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Eukaryotic chromosomes are organized into structural and functional domains with characteristic replication timings, which are thought to contribute to epigenetic programming and genome stability. Differential replication timing results from epigenetic mechanisms that positively and negatively regulate the competition for limiting replication initiation factors. Histone deacetylase Sir2 negatively regulates initiation of the multicopy (~150) rDNA origins, while Rpd3 histone deacetylase negatively regulates firing of single-copy origins. However, Rpd3's effect on single-copy origins might derive indirectly from a positive function for Rpd3 in rDNA origin firing shifting the competitive balance. Our quantitative experiments support the idea that origins compete for limiting factors; however, our results show that Rpd3's effect on single-copy origin is independent of rDNA copy-number and of Sir2's effects on rDNA origin firing. Whereas RPD3 deletion and SIR2 deletion alter the early S phase dynamics of singlecopy and rDNA origin firings in opposite fashion, unexpectedly only RPD3 deletion suppresses overall rDNA origin efficiency across S phase. Increased origin activation in $rpd3\Delta$ requires Fkh1/2, suggesting that Rpd3 opposes Fkh1/2-origin stimulation, which involves recruitment of Dbf4-dependent kinase (DDK). Indeed, Fkh1 binding increases at Rpd3-regulated origins in $rpd3\Delta$ cells in G1, supporting a mechanism whereby Rpd3 influences initiation timing of single-copy origins directly through modulation of Fkh1-origin binding. Genetic suppression of a DBF4 hypomorphic mutation by RPD3 deletion further supports the conclusion that Rpd3 impedes DDK recruitment by Fkh1, revealing a mechanism of Rpd3 in origin regulation.

Replication origin | Chromatin domains | Genome instability | Repetitive DNA | Histone deacetylase

The replication of functionally distinct chromosomal regions presents challenges to genome stability. For example, repeated and highly expressed DNA sequences, such as the ribosomal DNA genes, appear to be destabilized by transcription-replication collisions and unequal sister chromatid exchange or intrachromosomal recombination (reviewed in 1). In Saccharomyces cerevisiae, the ribosomal DNA (rDNA) gene cluster consists of about 150 direct repeats of the ~9.1-kbp rDNA sequence, each containing one potential origin (reviewed in 2), which represents a significantly higher origin density than the genome average among unique sequences (reviewed in 3). Thus, rDNA origins have the potential to impose a significant load on the replication system and impede the replication of other chromosomal regions, as several replication-initiation factors are present in limiting quantities (reviewed in 4). However, only ~20% of the rDNA origins fire in each S phase in wild-type (WT) cells (5, 6), and this has been attributed to an intrinsically weak autonomously replicating sequence (ARS) (7, 8), and to suppression of origin firing by the histone deacetylase Sir2 (9). Deletion of SIR2 results in a higher proportion of the rDNA origins initiating replication, which has been suggested to contribute to instability of the rDNA repeats. Histone deacetylase Rpd3, in contrast, has been implicated in the delayed initiation of late-firing, singlecopy origins dispersed throughout the genome. RPD3 deletion advances the firing of many late origins while increasing the levels of histone acetylation at these origins (10-12). Targeting of Gcn5 histone acetylase to a late replication origin advanced its timing, supporting the idea that histone acetylation regulates initiation rate (12). However, specific DNA-binding, protein-mediated recruitment of Rpd3 to origins has not been demonstrated (nor ruled out), so Rpd3 has been suggested to act directly at singlecopy origins through an untargeted (or "global") interaction mechanism (10, 12).

More recently, however, this model of Rpd3 function has been challenged by a study claiming that Rpd3 acts directly to stimulate rDNA origins and that effects of *RPD3* deletion on single-copy origins are an indirect consequence of reduced competition from rDNA origins for a limited pool of replication-initiation factors (13). Indeed, recent studies have identified several initiation factors as rate limiting for origin firing, leading to models in which unequal competition for limiting factors among all licensed

Significance

The faithful replication of eukaryotic genomes requires balancing the replication capacities of different genomic regions, such as repetitive versus single-copy genetic elements, which may compete for limiting replication resources, possibly leading to replication stress and genome instability. We examined the function of histone deacetylases Rpd3 and Sir2 in balancing replication between unique genome sequences and the multicopy ribosomal DNA genes. Our findings support prior conclusions that Sir2 directly suppresses early firing of rDNA origins, thereby enabling balanced replication of the genome. We further show that Rpd3's function in delaying firing of later-firing, single-copy origins is independent of Sir2 and rDNA load. Instead, Rpd3 appears to oppose the Fkh1/2 origin activation pathway by regulating binding of the origin-stimulator Fkh1.

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origins underlies their differential activities (14–16). In the Yoshida study, deletion of *SIR2* increased early firing of rDNA origins while decreasing early firing of single-copy origins, consistent with the competition model. Additional deletion of *RPD3* suppressed the effects of *SIR2* deletion, consistent with these factors acting in opposition at the rDNA. Thus, it was concluded that advanced firing of single-copy origins in $rpd3\Delta$ cells is a consequence of reduced Rpd3-stimulated rDNA origin firing competing for limiting factors, rather than a direct repressive effect of Rpd3 on single-copy origins. A previous study implicating Rpd3 in transcriptional silencing of the rDNA was also cited as supporting evidence for the idea that Rpd3 acts directly on rDNA origins (17).

The main conclusions of the earlier study relied on quantitative comparisons of independently generated bromodeoxyuridineimmunoprecipitation-sequencing (BrdU-IP-seq) datasets with WT, sir2 Δ , rpd3 Δ , and sir2 Δ rpd3 Δ strains. Such comparisons require signal normalization between the independently produced experimental datasets, which can be problematic, typically requiring arbitrary, if not dubious, assumptions about the data. More recently, we developed quantitative BrdU-seq (gBrdU-seq) (or "QBU") to overcome this barrier to an unambiguous, quantitative comparison of independent BrdU-IP-seq samples (18). Our method involves sample-specific barcoding of BrdU-labeled genomic DNA, followed by pooling for IP, amplification, and highthroughput sequencing. Additionally, prior to IP, a small aliquot of the pooled "input" sample is retained for amplification and sequencing in parallel, providing a direct reference for normalization of IP sequencing read counts based on actual input. QBU eliminates and/or corrects for most sources of technical or experimental error that may hinder (or at least cast doubt upon) guantitative comparison between individually processed samples.

In this study, we have re-examined the impacts of Rpd3 and Sir2 on genome replication, with particular focus on the origin competition model and Rpd3 function at single-copy versus rDNA origins. Our findings support the idea of competition between rDNA and single-copy origins, but not the notion that Rpd3 acts positively through rDNA origins to squelch single-copy origins. We have also probed more deeply into the mechanism of Rpd3 action at origins and present evidence that Rpd3 antagonizes single-copy origin firing by regulating Fkh1 binding and, hence, recruitment of Dbf4-dependent kinase (DDK) to origins. In accord, deletion of *RPD3* rescues defective DDK function.

