Compensatory Genetic and Transcriptional Cytonuclear Coordination in Allopolyploid Lager Yeast (Saccharomyces pastorianus)

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Abstract

Cytonuclear coordination between biparental-nuclear genomes and uniparental-cytoplasmic organelar genomes in plants is often resolved by genetic and transcriptional cytonuclear responses. Whether this mechanism also acts in allopolyploid members of other kingdoms is not clear. Additionally, cytonuclear coordination of interleaved allopolyploid cells/individuals within the same population is underexplored. The yeast Saccharomyces pastorianus provides the opportunity to explore cytonuclear coevolution during different growth stages and from novel dimensions. Using S. pastorianus cells from multiple growth stages in the same environment, we show that nuclear mitochondria-targeted genes have undergone both asymmetric gene conversion and growth stage-specific biased expression favoring genes from the mitochondrial genome donor (Saccharomyces eubayanus). Our results suggest that cytonuclear coordination in allopolyploid lager yeast species entails an orchestrated and compensatory genetic and transcriptional evolutionary regulatory shift. The common as well as unique properties of cytonuclear coordination underlying allopolyploidy between unicellular yeasts and higher plants offers novel insights into mechanisms of cytonuclear evolution associated with allopolyploid speciation.

Key words: cytonuclear coevolution, allotetraploid lager yeast, nuclear mitochondria-targeted gene.

Introduction

Polyploidy, or whole-genome duplication (WGD), is ubiquitous across eukaryotic kingdoms (Jiao et al. 2011; Choleva and Janko 2013; Soltis et al. 2015; Rodriguez and Arkhipova 2018), and has been important in fungal evolution (Albertin and Marullo 2012; Todd et al. 2017; Gorkovskiy and Verstrepen 2021; Marsit et al. 2021). Polyploidy also is known to have contributed to rapid adaptation and increase species diversity during the evolutionary history of the representative model budding yeast genus Saccharomyces (Selmecki et al. 2015). It is possible to analyze the genetic and transcriptional responses to WGD using autopolyploid (polyploids with multiple chromosome sets derived from a single taxon) or allopolyploid (polyploids with chromosomes derived from two or more diverged taxa) yeasts (Krogerus et al. 2016; Scott et al. 2017).

Lager yeast (Saccharomyces pastorianus) is a well-described, naturally evolved allopolyploid species that has been utilized for many centuries for bottom fermentation at low temperatures to produce lager beers (Nakao et al. 2009). Both allotriploid and allotetraploid lager yeasts have been identified (Monerawela and Bond 2018; Salazar et al. 2019). Previous work has suggested that the closest extant relatives of the allopolyploid genome donors to S. pastorianus involve strain(s) of Saccharomyces cerevisiae and an Saccharomyces eubayanus strain from Tibet (Libkind et al. 2011; Bing et al. 2014), with the mitochondrial donor being closely related to the Tibetan S. eubayanus strain (Okuno et al. 2016). There is a well-annotated mitochondrial genome of S. eubayanus (Baker et al. 2015), which is about 22 kb smaller (64.0 kb) than the 85.8 kb mitochondrial genome of S. cerevisiae. Together with a recent chromosome-level genome assembly of lager yeast (S. pastorianus CBS 1483; Salazar et al. 2019), this
system provides the genomic resources necessary for studying mitochondrial genes and identifying subgenomic nuclear homoeologs (genes duplicated via polyploidy) within allopolyploid lager yeasts.

Similar to allopolyploid plant species, allopolyploid lager yeasts need to overcome cytonuclear challenges accompanying the merger of nuclear genomes from two species with the organellar genome from only one (Gong et al. 2012, 2014; Barnard-Kubow et al. 2017; Sloan et al. 2018; Li et al. 2019, 2020; Timouma et al. 2021; Forsythe et al. 2022; Sharbrough et al. 2022). Taking allopolyploid lager yeasts as an example, the combination of divergent S. cerevisiae and S. eubayanus nuclear genomes but the inheritance of only S. eubayanus progenitor organellar mitochondrial genome could result in a challenge of coordination between mitochondrial-targeted S. cerevisiae homoeologs and the S. eubayanus mitochondria. Identification of fertility-related mitochondrial genes in the nuclear genome of hybrid (S. cerevisiae and S. eubayanus) yeast reflects the importance of mitochondrial–nuclear compatibility in yeast speciation (Lee et al. 2008; Solieri 2010; Leducq et al. 2017). Studies focusing on the rbcS-encoding, chloroplast-targeted small subunits of RuBisCo have characterized cytonuclear processes accompanying allopolyploid evolution in angiosperms (Gong et al. 2012, 2014; Wang et al. 2017). One characterized route reducing potential cytonuclear conflict is intergenic homogenization of rbcS homoeologs through “paternal-to-maternal” gene conversions (Gong et al. 2014; Li et al. 2019, 2020; Grover et al. 2022). At the transcriptional level, a second path involves biased expression of rbcS homoeologs in the direction predicted by the organellar origin in the allopolyploids (Gong et al. 2012, 2014; Wang et al. 2017). These and other coordinated cytonuclear responses are thought to stabilize or optimize RuBisCo holoenzyme activity.

Population growth in allotetraploid lager yeasts can be divided into four characteristic stages or phases, namely, lagging, exponential/log, stationary, and death, and are monitored by optical absorbance densities at 600 nm (short OD500). Similar to other microbial populations, Saccharomyces species exhibit unique physical and metabolic states across these different stages or phases (Johnson 1968; Ginovart et al. 2011, 2018; Olivares-Marin et al. 2018). For example, rapid multiplication of cells and consumption of sugar occur in the exponential/log phase, whereas the subsequent stationary phase involves flocculation (the act of yeasts aggregating into “clumps”) and preparation for nutritional constraints (i.e., running out of sugar supply). These characteristic population level features allow us to explore transcriptomic cytonuclear responses that characterize each specific phase. Importantly, allopolyploid yeasts living in the same habitat are interconnected via multiple metabolic and physiological routes, and thus individuals are not really independent (Maclean and Gudelj 2006; Bleoanca et al. 2013; Youk and Lim 2014; Laman Trip and Youk 2020). These interactions integrate the allopolyploid yeast individuals into an interleaved population, representing an interesting model for exploring cytonuclear coordination at the population level in the same habitat.

