for packaging into
YPR049C/ATG11	vesicles; required for recruiting other proteins to the (PAS)
	Protein of the Sec1p/Munc-18 family, essential for vacuolar protein sorting; required for the
VOL0050 // (D0.45	function of Pep12p and the early endosome/late Golgi SNARE Tlg2p; essential for fusion of
YGL095C/VPS45	Golgi-derived vesicles with the prevacuolar compartment
	Na+/H+ and K+/H+ exchanger, required for intracellular sequestration of Na+ and K+;
VDD450W/NH IV4	located in the vacuole and late endosome compartments; required for osmotolerance to
YDR456W/NHX1	acute hypertonic shock and for vacuolar fusion
	AAA-peroxin that heterodimerizes with AAA-peroxin Pex6p and participates in the recycling
VKI 107C/DEV1	of peroxisomal signal receptor Pex5p from the peroxisomal membrane to the cystosol;
YKL197C/PEX1	induced by oleic acid and upregulated during anaerobiosis
	Essential 88kDa subunit of the exocyst complex, which mediates polarized targeting of secretory vesicles to active sites of exocytosis; dimeric form of Sec6p interacts with Sec9p
YIL068C/SEC6	
TILUUUU/SEUU	in vitro and inhibits t-SNARE assembly

	One of two type V myosin motors (along with MYO4) involved in actin-based transport of cargos; required for the polarized delivery of secretory vesicles, the vacuole, late Golgi
YOR326W/MYO2	elements, peroxisomes, and the mitotic spindle
YNR045W/PET494	Mitochondrial translational activator specific for the COX3 mRNA, acts together with Pet54p and Pet122p; located in the mitochondrial inner membrane
YJR107W/LIH1	Putative protein of unknown function; has sequence or structural similarity to lipases
	Phospholipase C, hydrolyzes phosphatidylinositol 4,5-biphosphate (PIP2) to generate the signaling molecules inositol 1,4,5-triphosphate (IP3) and 1,2-diacylglycerol (DAG); involved
YPL268W/PLC1	in regulating many cellular processes
YJL062W/LAS21	Integral plasma membrane protein involved in the synthesis of the glycosylphosphatidylinositol (GPI) core structure; mutations affect cell wall integrity
YCR042C/TAF2	TFIID subunit (150 kDa), involved in RNA polymerase II transcription initiation

Supplementary Table 9. Annotations of top hit loci from barcoded RH-seq of thermotolerance. Shown are hits from thermotolerance mapping by barcoded RH-seq (Table S7) that met quality control thresholds and at which disruption of the *S. cerevisiae* allele compromised thermotolerance to a greater extent than did disruption of the *S. paradoxus* allele in the interspecific hybrid.