Results

Rpd3 Has Differential Effects on Single-Copy Origins. A rigorous test of the competition model between rDNA and singlecopy origins and of the roles of Rpd3 and Sir2 in regulating these origins requires an unambiguous, quantitative comparison of origin firing levels. Therefore, we performed QBU analysis of WT, $rpd3\Delta$, $sir2\Delta$, and $sir2\Delta rpd3\Delta$ strains. Cells were synchronized in G1 phase with α -factor and released into S phase in the presence of hydroxyurea (HU) to block replication and distinguish between early origins, which fire efficiently in HU, and late origins, which do not due to intra-S checkpoint inhibition (19). DNA content analysis of cells similarly synchronized and released into S phase without HU shows similar overall timing of S phase execution for the four strains, with $rpd3\Delta$ cells completing S phase slightly more rapidly than WT, as observed previously (*SI Appendix*, Fig. S1A) (12).

Distribution of the QBU values comparing the individual replicates of each strain shows high correlations, demonstrating the high experimental reproducibility of the QBU analysis (SI Appendix, Fig. S1B). Chromosomal plots of the QBU data show qualitatively similar results as observed previously in comparison of WT and rpd31 cells, with increased BrdU incorporation at several single-copy, later-firing origins in $rpd3\Delta$ cells (Fig. 1A). We determined the origin loci with significant BrdU incorporation and identified 235 in WT cells and 301 in $rpd3\Delta$ cells; a Venn diagram shows that almost all the origins identified in WT cells were also identified in $rpd3\Delta$ cells (Fig. 1B). Origins were assigned to timing quartiles based on their previously determined replication timings (T_{Rep}) among 625 confirmed or likely origin in the oriDB (20) and displayed as stack graphs of origin counts and pie charts of origin proportions (Fig. 1C). These results show that almost all of the additional origins detected to fire in $rpd3\Delta$ cells fall into later (Q2, Q3, and Q4) timing quartiles (in WT cells), consistent with Rpd3 acting to delay origin firing.

As BrdU signals at origins in HU-blocked cells inversely correlate with T_{Rep} (11), we also examined the relationship between the effect of RPD3 deletion and origin timing by plotting QBU signals for individual origins in $rpd3\Delta$ cells against their corresponding values in WT cells (Fig. 1D). The two-dimensional scatter plot reveals a relationship between WT BrdU levels and the effect of RPD3, with origins having higher QBU values in WT being decreased in $rpd3\Delta$ cells and origins having lower QBU values in WT being increased in rpd31 cells. To directly examine the relationship between the QBU values and origin timing, we plotted average QBU values for origins divided according to their assigned T_{Rep} quartiles (Fig. 1*E*). The results show that, in *rpd3* Δ cells, QBU values are lower for origins in the earliest T_{Rep} quartile, while QBU values are slightly higher for origins in the later quartiles, although only the difference in the first quartile is statistically significant (SI Appendix, Fig. S1 C). We also examined the effects of RPD3 deletion on specific origin classes by highlighting these in the scatter plots (Fig. 1D) and by producing distribution box plots for those groups (Fig. 1F); t tests of the distributions show that centromere (CEN)-proximal origins, which are early firing, are significantly decreased in activity, while Rpd3-repressed origins, which are later firing, show significantly increased replication activity in $rpd3\Delta$ cells (Fig. 1F). The relatively minor but significant reduction in early origin firing in $rpd3\Delta$ cells has not been previously reported, likely because normalization methods obscured the difference. The opposite effect on earlier versus later origins is consistent with the idea that origins compete for limiting factors and that increased competition from normally later-firing origins in $rpd3\Delta$ cells consequently reduces activation of normally earlier-firing origins.

In contrast to the results in $rpd3\Delta$ cells, the replication profiles of $sir2\Delta$ cells showed decreased BrdU incorporation at virtually all single-copy origins (Fig. 1 *A*, *D*–*F*), though the effect was smaller among origins with higher BrdU incorporation values (Fig. 1*D*). Only 192 origins were called as active in $sir2\Delta$ cells, and there was a decrease in origins called in each T_{Rep} quartile (Fig. 1*C*). Overall, these results suggest that Sir2 acts nonspecifically with respect to single-copy origins because almost all single-copy origins are decreased in activity.

Combined deletion of *SIR2* and *RPD3* resulted in a replication profile intermediate to the single-mutant strains, and similar to WT (Fig. 1 A and D), as supported by correlation analysis (*SI Appendix*, Fig. S1D). Two hundred twenty-six origins were detected in $rpd3\Delta$ sir2 Δ cells with almost all these origins also being detected in WT cells and having a similar T_{Rep} distribution as in WT cells (Fig. 1 B and C). Overall, QBU signals at individual origins were also more like WT than either single mutant, although more origins were reduced in activity as in sir2 Δ cells



Fig. 1. Rpd3 and Sir2 have different effects on single-copy origins. Strains CVy43 (WT), CVy44 ($rpd3\Delta$), YHy3 ($sir2\Delta$), and YHy6 ($sir2\Delta rpd3\Delta$) were synchronized in G1 phase and released into S phase without HU for DNA content analysis (*SI Appendix*, Fig. S1A) or with HU for 60 min for QBU analysis. Strains with reduced rDNA copies PP1758 (WT *rDNA20*) and JPy115 ($rpd3\Delta rDNA20$) were synchronized in G1 phase and released into S phase for 120 min in the presence of HU for QBU analysis. (A) QBU values averaged for three replicates and scale normalized across strains are shown for representative chromosome XI; origins and origin subgroups are indicated with color-coded circles below the x axis. (B) Venn diagrams depicting overlaps between origins identified as active in each strain; the union of 311 origins detected was used in subsequent analyses. (*C*) Origin distributions according to T_{Rep} quartiles shown as pie charts with proportions of identified origins in each strain and as stack graphs showing total counts of origins identified in each T_{Rep} quartile. (*D*) 2D scatter plots comparing QBU signals for 500-bp regions centered on 311 individual origins plus the rDNA origins, which are represented by two (overlapping) data points; origins and subgroups are color coded as indicated. (*E*) Average QBU signals for 311 origins detected with in parentheses; two-sided t tests were performed on all pairs of strains, and significant results are indicated as *P < 0.05 and **P < 0.01.

than increased in activity as in $rpd3\Delta$ cells (Fig. 1 D and F and SI Appendix, Fig. S1D). Comparison of QBU origin signals between $rpd3\Delta$ sir2 Δ and sir2 Δ cells and between $rpd3\Delta$ sir2 Δ and $rpd3\Delta$ cells shows that RPD3 deletion in sir2 Δ cells and SIR2 deletion in $rpd3\Delta$ cells affect origins similarly as they do in WT cells, consistent with RPD3 and SIR2 acting through independent mechanisms (Fig. 1 D and F and SI Appendix, Fig. S1D).