In the present study, we describe genome-wide cytonuclear coevolution for nuclear mitochondria-targeted (hereafter abbreviated as NMT) genes in fungal allopolyploid yeasts. We identified homologous NMT groups in allopolyploid S. pastorianus and models of its diploid genome donors S. cerevisiae and S. eubayanus. Based on the finding that the NMT homoeologs in S. pastorianus were under strong purifying selection, we characterized genomic and transcriptional coordination during micro-aerobic growth stages. Our work shows that cytonuclear evolutionary responses to allopolyploidy in lager yeasts bear similarities to those of plants, comprising both unidirectional homoeologous gene conversions and biased expression for homoeologs of S. eubayanus origin. Our findings provide novel insights into the general and unique features of cytonuclear coevolution in unicellular fungal allopolyploid species.

Results

NMT Genes are Under Strong Purifying Selection Pressure Conveyed from Cytoplasmic Mitochondria of Saccharomyces eubayanus Origin

Allopolyploid Saccharomyces pastorianus originated through complex hybridization events (allotriploidization and allotetraploidization), involving progenitors similar to modern S. eubayanus and S. cerevisiae. Based on the aligned ORFs of coding genes in mitochondrial DNAs (hereafter mtDNAs), mtDNAs maintained in S. pastorianus were confirmed to be inherited from the S. eubayanus-like parent (Okuno et al. 2016). To increase the resolution of phylogenetic analysis, we compared whole-genomic mtDNA SNPs (in both coding and non-coding regions) in all yeast strains studied to confirm the mitochondrial origin of our S. pastorianus strains. The phylogenetic placement of all S. pastorianus mtDNAs as sister to S. eubayanus confirms that the mitochondrial genomes of all three allopolyploid S. pastorianus strains were indeed contributed by S. eubayanus (supplementary fig. S1, Supplementary Material online).

Allopolyploid S. pastorianus exhibits low-temperature tolerance, enabled by its mitochondrial DNA inherited from cryotolerant S. eubayanus (Baker et al. 2015). Accordingly, it is hypothesized that specific mitochondrial genes (mtDNA genes) in both diploid S. eubayanus and allopolyploid S. pastorianus could be under selection for cold tolerance. In addition, considering the importance of NMT genes in mitochondrial activities (Bousquet et al. 1991; Malina et al. 2018), it is reasonable to hypothesize that selection also acts on NMT genes in cryotolerant yeasts as well. To test these hypotheses, we characterized mtDNA genes, non-NMT, and NMT homolog groups in both diploid S. eubayanus and allopolyploid S. pastorianus. Based upon sequence alignments of each gene homolog group, we estimated and compared their ratios of
nonsynonymous to synonymous substitutions ($d_N/d_S$ ratios) between species at the same and different ploidy levels and even between subgenomes within the same allopolyploid species.

In both diploid $S. eubayanus$ and allopolyploid $S. pastorianus$, all genes except $atp8$ harbored $d_N/d_S$ ratios less than 1.0, consistent with purifying selection (fig. 1A; supplementary table S2, Supplementary Material online). $atp8$ was an outlier with a high $d_N/d_S$ ratio resulting from its single nonsynonymous substitution. Importantly, the $cox1$ gene with a lower $d_N/d_S$ ratio had a specific allele type which was confirmed to confer cryotolerance in $Saccharomyces uvarum$ (Li et al. 2019). For the profiles of NMT and non-NMT genes in the genomes of diploid $S. cerevisiae$, $S. eubayanus$, and allopolyploid $S. pastorianus$ strain(s) were characterized by sequence similarity searches against genes in the fungal secretome and subcellular proteome knowledgebase (Meinken et al. 2014) (table 1), in which mitochondria-targeted is well annotated. A total of 1616 (26.90%) NMT and 4392 (73.10%) non-NMT genes were identified in $S. cerevisiae$, whereas 535 (9.95%) NMT and 4842 (90.05%) non-NMT genes were identified in $S. eubayanus$ (table 1). In principle, the differences in the abundance of NMT genes in $S. cerevisiae$ and $S. eubayanus$ could reflect differences in qualities of their genome assemblies/annotations and/or to actual evolutionary divergence in NMT gene composition in yeast species or strains. Given the numbers and relative abundances of NMT genes in the two subgenomes C ($S. cerevisiae$ contributed; 871 [23.09%], 1261 [23.04%], and 1365 [23.94%] genes) and E ($S. eubayanus$ contributed; 456 [9.30%], 429 [9.15%], and 465 [9.26%] genes) in multiple $S. pastorianus$ allopolyploid strains (table 1), we infer that the difference is real rather than due to annotation or genome assembly issues. Notably, the significantly higher abundance of NMT genes (or lower abundance of non-NMT genes) in $S. cerevisiae$ than those in both $S. eubayanus$ and the C subgenome of $S. pastorianus$ (Proportion test, $P$ value < 0.01; table 1) and absence of this pattern in $S. eubayanus$ and the E subgenome of $S. pastorianus$ (Proportion test, $P$ value > 0.05; table 1) implicate either amplification in the number of NMT genes in $S. cerevisiae$ or genome-wide loss of genes in $S. pastorianus$, consistent with their differences in genome size (85.8 vs 68.6 kb).

