Integrated genomic analysis reveals key features of long undecoded transcript isoform-based gene repression

Highlights

- Long-read RNA-seq reveals 5’-extended mRNAs as candidate LUTIs in yeast meiosis
- Majority of the LUTIs are regulated by a cell-fate-instructive transcription factor
- LUTI-based translational repression depends on uORFs
- Chromatin changes and LUTI expression predict LUTI-based transcriptional repression

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In brief

Tresenrider et al. conducted a genome-wide study to dissect an unconventional gene regulatory mechanism whereby transcription factors could function as repressors by inducing non-canonical mRNAs called LUTIs. Their findings reveal that LUTI-based gene repression depends on a combination of factors, including translational repression, chromatin state changes, and LUTI expression.
Integrated genomic analysis reveals key features of long undecoded transcript isoform-based gene repression

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SUMMARY

Long undecoded transcript isoforms (LUTIs) represent a class of non-canonical mRNAs that downregulate gene expression through the combined act of transcriptional and translational repression. While single gene studies revealed important aspects of LUTI-based repression, how these features affect gene regulation on a global scale is unknown. Using transcript leader and direct RNA sequencing, here, we identify 74 LUTI candidates that are specifically induced in meiotic prophase. Translational repression of these candidates appears to be ubiquitous and is dependent on upstream open reading frames. However, LUTI-based transcriptional repression is variable. In only 50% of the cases, LUTI transcription causes downregulation of the protein-coding transcript isoform. Higher LUTI expression, enrichment of histone 3 lysine 36 trimethylation, and changes in nucleosome position are the strongest predictors of LUTI-based transcriptional repression. We conclude that LUTIs downregulate gene expression in a manner that integrates translational repression, chromatin state changes, and the magnitude of LUTI expression.

INTRODUCTION

Gene expression programs determine cellular identity, form, and function through extensive regulation. Key regulators of such programs include transcription factors (TFs), which initiate cascades of gene expression that ultimately drive changes in cellular state. While TFs are generally implicated in gene activation, less is known about how gene repression events can be coordinated with TF-dependent waves of gene activation.

New insights into TF-mediated gene repression came from the studies of an essential gene called NDC80. NDC80 is tightly regulated during meiotic differentiation, the developmental program that generates gametes (Asakawa et al., 2005; Chen et al., 2017; Miller et al., 2012; Sun et al., 2011). Upon meiotic entry, a key TF complex, Ime1-Ume6, activates the distal NDC80 promoter (P2) to induce a 5′-extended mRNA isoform (Figure 1A). This transcript cannot be translated into a functional protein because the upstream open reading frames (uORFs) in its 5′ leader inhibit the translation of the coding sequence (CDS). Instead of producing Ndc80 protein, the “long undecoded transcript isoform” or “LUTI” serves to inactivate the CDS-proximal NDC80 promoter (P1). As a result, the protein-coding NDC80 transcript is downregulated in meiotic prophase (Chen et al., 2017; Chia et al., 2017). Reliance of the LUTI-based repression of NDC80 on Ime1-Ume6 couples gene activation and repression events to a common TF.

The LUTI-based mechanism is neither limited to NDC80 nor restricted to yeast meiosis. Genome-wide analysis revealed that at least 380 LUTIs are expressed throughout yeast meiosis, whereby transcript levels inversely correlate with protein expression (Cheng et al., 2018). During the endoplasmic reticulum unfolded protein response, Hac1 TF induces at least 15 LUTIs, most of which are required for downregulating genes involved in cellular respiration (Van Dalfsen et al., 2018). Finally, the MDM2 oncogene is regulated by a LUTI-based mechanism in human cells (Hollerer et al., 2019). The prevalence of LUTIs in yeast and their discovery in humans, along with the pervasive use of alternative transcription start sites (TSSs) and the frequency of uORFs all point to the conservation of LUTI-based gene repression across vast evolutionary time (Batut et al., 2013; Calvo et al., 2009; Chew et al., 2016; Johnstone et al., 2016; Kimura et al., 2006).

Many questions remain to be answered about LUTI-based gene repression. First, why do only some 5′-extended transcripts appear to repress gene expression while others do not? Is the difference due to lack of translational repression,
A) LUTI not expressed

B) TL-seq

C) Meiosis-specific gene

D) URS1 Consensus Motif

E) Fold-enrichment over input

F) Canonical (n=73)

G) Degree of conservation

H) Phastcons conservation

I) RNA-seq (log normalized counts)

J) (legend on next page)
transcriptional repression, or both? Second, when gene repression does occur, does it involve a common mechanism? Lastly, what are the key features of the LUTI-based mechanism on a global scale?