SIR2 Deletion Is Epistatic to *RPD3* Deletion for rDNA Origin Firing. Next, we examined the effects of *RPD3* and *SIR2* deletion on initiation levels of rDNA origins. In WT cells, the QBU signal for rDNA origins (QBU corrects for copy-number so the signal represents the average firing level per rDNA copy) was between that of single-copy origins in the first and second timing quartiles (Fig. 2A, compare with Fig. 1E), consistent with the average mid-S phase peak replication timing of these origins (18). In *rpd3* Δ cells, the rDNA QBU signal was strongly diminished (Fig. 2 A and B). In contrast, the rDNA QBU signal in *sir2* Δ cells was about twice the level in WT cells (Fig. 2 A and B), in rough agreement with previous reports on the effect of *SIR2* deletion on rDNA origin firing levels (9, 13). Unlike the additive effect of double deletion of *RPD3* and *SIR2* on single-copy



Fig. 2. *SIR2* deletion is epistatic to *RPD3* deletion for rDNA origin firing. See Fig. 1 legend for strains and experimental description. (*A*) Average QBU signal for the rDNA depicted as a single locus. (*B*) Cumulative ($^{+/-}$ SD) QBU signal (500-bp windows) for 311 single-copy origins (*Left panel*) and 100 rDNA origins (*Center panel*) was determined and plotted; two-sided *t* tests were performed on all pairs of strains, and significant results are indicated as **P* < 0.05 and ***P* < 0.01. Total QBU signal associated with rDNA origins glus the rDNA origins (*Right panel*). (*C*) 2D gel electrophoresis analysis probing for the rDNA was performed on the strains as described above. Two replicate sets are shown; the images framed with broken lines are darker images of the corresponding *rpd3*Δ image in each set. The arrows in the WT panels indicate the bubble arcs (filled arrowhead) and 1N spots (unfilled arrowhead) that are used for quantification (see *SI Appendix*, Fig. S2*B* for images with exact areas measured for quantification). The graph indicates quantification of the 2D gels by measuring intensity of "bubble arcs" in relation to the corresponding 1N spots; values are relative to WT arbitrarily set to 1.

origins, the rDNA QBU signal in $rpd3\Delta$ sir2 Δ was almost identical to the sir2 Δ cells (Fig. 2 A and B). This result suggests that SIR2 deletion is epistatic to, not additive with, RPD3 deletion with respect to rDNA origin firing, differing from previous conclusion (13). Whereas the significantly reduced early firing of rDNA origins in $rpd3\Delta$ cells is consistent with the idea that Rpd3 is a direct stimulator of rDNA origin firing, the high level of rDNA origin firing in sir2 Δ rpd3 Δ cells conflicts with this idea. Moreover, the effect of RPD3 deletion on single-copy origins in $rpd3\Delta$ sir2 Δ cells (Fig. 1D) occurred independently of any effect on rDNA origin firing (Fig. 2B), suggesting a direct effect of Rpd3 on single-copy origins.

We also quantified QBU signal from single-copy origins versus rDNA origins, after confirming that all four strains had similar rDNA copy counts, estimated at ~100 in WT (SI Appendix, Fig. S2A). BrdU incorporation into single-copy origins was significantly lower in the sir2 Δ and sir2 Δ rpd3 Δ strains versus WT, while $rpd3\Delta$ cells were like WT (Fig. 2B). We also calculated the absolute proportion of rDNA origin activity as a proportion of total genome origin activity for each strain, which shows the substantial shifts in balance between replication of the rDNA in relation to the genome resulting from deletion(s) of RPD3 and SIR2 (Fig. 2B, Right panel). Overall, the finding that *RPD3* deletion in *sir2* Δ cells affects single-copy (Rpd3-repressed) origins without affecting rDNA origins argues against the idea that the effect of RPD3 deletion on single-copy origin firing derives indirectly from diminished rDNA origin firing.

To confirm that the results of QBU at the rDNA accurately reflect origin initiation levels, we used two-dimensional agarose gel (2D gel) electrophoresis to directly examine replication initiation structures at rDNA origins. As above, cells were synchronized in G1 phase with α -factor and released into S phase in the presence of HU to evaluate early replication events. Comparison of the intensity of replication "bubble arcs" representing initiation structures versus the 1N spots representing total molecules yields a relative measurement of origin firing activity (Fig. 2*C* and *SI Appendix*, Fig. S2*B*). The 2D gel results mirror the QBU results with rDNA origin firing decreased significantly in *rpd3* Δ cells, increased significantly in *sir2* Δ cells, and similarly increased in *sir2* Δ *rpd3* Δ cells (Fig. 2*C* and *SI Appendix*, Fig. S2*B*). These results reinforce the conclusion that *SIR2* deletion is epistatic to, not additive with, *RPD3* deletion with respect to rDNA origin activation.

RPD3 Regulates Single-Copy Origins Independently of the rDNA Array. Yoshida et al. elegantly tested their model of competition between rDNA and single-copy origins by evaluating the effect of a reduction in rDNA copies on the suppression of single-copy origin firing resulting from *SIR2* deletion, the premise being that fewer rDNA origins would create less competition. In support of the competition model and a role for Sir2 in regulating the rDNA directly and single-copy origin indirectly, they showed that, in a strain with only ~20 copies of rDNA, the effect of *SIR2* deletion on single-copy origins was largely suppressed, as predicted. We adopted the same rationale and approach to test whether Rpd3 acts on single-copy origins through effects on the rDNA by evaluating the effect of *RPD3* deletion in the strain background with the reduced rDNA copies; we confirmed that the strains with reduced



Fig. 3. Rpd3 suppresses rDNA origin firing. (A) Strains described in Fig. 1 legend were grown logarithmically and harvested for 2D gel analysis and probing for the rDNA. Three replicate sets are shown. The arrows in the WT panels indicate the bubble arcs (filled arrowhead) and early Y-arcs (unfilled arrowhead) that are used for quantification (see *SI Appendix*, Fig. S3A for images with exact areas measured for quantification). The graph shows quantification of the 2D gels by measuring intensity of bubble arcs in relation to the corresponding Y-arcs (replication fork structures); values are relative to WT arbitrarily set to 1; two-sided *t* tests were performed on all pairs of strains, and results are indicated as *P < 0.05 and *P < 0.01. (B) WT, rpd3A, and *sir2* strains described above were synchronized in G1 phase, released into S phase, and incubated with BrdU for the following time intervals for QBU analysis: pulse 1 = 15–30 min, pulse 2 = 25–40 min, pulse 3 = 35–50 min, and pulse 4 = 45–60 min; 5-min overlap is included to account for lag time in BrdU entry to cells and incorporation into DNA. The stack graphs show total QBU signal for 625 potential single-copy origins (*Left*) and total QBU signal for all rDNA origins during each time interval. (C) Box plot distributions of QBU signals (500-bp regions) in the first pulse interval of the time course for the indicated origin groups.

rDNA copies contained about 25% of the rDNA complement as the normal copy strains (SI Appendix, Fig. S2A). Chromosome plots show that similar derepression of dormant/late-firing origins occurred in $rpd3\Delta$ cells compared with WT cells in the strains with ~ 20 rDNA copies (Fig. 1*A*). Analysis of QBU values for individual origins in the scatter plots revealed a similar relationship of RPD3 deletion on origins according to their QBU levels, with the greatest increases among those origins that normally have low QBU values, reflecting later-firing origins (Fig. 1D). Distribution box plots of QBU signals affirm the global increase in firing of Rpd3-repressed origins in $rpd3\Delta$ cells relative to WT (Fig. 1F), and correlation analysis demonstrates closer similarity of the $rpd3\Delta$ strain with reduced rDNA copies to the $rpd3\Delta$ strain with normal rDNA than the WT strain with normal rDNA (SI Appendix, Fig. S1D). Notably, rDNA origin activity level was only modestly reduced relative to WT in the $rpd3\Delta$ rDNA 20 strains. These results indicate that the effect of RPD3 deletion on single-copy origins does not rely on repression of a large array of rDNA origins. Instead, the findings support the conclusion that Rpd3 regulates origin firing directly at single-copy origins, rather than indirectly through the rDNA.