**Table 1.** The Number of NMT and Non-NMT Genes Identified in Diploid and Allopolyploid Yeast Strains.

| Species         | Type     | NMT | Non-NMT |
|-----------------|----------|-----|---------|
| $S. pastorianus$| CBS 1513 | 871 | 23.09%  | 2901 | 76.91% |
|                 | E 456    | 9.30% | 4445 | 90.70% |
| $S. pastorianus$| WS 34/70 | 1261 | 23.04%  | 4212 | 76.96% |
|                 | E 429    | 9.15% | 4262 | 90.85% |
| $S. pastorianus$| CBS 1483 | 1365 | 23.94%  | 4336 | 76.06% |
|                 | E 465    | 9.26% | 4554 | 90.74% |
| $S. cerevisiae$ | FostersO| 1616 | 26.90%  | 4392 | 73.10% |
|                 | Ale 1616 | 26.90% | 4392 | 73.10% |
| $S. eubayanus$  | NCYC1063| 1616 | 26.90%  | 4392 | 73.10% |
|                 | Stout 1616 | 26.90% | 4392 | 73.10% |
|                 | CDMF21L1 | 535  | 9.95%   | 4842 | 90.05% |

The bold values represent the relative proportion of NMT or Non-NMT genes within each diploid or polyploid subgenome.

C and E represent the $S. cerevisiae$- and $S. eubayanus$-subgenomes in $S. pastorianus$, respectively.

**Fig. 1.** Nonsynonymous to synonymous substitution ratios ($d_N/d_S$) of mitochondrial and nuclear genes. (A) $d_N/d_S$ ratios (y-axis) of homologous mitochondrial genes (mtDNA genes; x-axis) in $S. pastorianus$ and $S. eubayanus$ are summarized; (B) Each panel summarizes the $d_N/d_S$ ratios of all single-copy NMT and non-NMT genes characterized in diploid orthologs of $S. eubayanus$ and $S. cerevisiae$ and subgenomic homoeologs of allopolyploid $S. pastorianus$. 
Interestingly, the number of NMT and non-NMT genes varied across different allopolyploid S. pastorianus strains, with fewer C subgenomic NMT and non-NMT genes in the CBS 1513 strain (871 [23.09%] and 2901 [76.91%], respectively). To minimize potential artifactual (genome assembly/annotation) effects, we grouped single-copy biparental diploid orthologs with their respective subgenomic homoeologous copies (all homologs exist in single-copy status) for each allopolyploid S. pastorianus strain into a homolog group by OrthoFinder (Emms and Kelly 2019). These single-copy, homologous NMT and non-NMT gene groups were input into subsequent evolutionary analyses (table 2).

We compared $d_{N}/d_{S}$ ratios for diploid orthologs and their respective C and E subgenomic homoeologous groups with the majority of NMT and non-NMT genes being contributed by the E subgenomic homoeolog group (fig. 1B; supplementary table S3, Supplementary Material online). Of note, for parental diploid orthologs, the $d_{N}/d_{S}$ ratios of most NMT and non-NMT genes were zero (fig. 1B), implicating a dominant pattern of purification selection. After allopolyploidy, within both NMT and non-NMT gene groups, $d_{N}/d_{S}$ ratios of C subgenomic homoeologs were significantly higher than those of E subgenomic copies (fig. 1B). Intriguingly, for E subgenomic homoeologs, most $d_{N}/d_{S}$ ratios of NMT and non-NMT homoeologs were less than 1.0 while ratios of NMT group were lower than those of the non-NMT group (Wilcoxon Rank Sum test, $P$ value <0.01; fig. 2B). These changes were consistent with its mtDNAs being contributed by the S. eubayanus-like progenitor, which implicates cytonuclear coevolution; however, more S. cerevisiae-to-S. eubayanus gene conversions in non-NMT homoeologs could be attributed to some uncharacterized selective constraint rather than cytonuclear coevolution.

Table 2. Single-Copy NMT and Non-NMT Gene Groups in Diploid and Allopolyploid Saccharomyces Strains.

| Group Names | Member          | Number of Groups |
|-------------|-----------------|------------------|
| CBS 1513    | S. pastorianus  | 682              |
|             | S. cerevisiae   |                  |
|             | CBS 1513        | 2152             |
|             | FostersO        |                  |
|             | NCYC 1063       |                  |
| WS 34/70    | S. eubayanus    | 971              |
|             | S. pastorianus  |                  |
|             | S. cerevisiae   |                  |
|             | CDFM21L1        | 3015             |
|             | WS 34/70        |                  |
|             | CBS 1483        | 1087             |
|             | FostersO        |                  |
|             | NCYC 1063       |                  |
|             | CBS 1483        | 3226             |
|             | CDFM21L1        |                  |
|             | S. eubayanus    | 4313             |

Saccharomyces cerevisiae-to-Saccharomyces eubayanus Gene Conversions Among NMT Homoeologs

To evaluate possible gene conversion among homoeologs in allopolyploid yeast, we characterized nonsynonymous SNP changes (Materials and Methods; fig. 2A) for single-copy NMT and non-NMT homoeologous (table 2). These changes included conversions in the S. cerevisiae-to-S. eubayanus and S. eubayanus-to-S. cerevisiae directions (S. cerevisiae homoeologous SNPs were converted into S. eubayanus homoeologous SNPs, and vice versa) (supplementary table S5 and S6, Supplementary Material online). We found that the percentages of S. cerevisiae-to-S. eubayanus SNPs were higher in the NMT groups than in the non-NMT groups (binomial test, $P$ value <0.01; fig. 2B) in all three allopolyploid strains. More intriguingly, a higher degree of gene conversions in the S. cerevisiae-to-S. eubayanus direction occurred in the allotetraploid (CBS 1513) than in the allotetraploids (CBS 1483 and WS 34/70; binomial test, $P$ value <0.01; fig. 2B). A representative case of the NMT gene involving Mtx15 homoeologs (encoding membrane-associated mitoribosome receptor) and its respective S. cerevisiae-to-S. eubayanus and S. eubayanus-to-S. cerevisiae gene conversions are illustrated in supplementary fig. S2a, Supplementary Material online, respectively. Overall, the greater S. cerevisiae-to-S. eubayanus gene conversions than the reciprocal direction in the allopolyploid S. pastorianus were consistent with its mtDNAs being contributed by the S. eubayanus-like progenitor, which implicates cytonuclear coevolution; however, more S. cerevisiae-to-S. eubayanus gene conversions in non-NMT homoeologs could be attributed to some uncharacterized selective constraint rather than cytonuclear coevolution.