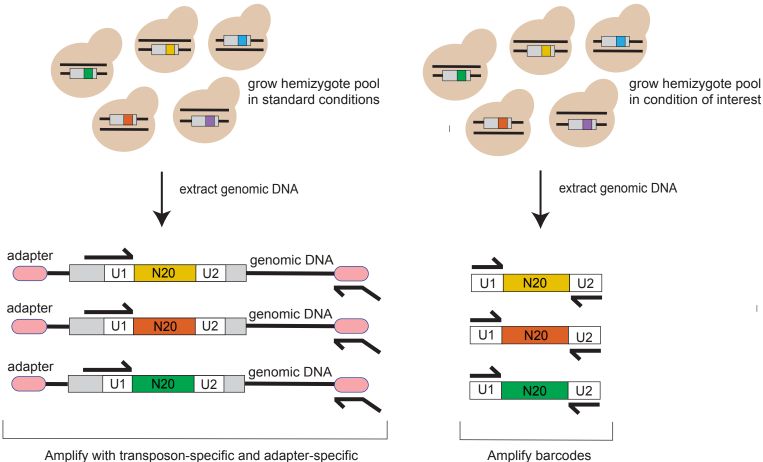
gene	Ds	Dn	Ps	Pn	NI	р	adjusted p
YBR136W/MEC1	293	38	770	363	3.63	5.71 x 10 ⁻¹⁵	1.69 x 10 ⁻¹²
YLR422W/DCK1	122	26	335	280	3.92	1.13 x 10 ⁻¹⁰	8.70 x 10 ⁻⁹
YOR326W/MYO2	35	0	1486	1273	inf	5.15 x 10 ⁻¹⁰	2.77 x 10 ⁻³
YAL026C/DRS2	32	1	1399	1595	36.48	5.81 x 10 ⁻¹⁰	2.97 x 10 ⁻⁸
YMR207C/HFA1	77	16	2215	2103	4.57	6.16 x 10 ⁻¹⁰	3.08 x 10 ⁻⁸
YOR371C/GPB1	79	18	347	323	4.09	1.42 x 10 ⁻⁸	4.02 x 10 ⁻⁷
YML099C/ARG81	96	16	351	247	4.22	1.61 x 10 ⁻⁸	4.43 x 10 ⁻⁷
YJL062W/LAS21	74	10	170	131	5.70	2.19 x 10 ⁻⁸	5.77 x 10 ⁻⁷
YPL268W/PLC1	112	16	245	146	4.17	4.57 x 10 ⁻⁸	1.05 x 10 ⁻⁶
YCR042C/TAF2	155	45	661	470	2.45	1.70 x 10 ⁻⁷	2.90 x 10 ⁻⁶
YPL254W/HFI1	76	10	150	106	5.37	1.66 x 10 ⁻⁷	2.90 x 10 ⁻⁶
YPR049C/ATG11	131	45	448	395	2.57	1.67 x 10 ⁻⁷	2.90 x 10 ⁻⁶
YNL049C/SFB2	32	1	773	587	24.30	3.22 x 10 ⁻⁷	4.88 x 10 ⁻⁶
YKL114C/APN1	44	3	120	87	10.63	9.80 x 10 ⁻⁷	1.19 x 10 ⁻⁵
YIL068C/SEC6	80	5	271	111	6.55	1.36 x 10 ⁻⁶	1.52 x 10 ⁻⁵
YGR198W/YPP1	79	25	295	282	3.02	1.97 x 10 ⁻⁶	2.10 x 10 ⁻⁵
YDR375C/BCS1	67	4	107	54	8.45	2.20 x 10 ⁻⁶	2.28 x 10 ⁻⁵
YKL017C/HCS1	78	13	219	143	3.92	3.56 x 10 ⁻⁶	3.28 x 10 ⁻⁵
YDR235W/PRP42	61	7	177	109	5.37	5.34 x 10 ⁻⁶	4.57 x 10 ⁻⁵
YDR180W/SCC2	173	72	523	431	1.98	6.47 x 10 ⁻⁶	5.35 x 10 ⁻⁵
YMR167W/MLH1	94	26	286	209	2.64	2.47 x 10 ⁻⁵	1.54 x 10 ⁻⁴
YKL197C/PEX1	132	58	415	351	1.92	1.55 x 10 ⁻⁴	6.57 x 10 ⁻⁴
YMR078C/CTF18	73	26	295	260	2.47	1.63 x 10 ⁻⁴	6.82 x 10 ⁻⁴
YGL095C/VPS45	67	14	202	122	2.89	3.72 x 10 ⁻⁴	1.30 x 10 ⁻³
YMR094W/CTF13	54	21	152	158	2.67	4.44 x 10 ⁻⁴	1.50 x 10 ⁻³
YNL132W/KRE33	75	4	187	52	5.21	5.13 x 10 ⁻⁴	1.69 x 10 ⁻³
YDR103W/STE5	93	56	344	373	1.80	1.57 x 10 ⁻³	4.16 x 10 ⁻³
YOR092W/ECM3	16	1	648	498	12.30	1.92 x 10 ⁻³	4.86 x 10 ⁻³
YNR045W/PET494	61	20	177	136	2.34	2.15 x 10 ⁻³	5.32 x 10 ⁻³
YJR107W/LIH1	19	1	102	61	11.36	2.40 x 10 ⁻³	5.79 x 10 ⁻³
YLR397C/AFG2	102	27	288	155	2.03	2.56 x 10 ⁻³	6.10 x 10 ⁻³
YGL082W/MIY1	41	8	132	84	3.26	2.60 x 10 ⁻³	6.17 x 10 ⁻³
YOR091W/TMA46	40	6	103	46	2.98	0.0212	0.0351
YDR456W/NHX1	87	13	187	59	2.11	0.0278	0.0443
YDR508C/GNP1	3	0	959	797	inf	0.2562	0.3009
YIL152W/VPR1	19	11	70	59	1.46	0.4184	0.4669

Supplementary Table 10. Whole-gene tests for evidence of non-neutral protein evolution at thermotolerance loci. Each row reports results from the McDonald-Kreitman test on sequences from strains of European populations of S. cerevisiae and S. paradoxus of the indicated top hit from barcoded RH-seq mapping of thermotolerance. D_s , number of sites of synonymous nucleotide divergence between species; number of sites of D_n , nonsynonymous nucleotide divergence between species; P_s , number of

1034	sites of synonymous nucleotide polymorphisms within species; Pn, number of sites of nonsynonymous
1035	nucleotide polymorphisms within species. NI, neutrality index. The sixth column reports the p-value from a
1036	Fisher's exact test on D _s , D _n , P _s , and P _n , and the seventh column reports the adjusted <i>p</i> -value after
1037	applying the Benjamini-Hochberg correction for multiple hypothesis testing. All loci exhibited NI > 1,
1038	corresponding to a dearth of divergent amino acid changes relative to synonymous changes and
1039	polymorphisms.

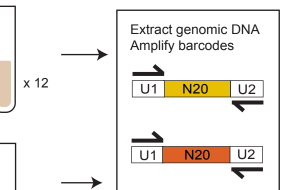
Tn-seq: map barcode to genomic location

Bar-seq: quantify fitness of mutants by sequencing barcodes



Amplify with transposon-specific and adapter-specific primers and sequence

В



and sequence