RPD3 Suppresses Pan-S Phase rDNA Origin Efficiency. The analyses above were all conducted in the presence of HU to evaluate the levels of origin firing occurring in an early S phase interval. To determine the impacts of *RPD3* and *SIR2* on the overall efficiency of rDNA origin firing throughout the duration of S phase, we used 2D gels to examine and quantify replication intermediates at the rDNA origins in unperturbed cycling cells (pan-S phase); here, we compared the intensities of initiation bubble structures with replication fork structures to

provide a relative measure of origin efficiency (Fig. 3A and SI Appendix, Fig. S3A). In striking contrast to the 2D gel results in early S phase (HU) described above, pan-S phase $rpd3\Delta$ cells exhibited modest but significantly increased overall efficiency of rDNA origin firing relative to WT cells (Fig. 3A). The overall increased firing of rDNA origins in $rpd3\Delta$ cells is inconsistent with the notion that RPD3 is a direct activator of rDNA origins. Interestingly, $sir2\Delta$ cells showed no difference in pan-S phase rDNA origin efficiency compared with WT (Fig. 3A), which contrasts with a previous report (9). Moreover, $rpd3\Delta$ sir2 Δ cells showed similar pan-S phase rDNA origin efficiency as $sir2\Delta$ and WT cells (Fig. 3A). Thus, the striking, and opposite, effects of SIR2 deletion and RPD3 deletion on rDNA origin firing levels in early S phase are not reflected in their overall efficiencies through the duration of S phase.

The differences in origin firing in the early-S versus pan-S phase analysis, particularly in $rpd3\Delta$ cells, suggests that alterations in rDNA origin usage early are subsequently compensated. For example, more early-firing of normally later-firing, single-copy, Rpd3-regulated origins should leave fewer such origins unfired in late S phase in competition with unfired rDNA origins, thus shifting the competitive balance. To more precisely examine the temporal dynamics of replication in the rDNA versus single-copy origins in $rpd3\Delta$ cells, we analyzed BrdU incorporation through a time series of pulse labeling in cells progressing synchronously through S phase. This analysis also eliminates possible perturbations due to HU. The analysis shows relative levels of BrdU incorporation at single-copy versus rDNA origins over the time course (Fig. 3B). Replication origin usage profiles are similar in the initial time pulse as in HU (SI Appendix, Fig. S3B). As expected, rDNA replication occurs throughout S phase in WT cells. In rpd3A cells, the relative proportions of replication at rDNA versus single-copy origins shift in favor of single-copy origins in earlier pulse periods and, conversely, in favor of rDNA origins in the later periods (Fig. 3*B*). The results are consistent with the idea that reduced usage of rDNA origins in early S phase preserves more rDNA origins to initiate later, likely with fewer unfired, singlecopy origins in competition. Analysis of single-copy origin firing in the earliest interval shows significantly increased activity of Rpd3-repressed origins, showing that the effect of *RPD3* deletion is similar in early S phase with or without HU (Fig. 3*C*).

Although deletion of *SIR2* had no effect on overall origin efficiency in pan-S phase analysis, examination of BrdU incorporation over time shows a striking shift toward earlier replication of the rDNA in *sir24* compared with WT cells (Fig. 3*B*), consistent with the early S phase results in HU. *SIR2* deletion also results in notably delayed replication of the total genome, while *RPD3* deletion's effect on the genome is minor and in the same direction as its effect on the rDNA in the earliest pulse (Fig. 3*B*). We also note that this type of global BrdU incorporation analysis does not distinguish active versus passive replication of origin sequences.

Origin Derepression in $rpd3\Delta$ **Cells Requires Fkh1/2.** *FKH1* and/or *FKH2* (*FKH1/2*) are responsible for the early activation of most non-CEN-proximal, early-firing origins throughout the genome (21). Fkh1 promotes early origin firing by recruiting limiting factor Dbf4 to stimulate execution of the DDKdependent Sld3-Cdc45 loading step of replication initiation (22, 23). Overexpression of Fkh1 stimulates earlier firing of many later-firing origins, including Rpd3-repressed origins (*SI Appendix*, Fig. S4A) (18), suggesting an antagonistic relationship between Fkh1 and Rpd3. Because Fkh1 levels can modulate firing of Rpd3-repressed origins, we tested whether Fkh1 protein level was altered in $rpd3\Delta$ cells and found no difference relative to WT (*SI Appendix*, Fig. S4*B*). To gain further insight into the relationship between Rpd3 and Fkh1/2 function in replication timing, we tested for genetic interaction by determining the effect of their combined mutation. We performed QBU with WT, $rpd3\Delta$, $fkh1\Delta$ fkh2-dsm, and $rpd3\Delta$ $fkh1\Delta$ fkh2-dsm strains; the fkh2-dsm allele was used rather than $fkh2\Delta$ because it is virtually null in replication timing control but functional in regulating gene expression and, therefore, does not, in combination with FKH1 deletion, exhibit more-severe phenotypes of $fkh1\Delta$ $fkh2\Delta$ cells, such as pseudohyphal growth, which complicates analysis and potential interpretation (24). QBU values comparing the individual replicates of each strain are well correlated (*SI Appendix*, Fig. S4*C*).

The results show that derepression of Rpd3-repressed origins in $rpd3\Delta$ cells was largely eliminated in $rpd3\Delta$ fkh1 Δ fkh2-dsm cells (Fig. 4). Analysis of Rpd3-repressed origins shows no significant increase in BrdU incorporation in $rpd3\Delta$ fkh1 Δ fkh2dsm versus fkh1 Δ fkh2-dsm cells (Fig. 4B). This finding suggests that Rpd3 opposes origin stimulation by Fkh1/2. As origin stimulation by Fkh1/2 involves its direct binding to DNA sequences at origins, we posited that repression by Rpd3 likely involves direct effect on the origin.