To determine the extent to which gene conversions occurred in NMT homoeologs, we estimated the proportion of nonsynonymous SNPs involved in gene conversions for each NMT homoeolog group, which we defined as "conversion level" (Materials and Methods; fig. 2C). Notably, the majority of NMT homoeologs were found to exhibit nonsynonymous S. cerevisiae-to-S. eubayanus gene conversions (S. cerevisiae-to-S. eubayanus conversion level = 1.00), meaning that all polymorphic amino acids of most S. cerevisiae NMT homoeologs were replaced by S. eubayanus amino acids (fig. 2C). A representative NMT homoeolog group, PUT2, exhibiting complete S. cerevisiae-to-S. eubayanus conversion, is illustrated in supplementary fig. S2b, Supplementary Material online. We note that some NMT homoeologs harbored a lower level of gene conversion or did not have any nonsynonymous S. cerevisiae-to-S. eubayanus conversions (S. cerevisiae-to-S. eubayanus conversion level = 0) (fig. 2C).

Temporal Transcriptional Expression Preference for Saccharomyces eubayanus NMT Homoeologs in Saccharomyces pastorianus

Cytonuclear coevolution in plant species often entails some degree of homoeologous expression bias favoring the parent contributing the allopolyploid organellar
Fig. 2. Genetic cytonuclear coordination in three allopolyploid S. pastorianus strains. (A) Schematic diagram illustrating the S. cerevisiae-to-S. eubayanus (S. c.-to-S. e.) and S. eubayanus-to-S. cerevisiae (S. e.-to-S. c.) gene conversions (in right and left dotted boxes). Single-copy gene orthologs and homoeologs (C and E subgenomes) are aligned into quartets. Changes of amino acids and nucleotides involved in gene conversion are specified in each box. Labels 1-8 describe the criteria of defining nonsynonymous gene conversions among homoeologs (see Materials and Methods); (B) Number and relative percentage of S. cerevisiae-to-S. eubayanus (S. c.-to-S. e.) and S. eubayanus-to-S. cerevisiae (S. e.-to-S. c.) gene conversions in NMT and non-NMT homoeologs in three S. pastorianus strains are summarized in each column. (C) Density distribution of gene conversion levels of identified S. cerevisiae-to-S. eubayanus (S. c.-to-S. e.) gene conversions in each S. pastorianus strain.

DNAAs (Adams and Wendel 2005; Gong et al. 2012, 2014; Li et al. 2020). As the unicellular allopolyploid yeast enters exponential growth in culture, there are increasing demands for energy and hence mitochondrial activity (Johnson 1968). We hypothesized that NMT homoeologs in S. pastorianus exhibit biased transcriptional responses consistent with selection for metabolic optimization.

To test this hypothesis (supplementary fig. S1, Supplementary Material online), we designed a microaerobic growth experiment, in which the S. pastorianus WS 34/70 strain was incubated for 84 h in YPD (yeast extract–peptone–dextrose) medium. Samples of mixed yeast cells were collected at intervals (three replicates at each sampling point from T1 to T37; 111 samples in total) for RNA-sequencing, covering the exponential and stationary phases (Materials and Methods; fig. 3A). The overall expression levels of all allotetraploid gene homoeologs were mostly similar within each growth phase, whereas among-phase expression differences were significantly higher than within-phase differences (ANOVA test, P value < 0.01; supplementary fig. S3, Supplementary Material online). Subgenomic expression of each NMT and non-NMT homoeolog pair (S. cerevisiae and S. eubayanus homoeologs) was quantified and compared for the replicates at each sampling point. The proportion of gene pairs displaying biased expression towards the S. eubayanus homoeolog was evaluated for NMT and non-NMT homoeolog pairs (fig. 3B). Of note, the extent of biased expression of S. eubayanus homoeologs in NMT and non-NMT homoeologs were similar throughout the stationary stage (fig. 3B); however, in the exponential growth phase, the number of NMT homoeolog pairs exhibiting biased expression favoring S. eubayanus homoeologs was significantly increased relative to those of non-NMT homoeolog pairs, which remained unchanged (fig. 3B).

The increased relative preference for the expression of S. eubayanus NMT homoeologs in the exponential phase could be attributed to the increased expression of S. eubayanus homoeologs and/or decreased expression of S. cerevisiae homoeologs. To investigate this further, we employed a method to categorize NMT homoeologs into clusters (Materials and Methods) (Abu-Jamous and Kelly 2018). The largest cluster (Cluster 0) contained 110 NMT homoeolog pairs that displayed concordant biased expression towards S. eubayanus homoeologs (fig. 3C), with most biased expression reflecting upregulation of S. eubayanus NMT homoeologs (ANOVA test for orthogonal contrasts, P value < 0.01; fig. 3D).

Compensatory Genetic and Transcriptional Cytonuclear Coordination in Saccharomyces pastorianus and Gene Ontology Categorization

The question arises as to whether the observed genetic and transcriptional cytonuclear responses of NMT genes in S. pastorianus are independent, collaborative, or compensatory. Briefly, if genetic and transcriptional cytonuclear co-evolution are independent, those responsive genes are supposed to occupy most proportion of all cytonuclear coordinate genes and be non-overlapping if they are collaborative, there should be many common genes making both genetic and transcriptional cytonuclear co-evolutionary responses; if they are compensatory, those cytonuclear responsive genes are expected to be mutually exclusive.

To address the foregoing question, we examined the overall mutual exclusion/inclusion status of NMT genes exhibiting genetic and/or transcriptional cytonuclear responses (fig. 4), via categorizing the NMT homoeolog pairs (fig. 2C) into two groups according to having either high or low levels of S. cerevisiae-to-S. eubayanus gene conversions (“high” and “low”; fig. 4A). We also quantified expression bias toward S. eubayanus homoeologs at both exponential and stationary phases (fig. 4A). Of note, the low NMT
subgroup exhibited enhanced expression bias toward *S. eubayanus* homoeologs at the exponential stage, which was generally maintained until the stationary stage (fig. 4A); by contrast, the high NMT subgroup maintained a relatively stable homoeologous expression bias during all growth stages (fig. 4A).