Rpd3 Modulates Fkh1 Binding to Origins. Given the opposing activities of Fkh1/2 and Rpd3 in origin activation, we considered the possibility that Rpd3 might regulate binding of Fkh1/2 to origins, particularly Rpd3-regulated origins, including the possibility of differential regulation of Fkh1/2 binding to rDNA origins versus single-copy origins. Our previous studies have shown Fkh1/2 binding to Fkh-activated origins in G1 phase (25). In that study, we also observed more-extensive origin binding in S-phase-arrested cells, suggesting that Fkh1/2



Fig. 4. Origin derepression in $rpd3\Delta$ cells requires Fkh1/2. Strains CVy43 (WT), CVy44 ($rpd3\Delta$), OAy1114 ($fkh1\Delta$ fkh2-dsm), and YHy13 ($fkh1\Delta$ fkh2-dsm $rpd3\Delta$) were synchronized in G1 phase and released into S phase with HU for 90 min for QBU analysis. (A) Average QBU values are shown for representative chromosome XI; origins and origin subgroups are indicated with color-coded circles below the x axis. (B) Box plot distributions of QBU counts for 500-bp regions aligned on origins of the indicated subgroups; n of origins in each group is indicated within parentheses; two-sided t tests were performed on all pairs of strains, and results are indicated as *P < 0.05 and **P < 0.01. (C) 2D scatter plots comparing QBU signals centered on 311 individual origins plus the rDNA origins represented by two (overlapping) data points; origins and subgroups are color coded as indicated.

may bind additional origins during S phase to stimulate activation of later origins. Therefore, we hypothesized that Rpd3 regulates Fkh1 binding to later-firing origins in G1 and/or early S phase and performed chromatin immunoprecipitation sequencing (ChIP-seq) of Fkh1-3xFLAG in WT and rpd3A cells synchronized in G1 to test this idea. Comparison of ChIP signal from experimental replicates shows high correlation, demonstrating high reproducibility of the data (SI Appendix, Fig. S5A). In WT cells in G1, heatmaps of average ChIP signal show local Fkh1 enrichment at previously identified Fkh1 binding sites, including Fkh-activated origins, as expected, as well as some enrichment at Rpd3-repressed origins, previously unnoticed (Fig. 5A). In rpd3A cells in G1, Fkh1 profiles are similar; however, box plot analysis and statistical testing show there is a modest but significant increase in Fkh1 enrichment at Rpd3-repressed origins, but not at other origin groups or Fkh1 binding sites more generally (Fig. 5 A and B). Regression analysis of the ChIP enrichment values for Rpd3-repressed origin correlates with the increased activation of Rpd3-repressed

origins in early S phase in $rpd3\Delta$ cells (Fig. 5C). These results are consistent with the idea that the mechanism of Rpd3 in regulation of origin initiation is to oppose origin binding of Fkh1.

Fkh1 binding at origins occurs in G1 and S phases and is dependent on the prereplication complex (pre-RC), defined by origin-loaded MCM complexes (25, 26). Thus, a plausible mechanism for the increased Fkh1 binding to Rpd3-repressed origins in $rpd3\Delta$ cells might derive from an effect of Rpd3 on pre-RC levels. To address this possibility, we performed ChIPseq of Mcm4-3xHA in WT and $rpd3\Delta$ cells synchronized in G1. The results show robust enrichment for Mcm4 at origins of various classes, as expected; however, no significant difference in Mcm4 enrichment is detected at Rpd3-repressed origins or at the other groups examined (*SI Appendix*, Fig. S5). Taken together, these results indicate that Rpd3 modulates Fkh1 binding without altering pre-RC levels and are consistent with the idea that Rpd3 acts downstream of pre-RC assembly to impede Fkh1 binding.



Fig. 5. Rpd3 modulates Fkh1 binding to origins. Strains OAy1112 (*FKH1-3xFLAG*) and YHy17 (*rpd3* Δ , *FKH1-3xFLAG*) were synchronized in G1 phase and subjected to ChIP-seq. (A) Heatmaps of ChIP-seq enrichment at selected origin and Fkh1 binding loci; n of each group is indicated within parentheses. The "non-origins" group refers to Fkh1-binding sites called as peaks by MACS in the Fkh1 ChIP-seq dataset that do not overlap with a replication origin. (*B*) Box plots distributions of ChIP-seq enrichments for 500-bp regions aligned on origins of the indicated subgroups; n of origins or Fkh1 binding loci in each group is indicated within parentheses. Two-sided *t* tests were performed on all pairs of strains, and results are indicated as **P* < 0.05 and ***P* < 0.01. (*C*) 2D scatter plots of ChIP-seq signals (500-bp regions) for 174 Fkh1-binding sites (called as peaks by MACS as described above but including origins) plus the indicated origin subgroups, color coded as indicated. A linear regression deriving the best-fit line is shown for each origin group.

RPD3 Deletion Suppresses DBF4 Deficiency Implicating the Fkh1-DDK Pathway. As a further test of our hypothesis that Rpd3 acts in opposition to Fkh1/2-dependent stimulation of Dbf4 recruitment, we tested whether deletion of RPD3 suppresses the temperature sensitivity of a carboxyl-terminally truncated DBF4 allele ($dbf4\Delta C$) (27). Deletion of FKH1 (or FKH1 and FKH2) does not cause temperature sensitivity, indicating that Dbf4 Δ C is defective in function beyond its proposed defect in interaction with Fkh1. In attempting to isolate a fkh1 Δ dbf4 Δ C strain through meiosis and sporulation of a heterozygous diploid, we determined that *FKH1* deletion causes lethality in combination with $dbf4\Delta C$ (at normally permissive temperature for $dbf4\Delta C$, strongly suggesting that Dbf4 ΔC depends on Fkh1 for recruitment to origins (Fig. 6A). Thus, we created a diploid heterozygous for RPD3/rpd31 and DBF4/ $dbf4\Delta C$ and induced meiosis and sporulation at permissive temperature. Individual spores from mature tetrads were dissected onto rich medium, incubated at 30 °C (restrictive temperature for $dbf4\Delta C$, and imaged and genotyped. The results show that *RPD3* deletion suppresses the lethality of $dbf4\Delta C$ at 30 °C, as virtually no $dbf4\Delta C$ segregants were recovered while the expected quantity of $rpd3\Delta \ dbf4\Delta C$ segregants were recovered, although these formed much smaller colonies than WT or $rpd3\Delta$ isolates (Fig. 6B). Out of 13 tetrads dissected, 37 viable spores were genotyped, resulting in 15 WT, 9 $rpd3\Delta$, 12 $rpd3\Delta$ $dbf4\Delta C$, and 1 $dbf4\Delta C$ viable isolates. Fisher's exact test indicates that the RPD3 and DBF4 genotypes being tested are not independent (P = 0.00164), strongly supporting the notion that loss of Rpd3 function restores viability of Dbf4-deficient cells, likely by facilitating Dbf4 Δ C recruitment to origins by Fkh1.

Discussion

A full understanding of genome-wide replication dynamics requires the application of methods that accurately quantify the process and that may be accurately compared across independent samples. The QBU method has now allowed us to re-examine how the Sir2 and Rpd3 histone deacetylases affect genome replication, particularly the balance between dispersed, single-copy origins and the tandemly repetitive rDNA origins, which has been identified as a key factor in genome stability (28). Overall, our results support the current concept of differential replication timing among origins as the consequence of the differential accrual of limiting replication initiation factors. Deletions of regulators, such as Sir2 and Rpd3, that affect the differential accrual of factors disrupt the existing balance, potentially creating conflicts between different genomic regions. Thus, understanding how different chromatin regulators help to balance the activity of origins intrinsically in competition with each other is an important goal.

A clear conclusion of our study is that Rpd3 acts independently of Sir2 and of the rDNA in exerting its effects on singlecopy origins. By quantifying and normalizing for the absolute levels of replication associated with single-copy versus the rDNA origins, we showed that the effects of *RPD3* deletion on single-copy origins are additive with, and hence independent of, SIR2 deletion, whereas SIR2 deletion's effect on the rDNA origins is epistatic to RPD3 deletion. Specifically, RPD3 deletion in *sir2* Δ cells affects single-copy origins without altering rDNA origin firing, which remains elevated in the sir24 background. This is not consistent with Rpd3 acting as an rDNA origin activator but is consistent with Rpd3 acting as a repressor of single-copy origins. Indeed, the increased pan-S phase efficiency of rDNA origins in $rpd3\Delta$ cells is consistent with Rpd3 acting as a repressor of rDNA origins also. Moreover, we showed that the effect of RPD3 deletion on single-copy origins occurred in strains with much reduced rDNA complement, even while the effect on rDNA origins was also reduced. Thus, deregulation of single-copy origins in $rpd3\Delta$ cells does not appear to derive from primary effects on rDNA origins, though we cannot rule out the possibility of direct effect(s) of Rpd3 on rDNA origins.

Our quantitative conclusions on the effects of Rpd3 and Sir2 on rDNA origin firing from QBU were confirmed by the "gold standard" method of 2D gels for analysis of origin initiation levels. First, we showed that the differences in early firing levels (in HU) matched those determined by QBU, with significantly higher initiation of the rDNA origins in *sir2* Δ and *sir2* Δ *rpd3* Δ cells but lower initiation in *rpd3* Δ cells compared with WT.



Fig. 6. *RPD3* deletion suppresses *DBF4* deficiency. (*A*) Diploid strain YHy38 (*FKH1/fkh1*Δ *DBF4/dbf4*Δ*C*) was induced into meiosis and sporulation. Tetrads were dissected onto rich medium and incubated at 23 °C and imaged after 3 days and genotyped. Tetrads are arranged in columns; determined haploid genotypes are indicated on the panel to the right as F/f for *FKH1/fkh1*Δ and D/d for *DBF4/dbf4*Δ*C*; *FKH1* and *DBF4* are unlinked. Genotypes in red for inviable spores were inferred based on assumption of 2:2 segregation of alleles. The "fd" hypotheticals are circled. (*B*) Diploid strain OAy1188 (*RPD3/rpd3*Δ *DBF4/dbf4*Δ*C*) was induced into meiosis and sporulation. Tetrads *dbf4*Δ*C*) was induced into meiosis and sporulation. Tetrads were dissected onto rich medium and incubated at 30 °C and imaged after 3 days. Tetrads are arranged in columns; determined haploid genotypes are indicated on the panel to the right as R/r for *RPD3/rpd3*Δ *DBF4/dbf4*Δ*C*] was induced into meiosis and sporulation. Tetrads were dissected onto rich medium and incubated at 30 °C and imaged after 3 days. Tetrads are arranged in columns; determined haploid genotypes are indicated on the panel to the right as R/r for *RPD3/rpd3*Δ and D/d for *DBF4/dbf4*Δ*C*; *RPD3* and *DBF4* are unlinked. Genotypes in red for inviable spores were inferred based on assumption of 2:2 segregation of alleles. The "rd" isolates and hypotheticals are circled; the "Rd" hypotheticals are boxed.

Table 1. Sequences of DNA oligonucleotides used in this study

Oligonucleotides	Sequence
SIR2-deletion-F	ACATCTAGCACTCCTTCCAAC
SIR2-deletion-R	ACCTGCCCTTCTTACATTAAGC
RPD3-deletion-F	TCGCGGGCTGAACTGAATC
RPD3-deletion-R	GCTTTATCAACAGCGGTGGG
sir2-del-up	TTACTTGTAGCCTGCAACTCC
rpd3-del-up	TCAGCATAACGAATTGACGG
sir2-del-down	ATTCGACTTCTTTCCTTCGTTGT
rpd3-del-down	TGCAATTAGAAGAGAGTGAATC
int-sir2-down	ATCACAGGGTTCAATGTCGG
int-rpd3-down	TAGTGTTCAGTTGAATCACAC
SIR2-del-TRP-F	ATTCAAACCATTTTTCCCTCATCGGCACATTAAAGCTGGGACAAGTAACTGCAGGAATT
SIR2-del-TRP-R	ATATTAATTTGGCACTTTTAAATTATTAAATTGCCTTCTACACTATAGGGCGAATTGGGT
HIS-F	AGAATACCCTCCTTGACAGTC
HIS-R	TAGTATCGAATCGACAGCAG
HML-alpha-del-F	TATAGGGCAGTGTGTGACTTATGAATTGTTGTAGAAGGACgacatggaggcccagaatac
HML-alpha-del-R	ATGGCACAAGGAACACGCATTTTCCCAAGGCTTAGTATACcggcgttagtatcgaatcga
extend-HML-F	TTTTGGGACGATATTGTCATTATAGGGCAGTGTGTGACTT
extend-HML-R	TTTTATGAAGTAGCTTTCGGATGGCACAAGGAACACGCAT
probe-F	CTCACACTTGTACTCCATGA
probe-R	TGATGTGGAGAATAAGGTGC
2xL-3xFLAG-FKH1-5'	TTTCTACTACGACATCCATGGACGTAACAACAAACGCAAACGTGAACAATTCCTCTCTGAGTAGGGAACAAAAGCTGGAG
2xL-3xFLAG-FKH1-3'	TTCTTAACGGGTCTTTGTTCTTTATTGTTTAATAATACATATGGGTTCGACGACGCTGAATTCTATAGGGCGAATTGGGT
pAG61-Fkh1∆5′	AATAATAGTGTGTAAATTGTGCGTTCAATTAGCAAAGAAAgacatggaggcccagaatac
pAG61-Fkh1∆3′	TATTGTTTAATAATACATATGGGTTCGACGACGCTGAATTcggcgttagtatcgaatcga
Fkh2-up400bp	CATTACCGAAAATCTTCGATTTCGC
Fkh2-down375bp	CCGAAGCGTTGAGAAACAGC
Dbf4∆C-F1	AGCACAGACAGCAGCAGCGGTGAAGAAAGAAACGGTAtgacggatccccgggttaattaa
Dbf4∆C-R1	GATTTTATCACTAAAAGCTACTGCACTTTACGTCGTGTCCcggcgttagtatcgaatcga
3HA-MCM4-F	CGAGGGTGTAAGGAGATCAGTTCGCCTGAATAACCGTGTCCGGATCCCCGGGttaattaac
3HA-MCM4-R	GATTTTATCACTAAAAGCTACTGCACTTTACGTCGTGTCCcggcgttagtatcgaatcga

Oligonucleotides were obtained from International DNA Technologies.