Relative gene ontology (GO) enrichment analyses revealed that the high NMT genes were enriched in oxidoreductase activity (molecular function [MF] of GO), mitochondrial outer- and inter-membrane (cell component [CC] of GO), intrinsic components of mitochondrial membrane and envelop lumen (CC of GO), and oxidation–reduction process (biology process [BP] of GO; fig. 4B), whereas the low NMT genes were enriched in cytochrome complex assembly (BP of GO), mitochondrial RNA processing (BP of GO), and mitochondrial nucleoid (CC of GO; fig. 4B). These results indicate that the high conversion NMT genes with significant signals of gene conversion were involved in specialized mitochondrial function, concordant with cytonuclear coevolution.

To further support foregoing analysis, we determined the genetic cytonuclear status of the transcription-responsive NMT genes. Notably, most NMT genes with enhanced expression bias toward *S. eubayanus* homoeologs in Cluster 0 harbored fewer genes with complete nonsynonymous *S. cerevisiae*-to-*S. eubayanus* conversions (*S. cerevisiae*-to-*S. eubayanus* conversion level = 1.00; fig. 4C) than did other NMT genes. Altogether, from both viewpoints, genetic and transcriptional cytonuclear joint responses were mutually compensatory to some extent.

### Discussion

Challenges faced during allopolyploid speciation involves nuclear coordination between parental subgenomes and that between nuclear and organellar genomes. As for the first challenge, loss of chromosomes and gene homoeologs and combination of subgenomic genes into one homoeolog have been characterized extensively in both plant and yeast allopolyploids (Birchler and Veitia 2012; Monerawela et al. 2015; Monerawela and Bond 2018; Salazar et al. 2019; Timouma et al. 2021; Grover et al. 2022; Birchler and Yang 2022). Another challenge is the necessity of cytonuclear coordination among divergent proteins encoded by biparental-nuclear genomes and uniparental-cytoplasmic organellar genomes. To explore the mechanisms whereby such coordination might be achieved, researchers have employed both specific organelle-targeted nuclear genes (e.g., *rbcS*, *Cox5*, and etc.) and/or global re-sequencing and RNA-seq data to characterize the genetic and/or transcriptional
cytonuclear responses in natural and synthetic allopolyploid plants (Huang et al. 2014; Sehrish et al. 2015; Roux et al. 2016; Barnard-Kubow et al. 2017; Bruun-Lund et al. 2017; Wang et al. 2017; de Carvalho Ferreira et al. 2019; Li et al. 2019, 2020; Forsythe et al. 2021; Xu et al. 2021). Nevertheless, it is still unclear to what extent optimal cytonuclear reconfigurations entail changes at genetic or transcriptional levels. To broaden our understanding of cytonuclear evolutionary processes accompanying allopolyploidy across the tree of life, the present study was conducted using the allopolyploid lager yeast S. pastorianus, which also allowed us to characterize cytonuclear coordination of interleaved allopolyploid yeast cells within the same population. Our findings provide novel insights into the general and unique features of cytonuclear coevolution in unicellular fungal allopolyploid species.

**Gene Conversion in Nuclear Mitochondria-Targeted Genes**

Mitochondria play central metabolic roles in yeast as in all aerobic organisms, but in yeast this has an economical vital function in fermentation (Kitagaki and Takagi 2014; Malina et al. 2018). The lager yeast S. pastorianus is known to have acquired cold tolerance (cryotolerance) from S. eubayanus, and hence retains its mitotype (mitochondrial DNA sequences) (Baker et al. 2015). Accordingly, S. eubayanus-contributed mitochondria may have been the target for natural selection in allopolyploid lager yeast favoring cytonuclear or mitonuclear interactions or function. Given the close intracellular communications between the eukaryotic nucleus and cytoplasmic organelles (Storz 2006; Quirós et al. 2016; Guaragnella et al. 2018; Malina et al. 2018), it is conceivable that selection on mtDNA also entails corresponding nuclear genomic responses. Although empirical studies have documented the occurrence of post-homoploid hybridization (hybrid at the same ploidy level) in the diploid yeast nuclear genome (Lee et al. 2008; Paliwal et al. 2014; Hsu and Chou 2017; Jhuang et al. 2017; Baker et al. 2015; Nguyen et al. 2020; Bendixsen et al. 2021), this aspect remains unexplored in allopolyploid yeast.

**Fig. 4.** Compensatory cytonuclear coordination between genetic and transcriptional responses in S. pastorianus strain WS 34/70. (A) Dynamic transcriptional expression bias for S. eubayanus homoeologs with high and low gene conversion levels (abbreviated as high and low at each sampling point in the growth process. The solid curves illustrate linear fitting of expression bias of S. eubayanus NMT homoeologs in each homoeolog group for different gene conversion levels, respectively; (B) GO term enrichments (biological process [BP], cellular component [CC], and molecular function [MF]) for homoeologs with high and low conversion levels (high and low). GO terms specifically enriched in homoeologous groups of high or low conversion level are specified in the diagram; (C) The density distribution of S. cerevisiae-to-S. eubayanus gene conversion levels for homoeologs categorized in Cluster 0 and for all other NMT genes identified in S. pastorianus strain (background).
We propose that the data provided here constitute evidence for natural selection operating on functional cytonuclear optimization. Specifically, some specific mtDNA genes, including cox1, functioning in cellular respiration and mediating cytotolerance in other yeast relatives (Li et al. 2019), appear to have been under stronger purifying selection than other mtDNA genes (fig. 1A). We draw attention to the signals of selection were detected in NMT genes (fig. 1B) encoding proteins involved in mitochondrial activities. Moreover, the preferential occurrence of \textit{S. cerevisiae}-to-\textit{S. eubayanus} conversions in NMT homoeologs, favoring the direction of the mitochondrial genome donor, may reflect selection for optimization of function for interacting proteins encoded by mitochondrial and nuclear genes (fig. 2B). Specific gene homoeologs involved in the mitochondrial oxidation–reduction process (e.g. \textit{PUT2}), exhibited complete or a high level of \textit{S. cerevisiae}-to-\textit{S. eubayanus} gene conversion, which we propose reflects optimization of cytonuclear function in oxidation/reduction in \textit{S. pastorianus} (figs. 2C, 4B, and supplementary fig. S2B, Supplementary Material online).