Remarkably, we found that $rpd3\Delta$ cells showed a higher level of rDNA origin firing than WT in a pan-S phase culture and conversely found that $sir2\Delta$ and $sir2\Delta$ $rpd3\Delta$ cells were unchanged from WT in this analysis. In both cases, it appears that changes in origin usage subsequent to the levels observed in the early-Sphase-arrested cells, compensate for the earlier imbalances We used a BrdU pulse-labeling approach throughout S phase that provided evidence in support of the idea that the early reduction in rDNA origin firing in $rpd3\Delta$ cells is followed by an increase in later rDNA origin firing relative to WT cells. We note that the levels of early- versus late-origin usage are intrinsically linked in that reduced usage early reserves a larger complement of origins for later usage and vice versa. It is also anticipated that, as replication progresses, the abundance of replication factors will increase relative to the remaining unfired origins, while passive replication of potential origins by active forks simultaneously plays an important role in these dynamics (29, 30). In considering passive replication of the rDNA, it is notable that most replication is unidirectional due to the presence of the replication fork barrier (31), so passive replication of rDNA origins is reduced, potentially reserving more origins for later firing. Interestingly, the reduced early replication of rDNA in rpd3A cells ultimately is associated with a higher overall efficiency, which we think is quite unusual and are unaware of similar examples.

The exact mechanism involved in the repression of origin firing by Rpd3 has remained elusive and is generally thought to act through deacetylation to reduce access to chromatin (32). Several potential targets have been proposed among replication-initiation factors, including those acting in both cyclin-dependent kinase (CDK)-dependent and DDK-dependent initiation pathways (10, 12, 14). For example, we showed previously that RPD3 deletion partially suppresses the late-origin initiation defect of $clb5\Delta$ cells (10), presumably facilitating access of Clb6-Cdk1, which we showed was limiting in *clb5* Δ cells (33). Mantiero and colleagues showed that RPD3 acts in opposition to simultaneous overexpression of multiple initiation factors on both initiation branches (14). More recently, we showed that overexpression of Fkh1/2 alone stimulates significantly more origins to detectably fire earlier (18). As origin stimulation by Fkh1/2 involves Dbf4 recruitment directly to origins (22, 23), we tested whether RPD3 affects the DDK pathway through impedance of Fkh1 access to bind origin chromatin. ChIP analyses indicate that RPD3 deletion leads to a modest but significantly increased enrichment of Fkh1 at Rpd3repressed origins in G1 phase, anticipating the modest increases in QBU signals at these origins in early S phase. Together with the dependence of increased origin firing in $rpd3\Delta$ cells on FKH1/2, these findings support a mechanism whereby RPD3 regulates origin function by impeding Fkh1 binding, to varying degrees at different loci. The relatively subtle effects of RPD3 deletion on Fkh1 binding may reflect the limiting nature of Fkh1 being distributed across numerous potential binding loci differing in the No. and/or distribution of Fkh1/2 binding sites and/or in local recruitment of Rpd3. In fact, Fkh1/2 were originally identified as potential origin regulators due to anticorrelation of their binding with Rpd3-regulated origin loci (21). We also showed previously that Fkh1 binding is associated with decreased nucleosome occupancy (25), and Soriano et al. reported changes in nucleosome occupancy at Fkh1-binding sites in $rpd3\Delta$ cells (32).

Finally, we presented evidence that Fkh1 and Rpd3 have opposing functions in relation to Dbf4, as deletion of *FKH1* causes lethality of *dbf4* Δ *C* cells, whereas elimination of *RPD3* function rescues the viability of cells having the *dbf4* Δ *C* allele, which is hypomorphic for DDK function. The implication is that *RPD3* deletion facilitates Fkh1 recruitment to origins, which becomes critical in Dbf4-deficient cells. This finding reinforces the model we have proposed wherein Rpd3 acts by impeding Fkh1, though the results do not exclude other factors from being similarly affected, such as implied for the CDK pathway by suppression of *clb5* Δ defects as discussed above. Given the pervasive role of histone deacetylases in most aspects of genome regulation, the mechanistic insights gained here in yeast are expected to serve as paradigms for derived functions in more-complex systems.

Concluding on the possible evolutionary and developmental roles of Fkh1/2 in regulation of rDNA origin activity, we note that rDNA dynamics, such as expansion and contraction of repeats, and regulation of gene expression are both potentially influenced by rDNA origin activity and replication timing. For example, higher initiation levels are expected to lead to higher levels of blocked forks and thus increased potential for DNA breaks, leading to changes in rDNA copies. Increased early firing of the rDNA may also contribute to higher rRNA expression levels by increasing template copies early in the cell cycle and confer a growth advantage. Intriguingly, deletion of both *FKH1* and *FKH2* leads to pseudohyphal growth, which is

normally a starvation response, on rich medium, suggesting that *FKH1/2* play a central role to integrate growth signals with cell-cycle progression and DNA replication, which will be interesting to study further.

Materials and Methods

Yeast Strain Construction. Strain constructions were carried out by genetic crosses or lithium acetate transformations with linearized plasmids or PCR products generated with hybrid oligonucleotide primers having homology to target loci (34, 35); primer sequences for strain constructions are given in Table 1. Yeast strains are described in Table 2. All strains used for experiments are congenic with W303 background, and most are derived from BrdU-incorporating strains CVy43, CVy61, and CVy68, which are derived from SSy161 (36). OAy1096 was derived from a cross of OAy1069 and OAy1070 (18). OAy1100 was created by transformation of SSy161 with PCR product of primers 2xL-3xFLAG-FKH1-F and 2xL-3xFLAG-FKH1-R using plasmid p2xL-3xFLAG(TRP1) as template (gift from T. Tsukiyama); expression of Fkh1-3xFLAG was confirmed by Western blotting with anti-FLAG M2 at 1:1,000 (Sigma F1804). OAy1102 is an ADE2 derivative of OAy1100. The clean replacement of FKH2 with fkh2-dsm was constructed in two steps: FKH2 was deleted in CVy68 using primers pAG61-Fkh2 Δ 5' and pAG61-Fkh2 Δ 3' with URA3MX to create strain OAy1107, which was transformed with fkh2-dsm DNA amplified using primers Fkh2-up400bp and Fkh2-down375bp from p405-Fkh2-dsm (24), followed by selection of 5-fluoroorotic acid to yield strain OAy1109. OAy1114 is haploid segregant from a cross of strains OAy1109 and OAy1096. Strain YHy36 was constructed by PCR amplification of HIS3MX from pFA6a-HIS3MX6 (35) with primers HIS-F and HIS-R and transformation into JHy4. RPD3 and SIR2 deletions were constructed by