Further evidence is required to confirm a functional and hence possibly adaptive role for the genetic changes we report here for allopolyploid lager yeast. Taking potential participation in cytotolerance as an example, one might reverse specific \textit{S. cerevisiae}-to-\textit{S. eubayanus} changes by genetic engineering to study possible attenuation or abrogation of cold tolerance. In addition, it would be intriguing to elucidate the pace at which these types of evolutionary changes arise. For example, one might synthesize fast-cycling synthetic allopolyploid yeasts by mating diploid vegetative cells (Sipiczki 2008; Turgeon et al. 2021) and monitor genetic changes across generations.

Temporal Dynamics of Gene Expression and Possible Connections to Cytonuclear Function

In addition to gene conversions, duplicated genes could undergo transcriptional sub-functionalization via biased usage or expression of specific homoeologous copy (Hu et al. 2020; Nieto Feliner et al. 2020; Birchler and Yang 2022). Lager yeast cells propagated during different growth stages allows investigation of the temporal dynamics of transcriptional sub-functionalization and how it relates to possible cytonuclear selection. Here, we showed that: 1) in the early exponential growth stage, biased expression of \textit{S. eubayanus} homoeologs for NMT genes was highest (fig. 3B) (Mishra and Chan 2014). The absence of these dynamics for non-NMT genes suggests a possible optimization for NMT genes during this vigorous propagation stage; 2) upon reaching the stationary phase, relatively high biased expression toward \textit{S. eubayanus} NMT genes is maintained (fig. 3B), perhaps indicating a continued optimization of cytonuclear expression levels even under conditions of limited nutrition and mitotic energetic demand (Leitao and Kellogg 2017; Laman Trip and Youk 2020). Altogether, our data suggest that temporal dynamic transcriptional cytonuclear coordination could play an important fine-tuning role in the growth and propagation of allopolyploid lager yeasts. In addition, the findings of the present study suggest that allopolyploid yeasts commonly grow in an interactive and communicative manner in the same habitat, indicating that we should consider cytonuclear coordination in allopolyploids from a consolidated spatial and temporal perspective. For example, further analyses of allopolyploid lager yeasts growing in cold temperature and sampled in similar multiple growth stages will allow us to characterize the contribution of transcriptional cytonuclear coevolution in adaptation to cold tolerance.

Genetic and Transcriptional Cytonuclear Coordination Mechanisms in Fungi and Plants

Genetic and transcriptional changes represent two different evolutionary responses that might result from selection for cytonuclear functional optimization following allopolyploidy. Conceivably, complete or high-level gene conversion (\textit{S. cerevisiae}-to-\textit{S. eubayanus}) of most NMT homoeologs (fig. 2C) became fixed early in the hybridization and/or polyploidization process, through directional selection to optimize cytonuclear function (e.g., for cryotolerance in low-temperature fermentation and/or growth habitat). As for NMT homoeologs exhibiting lower or no cytonuclear gene conversions, it is possible that this reflects either insufficient evolutionary time since the formation of the \textit{S. pastorianus} strains, and/or the acquisition of physiological optimization via transcriptional cytonuclear responses during growth via differential transcriptional regulation of homoeologs (Johannesen and Hansen 2002; Sood et al. 2017; Hovhannisyan et al. 2020). Our findings implicate orchestrated compensatory genetic and transcriptional mechanisms to resolve cytonuclear conflict or optimize cytonuclear function.

In principle, transcriptional changes might be hypothesized to arise more quickly than genetic changes, with the latter relying on a presumably slower fixation process of favorable genome-wide mutations, whereas the former may be impacted by relatively fewer mutations in regulatory networks that might affect many genes simultaneously. In the present instance, it would be interesting to study newly synthesized allopolyploid yeasts, to determine the pace and evolutionary dynamics of expression-level changes vs. gene conversion over the generations.

As noted above, both gene conversions (in maternal-to-paternal direction) of organelle-targeted nuclear genes and biased expression for maternal homoeologs appear to be common cytonuclear responses following angiosperm allopolyploid speciation (Gong et al. 2014; Li et al. 2019, 2020). It will be interesting to explore the extent to which there are “rules” or commonalities in cytonuclear responses in allopolyploid yeasts of different ploidy levels. As the present study is the first to examine cytonuclear evolution following allopolyploidy in yeasts, it is an open question whether our findings will apply to other fungal systems, and hence how the cytonuclear evolutionary process differs or is
the same in angiosperms and fungi. We note that one aspect of our results, namely, the largely compensatory relationship between transcriptional vs. genetic alteration, has not been observed at a genome-wide scale in plant allopolyploids. It may be that this will indeed be revealed with further study, or alternatively, it is also possible that these two mechanisms often work in concert in higher plants, as exemplified by rbcS in some plant polyploids (Gong et al. 2012, 2014).

A final comment concerns the overall higher degree of S. cerevisiae-to-S. eubayanus gene conversion in allotriploid than allotetraploid yeast (fig. 2B). Perhaps this reflects the imbalance in ratios of S. eubayanus (mitochondrial donor) origin (1:2 and 1:1 ratios of S. cerevisiae vs. S. eubayanus homoeologous chromosomes in allotriploid and allotetraploid lager yeast, respectively), thus facilitating more frequent conversions using the extra homoeologs. Because there are plant analogs to this fungal set-up, involving unbalanced hexaploids, for example, formed via doubling of interspecific triploid intermediates, one wonders whether gene conversion will be similarly more strongly biased in these types of allotriploidic plants as they are in this single fungal system. Again, further study of multiple evolutionary systems is needed if we are to develop a general understanding of cytonuclear coevolution in polyploid eukaryotes.

Materials and Methods

Yeasts and Culture Medium
Saccharomyces pastorianus strain Weihenstephan 34/70 was obtained from China General Microbiological Culture Collection Center. The yeast was cultured in the YPD medium (w/v, 1% yeast extract, 2% peptone, and 2% D-glucose) supplemented with 100 μg/mL chloramphenicol.