Name	Genotype	Source
SSy161	MATa ade2-1 ura3-1 his3-11,15 trp1-1 leu2-3,112 can1-100 bar1∆::hisG	Viggiani et al., 2006
CVy43	ura3::BrdU-inc(URA3)	
CVy61	trp1::BrdU-inc(TRP1)	"
CVy68	MATα leu2::BrdU-inc(LEU2)	и
CVy44	rpd3∆::KanMX ura3::BrdU-inc(URA3)	Knott et al., 2009
JHy3	his3-1 leu2-0 ura3-0 met15-0 sir2∆::KanMX	Mark Rose
JHy4	his3-1 leu2-0 ura3-0 met15-0 rpd3∆::KanMX	"
PP1758	fob1Δ::URA3 ura3::7xTK (URA3) rDNA20 (~20copies)	Philippe Pasero
JPy115	rpd3Δ::KanMX fob1Δ::URA3 ura3::7xTK (URA3) rDNA20 (~20copies)	This study
MPy55	FKH1-3xFLAG(TRP1) ADE2::FLOPv2x2	Ш
MPy102	MCM4-3HA(KanMX) TRP1::HOPv1	"
MPy188	fkh1Δ::URA3	"
OAy1096	fkh1Δ::KanMX fkh2Δ::HIS3MX ars305Δ::BrdU-inc(URA3)	и
OAy1100	FKH1-3xFLAG(TRP1)	11
OAy1102	FKH1-3xFLAG(TRP1) ADE2::FLOPv1	"
OAy1107	MATα fkh2Δ::URA3MX leu2::BrdU-inc(LEU2)	"
OAy1109	MATα fkh2-dsm leu2::BrdU-inc(LEU2)	"
OAy1114	fkh1Δ::KanMX fkh2-dsm ars305Δ::BrdU-inc(URA3)	"
OAy1186	MATα rpd3ΔKanMX leu2::BrdU-inc(LEU2)	"
OAy1188	OAy1186 x YHy19 diploid	"
OAy1189	dbf4 Δ C::HIS3MX leu2::BrdU-inc(LEU2)	"
YHy3	sir2Δ::KanMX hmlɑΔ::URA3MX trp1::BrdU-inc(TRP1)	"
YHy6	sir2Δ::KanMX rpd3Δ::HIS3MX hmlaΔ::URA3MX trp1::BrdU-inc(TRP1)	и
YHy13	fkh1Δ::KanMX fkh2-dsm rpd3Δ::HIS3MX ars305Δ::BrdU-inc(URA3)	и
YHy17	rpd3Δ::HIS3MX FKH1-3xFLAG(TRP1) ADE2::FLOPv1	и
YHy19	dbf4∆C::HIS3MX ura3::BrdU-inc(URA3)	и
YHy27	rpd3Δ::HIS3MX FKH1-3xFLAG(TRP1) ADE2::FLOPv2x2	и
YHy33	sir2∆::KanMX trp1::BrdU-inc(TRP1)	и
YHy36	his3-1 leu2-0 ura3-0 met15-0 rpd3∆::HIS3MX	Ш
YHy37	rpd34::HIS3MX MCM4-3HA(KanMX) TRP1::HOPv1	Ш
YHy38	OAy1189 x MPy188 diploid	11

Table 2. Genotypes of S. cerevisiae strains used in this study

All strains are in the W303 (*RAD5*) background; the parental genotype of SSy161 is shown, whereas for other strains, only differences from this genotype are given, with the exceptions of strains JHy3, JHy4, and YHy36, which are in the BY4741 background and only used as source DNA for strain constructions.

PCR amplification of rpd3A::KanMX from the CVy44 or JHy4 genomes or rpd3A::HIS3MX from the YHy36 genome using primers RPD3-del-up and RPD3del-down or sir2A::KanMX from the JHy3 genome using primers SIR2-del-up and SIR2-del-down, followed by transformations into PP1758, CVy61, MPy102, OAy1102, OAy1114, and YHy3 to create strains JPy115, YHy33, YHy37, YHy17, YHy13, and YHy6, respectively (see Table 2). HML α deletions were constructed by transformation of YHy33 with PCR-amplified hmla∆::URA3MX from pAG61 (37), using primers HMLalpha-del-F and HMLalpha-del-R to create YHy3. MCM4 was carboxyl-terminally tagged with 3xHA epitope by transformation with PCR product of primers 3HA-MCM4-F and 3HA-MCM4-R and template pFA6-3HA-kanMX6 to create strain MPy102; expression of Mcm4-3xHA was confirmed by Western blotting with anti-HA (16B12 or 12CA5) at 1:2,000. DBF4 carboxyl-terminal truncation $(dbf4\Delta C)$ was created using primers Dbf4 Δ C-F1 and Dbf4 Δ C-R1 to amplify a DNA fragment for insertion of a stop codon with HIS3MX into a diploid strain based on crossing CVy43 and CVy68. Haploid segregants were generated to yield YHy19. CVy44 was backcrossed with CVy68 to generate OAy1186, which was mated with YHy19 to create diploid OAy1188. Genomic alterations were confirmed by PCR analysis and/or DNA sequence analysis as appropriate.

Other Methods. Cultures were grown at 23 °C, and synchronization was performed as described previously (38). Five micrograms per milliliter α -factor was used for synchronization of *BAR1* strains and 5 ng/mL for *bar1* Δ strains. QBU analysis was performed as described (38) using KAPA Hyper Prep Kit (KK8504). DNA content analysis by flow cytometry (FACScan) has been described previously (10). 2D gels were performed as described previously (39), except that only the rDNA fraction in the cesium chloride gradient was isolated and benzoylated naphthoylated diethylaminoethyl cellulose enrichment was omitted. One microgram rDNA was digested with HindIII; the probe for the 2D gel was generated as described previously by radioactive labeling of a DNA fragment produced by PCR with primers Probe-F and Probe-R using genomic DNA as template. Radioactive signal was captured on phosphor screens, scanned with Typhoon scanner, and imaged and quantified with ImageQuant software (Bio-Rad). ChIP-seq was performed as described (40) using KAPA Hyper Prep Kit (KK8504) using anti-FLAG M2 monoclonal antibody (Sigma F1804) at 1:100 and anti-HA monoclonal antibodies 12CA5 (Millipore-Sigma ROAHA) and 16B12 (BioLegend MMS-101P) at 1:100. High-throughput DNA sequencing was performed on one of several Illumina platforms at several different facilities.

Computation and Statistics. All sequencing data were binned (50 bp) and median smoothed over a 1-kb window. BrdU peaks were called by MACS 1.4.2 with no-model mode (P < 0.01). The called peaks in each set were cross-referenced against origins listed in OriDB to eliminate any peaks not aligning

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with a "confirmed" or "likely" origin. Overlapping origins between the different datasets were determined by bedtools 2.25.0 using intersect function; from these results, the union of origins identified in WI and $rpd3\Delta$ cells was determined for use as the total origins list of 306 only for the following purpose: Rpd3-repressed origins were called by applying a two-sample, two-tailed t test (P < 0.001) comparing WT and $rpd3\Delta$ QBU signals within a 5-kb window centered on these 306 replication origins; 80 differential origins were detected, of which 74 were increased in $rpd3\Delta$ and thus denoted Rpd3 repressed (SI Appendix, Table S1). The list of origins and their T_{Rep} values and set assignments (FKH activated; n = 94, excludes rDNA origin) are from refs. (20, 21). CEN-proximal origin refers to the closest origin on each side of each centromere (SI Appendix, Table S2). FKH1-OE-activated origins were taken from ref. (18) (SI Appendix, Table S3). Matlab was used for generation of most data displays and analyses. ChIP-seq data were normalized against data untagged control strains. Fkh1 ChIP peaks were called by MACS 2.2.7.1 with no-model mode and instead an extension size of 100 bp. ChIP data files were compared with control data files (untagged strain), allowing for background to be estimated (false discovery rate adjusted P < 0.05).

Data, Materials, and Software Availability. [DNA sequencing] data have been deposited in [GEO](GSE200351) (41).

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