Data Collection
The reference genomes of model S. cerevisiae, S. eubayanus, and S. pastorianus strains and re-sequencing data were downloaded from National Center for Biotechnology Information (NCBI). Strain names and respective accession numbers are tabulated in supplementary table S1, Supplementary Material online.

Genome Update and de Novo Gene Annotations
Whole-genome re-sequencing raw reads of S. eubayanus strain CDFM21L1 were converted from SRA to FASTQ file format using the fastq-dump command provided by the SRA Toolkit (v2.8.0) with default filter settings (Leinonen et al. 2011). Fastp (v0.20.1) was used to filter the low-quality raw sequencing reads (Chen et al. 2018). Sambamba (v0.7.1) was used to remove PCR duplication (Tarasov et al. 2015). Clean paired reads were mapped to the reference genome (FM1318; Baker et al. 2015) using Bowtie2 (v2.4.2) with default settings (Langmead and Salzberg 2012). These data analysis pipeline were used for the other two diploid strains, S. cerevisiae stout strainNCYC 1063 and S. cerevisiae ale strain FostersO (using S288c as the reference genome; Cherry et al. 1997).

Updates for the genome assembly and annotation of our three diploid strains were obtained using RGAAT (v2.0) (Liu et al. 2018). Concerning gene annotations for S. pastorianus strains WS 34/70 and CBS 1513, we employed the online tool (WebAUGUSTUS, http://bioinf.uni-greifswald.de/webaugsustus/; default parameters) (Stanke et al. 2006) to conduct de novo gene annotation using a training gene set from S. pastorianus CBS 1483 strain (Salazar et al. 2019). BUSCO (v4.1.4) was used to evaluate the completeness of genome assembly and the quality of annotations (supplementary fig. S4, Supplementary Material online) (Simão et al. 2015).

Mitochondrial genomes of S. pastorianus strains of CBS 1513 and WS 34/70 were de novo assembled using GetOrganelle toolkit (v1.7.5) (Jin et al. 2020). Mitochondrial genes (mtDNA genes) of S. pastorianus and S. eubayanus were predicted by the MITOS2 (v2.0.8) pipeline based on the NCBI ReSeq 89 Fungi database and using the yeast genetic code (Bernt et al. 2013).

Construction of Mitochondrial Phylogenetic Tree
To minimize the effect of AT-enrichment and long presence/absence in sequence alignment of whole-genomic mitochondrial DNAs (mtDNAs), mtDNAs of each strain were aligned with S. cerevisiae S288c mtDNA using Snippy (v4.6.0) in contig mode, in which core sites (genomic positions commonly shared in all strains) enclosing monomorphic nucleotide and polymorphic core SNPs will ignore the complications of long presence/absence sequences variant types and possible recombination which are frequently observed in introns and AT/TC clusters. Additionally, its contig mode also discards AT-enriched repeat regions by only maintaining regions covered by uniquely mapping reads (Seemann. T 2015). The multiple alignments were generated using MAFFT (v7.475) (Katoh and Standley 2013). A phylogenetic tree using the Neighbor-Joining method was obtained with MEGA-CC (v11) (Kumar et al. 2012), and visualized using iTOL (https://itol.embl.de/).

Prediction of Nuclear Mitochondria-Targeted Genes
Sequences of nuclear genes encoding S. cerevisiae proteins targeted to subcellular structures of mitochondria (mitochondrial membrane and nonmembrane) were downloaded from the fungal secretome and subcellular proteome knowledgebase (FunSecKB2, http://proteomics.ysu.edu/secretomes/fungi2/index.php) (Meinken et al. 2014). Proteins encoded by NMT genes of all yeasts in this study were identified by BLASTP (v2.6.0) against the downloaded mitochondrial subcellular protein database with similarity greater than 90%, length greater than 65%, and e-value cutoff at 10^{-12} (Camacho et al. 2009).

Calculation of the Nonsynonymous to Synonymous Substitution Ratios (dN/dS)
To assign the genes of S. pastorianus into homoeologous subgenomes (C and E), we used OrthoFinder (v2.5.2)
(default settings) (Emms and Kelly 2019) with BLASTP (v2.6.0) to compare the similarity of amino acid sequences of *S. pastorius* genes with their two parental diploid genes. Single-copy orthologs were identified for all yeast strains with default parameters. A single-copy ortholog involving at least one NMT gene in a certain yeast strain was identified as a single-copy NMT group. Amino acid sequences and CDSs from the same single-copy NMT orthogroup were aligned using ParaAT (v2.0) (Zhang et al. 2012). The $d_{N}/d_{S}$ ratio of each aligned single-copy NMT orthogroup was calculated by KaKs_Calculator package (v2.0) (Wang et al. 2010). Single-copy NMT and non-NMT orthogroups identified in *S. cerevisiae* (S288c, NCYC 1063, and FostersO strains), *S. eubayanus* (FM1318 and CDFM21L.1 strains), and *S. pastorianus* (CBS 1513, WS 34/70, and CBS 1483 strains) were input into above procedure to calculate respective $d_{N}/d_{S}$ ratio in each diploid genome and allopolyploid subgenomes (supplementary table S3, Supplementary Material online). This method was also used for the calculation of $d_{N}/d_{S}$ ratios for aligned mitochondrial genes (mtDNA genes) of *S. eubayanus* and *S. pastorius*.

**Identification of Nonsynonymous Homoeologous Gene Conversions Based on Homologous Gene Quartets**

Subgenomic single-copy homologs in each allopolyploid yeast strain were aligned with their respective parental diploid orthologs identified in the same orthogroup using MAFFT (v7.475) (Katoh and Standley 2013). Single-copy gene quartets with C and E homoeologs and orthologs of *S. cerevisiae* and *S. eubayanus* were generated.

Criteria for defining nonsynonymous gene conversions among homoeologs are described below corresponding to labels 1-8 in fig. 2A: site 1 denotes *S. eubayanus*-to-*S. cerevisiae* amino acid conversion; site 2 indicates *S. cerevisiae*-to-*S. eubayanus* amino acid conversion; site 3 denotes a gap in the alignment that is not considered in conversion analysis; sites 4 and 5 denote the specific nucleotide changes involved in nonsynonymous *S. eubayanus*-to-*S. cerevisiae* gene conversion; site 6 indicates an automorphic nucleotide mutation occurring in an E homoeolog; site 7 denotes a synonymous substitution occurring in the E homoeolog; site 8 is a specific nucleotide change involved in *S. cerevisiae*-to-*S. eubayanus* gene conversion.

**Calculation of Gene Conversion Level**

Conversion level represents the proportion of *S. cerevisiae*-to-*S. eubayanus* nonsynonymous substitutions occupied in the total number of nonsynonymous substitutions in a given single-copy orthogroup, calculated as follows:

Conversion Level = (number of "S. cerevisiae"-to-"S. eubayanus" nonsynonymous substitutions)/(sum of all nonsynonymous substitutions). Orthologs with conversion levels not less than 0.95 were defined as having the high conversion level, while others with conversion levels equal to or less than 0.05 were defined as the low conversion.

**Microaerobic Growth Experiments**

A suitable amount of *S. pastorius* strain WS 34/70 glycerin mixture was initially drawn from a cryopreserved tube and added to 50 mL YPD liquid medium, and grown overnight in a 28°C shaker at 180 rpm to activate the strain. The cultures were evenly streaked on YPD agar plates, which were incubated upside down at 28°C until colonies appeared. Three colonies were selected as biological replications and inoculated in 30 mL YPD liquid medium, respectively. We cultured the strains overnight in a 28°C shaker with 180 rpm. For each biological replication, the yeast strain was inoculated in 37 conical flasks containing 30 mL YPD liquid medium at 1000:1 ratio. 111 conical flasks were cultured at 28°C with shaking at 180 rpm for 84 h. The cultures were sampled for OD$_{600}$ measurements and RNA extraction at the following time points: every 1.5 h before 27 h, and every 3 h after 27 h. Three conical flasks were taken out as sample replications each time. Each sample was centrifuged at 5000 × g for 1 min at room temperature until 0.3 g yeast cells was collected in 1.5 mL microcentrifuge tube. Accordingly, yeasts were sampled at 37 time points in total for each replicate (supplementary table S4, Supplementary Material online).

**RNA Extraction and Transcriptomic Sequencing**

For each biological replicate, 500 μL TriPure Isolation Reagent (Roche Diagnostics GmbH, Mannheim, Germany) and 0.3 g acid-washed glass beads (SIGMA, USA) rapidly were added. After vibrating on the vortex oscillator at the maximum speed for 1 min, samples were put into liquid nitrogen for rapid freezing. This freezing and thawing vibration was repeated five times. After complete melting, the tube was incubated on ice for 5 min to lyse the yeast cells. The total RNA was purified using UNIQ-10 Column Trizol Total RNA Isolation Kit (Sangon Biotech, Shanghai, China).

RNA was treated by DNase I and purified using the RNA clean Kit (Qiagen, Germany). High-quality RNA was used for the subsequent library constructions. Ribosomal RNA (rRNA) was removed by Epicentre Ribo-zeroTM rRNA Removal Kit (Epicentre, USA), and rRNA free residue was cleaned up by ethanol precipitation. Strand-specific sequencing libraries were constructed by the NEBNext Ultra Directional RNA Library Prep Kit for Illumina (NEB, USA). The resulting library was sequenced on the Illumina Nova 6000 platform in PE150 bp mode at the Novogene Company (Beijing, China).

**Differentially Expressed Genes Analysis**

Fastp (v0.20.1) was used to filter the low-quality raw sequencing reads (Chen et al. 2018). The reads from ribosome RNA were removed using Bowtie2 (v2.4.2) (Langmead and Salzberg 2012) based on SILVA (https://www.arb-silva.de/) (Quast et al. 2012) and Pfam (Mistry...
Clean reads were mapped to the S. pastorianus strain WS 34/70 reference genome (Okuno et al. 2016) using HISAT2 (v2.2.1) (Kim et al. 2019). The reads counts of each sample were quantified by featureCounts (v2.0.1) with the parameters set as “-p -s 2” (Liao et al. 2014). Differentially expressed genes (DEGs) in two subgenomes (S. eubayanus homoeologs as the numerator and S. cerevisiae homoeologs as the denominator) were identified using DESeq2 (v1.30.1) based on false discovery rate adjusted P value < 0.05. Clust (v1.12.0) was employed to cluster the subgenomic DEGs in terms of their similarity in expression pattern in the temporal course of the lag and exponential phases (T1 to T17) (Abu-Jamous and Kelly 2018).

### Gene Ontology Analysis

GO term enrichment analysis was performed using the compareCluster command provided by the R package clusterProfiler (v3.18.1). All genes of S. pastorianus strain WS 34/70 were used as the reference background.

### Calculation of Transcriptional Expression Bias for Saccharomyces eubayanus Homoeologs

The proportion of highly expressed S. eubayanus homoeologs was calculated as follows:

\[ \text{Transcriptional expression preference for } S. \text{ eubayanus} \text{ NMT homoeologs} = \frac{\text{Number of groups with significantly higher expression in } S. \text{ eubayanus homoeologs}}{\text{Sum of total DEGs groups}}. \]

### Supplementary Material

**Supplementary data** are available at Molecular Biology and Evolution online.

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### Author Contributions

Conceptualization: B.L. and L.G., Methodology: Z.Z. and T.W., Investigation: K.Z., J.L., G.L. and Y.Z., Experiments: K.Z., J.L., Y.Z., W.S., J.W., J.Y. and Y.M., Data Analysis: K.Z., J.L., Y.Do. and H.W., Supervision: L.G., Writing—original draft: K.Z. and J.L., Writing—review & editing: J.F.W., B.L., and L.G.

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### Data Availability

RNA-sequencing data have been deposited and are available in the NCBI (PRJNA808674). All other data are available in the paper and/or the Supplementary files.

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