In this study, we used transcript leader sequencing (TL-seq) in conjunction with RNA sequencing on nanopore arrays (direct RNA sequencing [RNA-seq]) to identify seventy-four 5′-extended transcript isoforms in meiotic prophase. A total of 80% of these transcripts are directly regulated by a meiotic TF with consensus-binding motifs conserved across the budding yeast sensu stricto genus. Importantly, uORFs are present in 97% of the 5′-extended transcripts and almost all of them appear to be bona fide LUTIs based on uORF-dependent translational repression. Transcriptionally, the outlook is more complex: LUTI transcription leads to the downregulation of the CDS-proximal transcript in 50% of cases. Chromatin modifications, specifically histone 3 lysine 36 trimethylation (H3K36me3), and changes in nucleosome position are among the strongest predictors of LUTI-based repression. Finally, higher LUTI expression leads to a greater repression of the CDS-proximal transcripts. We conclude that translational repression, chromatine state changes, and degree of LUTI expression act in a combinatorial manner to determine the extent of LUTI-based repression. Our findings provide a roadmap to uncover similar types of gene regulation in other biological contexts.

RESULTS

Transcript leader and nanopore sequencing identifies 74 potential LUTIs in meiotic prophase

To determine the prominent features of LUTI-based repression, we sought to identify meiotic mRNAs with 5′ extensions (Figure 1B). Using a previously published early meiotic cell synchronization system, we aimed to uncover all 5′-extended mRNAs that are expressed in meiotic prophase, but not in the premeiotic phase (Berchowitz et al., 2013). Identification of the 5′-extended isoforms relied on data from two complementary sequencing techniques. The first method, TL-seq, allowed for the identification of TSSs (Figures 1B and 1C; Arnbere and Gilbert, 2013; Malabat et al., 2015). The second technique, nanopore sequencing, confirmed the instances in which TSSs identified by TL-seq produced transcripts that elongate across an entire CDS (Figure S1; Garalde et al., 2018). To distinguish LUTIs from the canonical meiotic mRNAs, we restricted our calls to loci in which the TSS identified by TL-seq was upstream of a second, CDS-proximal TSS (Figures 1B and 1C). These analyses revealed 74 potential LUTIs, 28 antisense transcripts, 65 intergenic transcripts, and 74 intragenic transcripts in meiotic prophase (Tables S1 and S2).

The vast majority of putative LUTIs are regulated by the same meiotic TF

We next looked for common regulatory features near candidate LUTI promoters and found a single significant hit matching the URS1 consensus motif (Figure 1D; Table S2; combined match p < 0.05) (Sumrada and Cooper, 1987). In mitosis, this motif is bound by Ume6, which functions as a transcriptional repressor (Park et al., 1992). However, Ume6 becomes a transcriptional coactivator upon interaction with the early meiotic TF Ime1, culminating in the expression of early meiotic genes (Bowdish et al., 1995).

To investigate how many URS1 sites are bound by Ume6, we performed chromatin immunoprecipitation followed by deep sequencing (ChIP-seq). Ume6 was enriched at 61 of the 74 candidate LUTI promoters (q < 0.001, fold enrichment > 4). The extent of enrichment was similar to a set of previously identified Ume6 targets (canonical, Figures 1E–1H; Williams et al., 2002). Both groups were more enriched with Ume6 than genes not in these lists (Figures 1E, 1F, and S2A; Table S2).

We further examined URS1 motif conservation in putative LUTI promoters. Using an alignment of 5 yeast species in the sensu stricto genus, conservation of the region around the URS1 motif was calculated for all of the sites that had both a URS1 motif and were bound by Ume6. For the previously defined canonical Ume6 targets and the regions around the newly identified 5′ extensions, conservation sharply increased around the URS1 motif (Figures 1G and 1H), providing strong evidence that 33 of the identified 5′-extended transcript isoforms have strongly conserved URS1 binding sites (Table S2).

As a final test, we performed RNA-seq in cells containing a mutant allele of UME6 (T99N). The T99N mutation inhibits the interaction between Ume6 and Ime1, preventing meiosis-specific gene activation without compromising Ume6 function during mitosis (Bowdish et al., 1995). In these cells, only 2.7% of

Figure 1. LUTI discovery and analysis of TF-based regulation of LUTIs

(A) A schematic of LUTI-based gene regulation. The terms ORF and CDS are used interchangeably throughout the text and figures.

(B) An overview of the discovery pipeline.

(C) Genome browser views of TL-seq for the meiosis-specific gene SPO11 and the NDC80 locus, which contains a LUTI.

(D) The URS1 motif found in LUTI promoters and the consensus-binding motif observed 300 bp ± distal TSSs.

(E and F) Ume6 ChIP-seq from (UB3301) grown in BYTA to saturation. One of 3 replicates is shown.

(G and H) For genes with both Ume6 enrichment and a URS1 motif, the degree of conservation across the 100 bp ± the URS1 motif center was determined by phastCons within the sensu stricto genus.

(I) Metagene analysis for degree of conservation in the promoters of previously identified Ume6 targets compared to the promoters of the 5′-extended transcripts.

(J) Heatmap of Ume6 fold enrichment over input in the promoters of previously identified Ume6 targets and in the promoters of the 5′-extended transcripts.

(k) RNA-seq from strains expressing either WT Ume6 (UB20649) or Ume6 with a T99N mutation (UB22629). Cells harboring pCUP1-IME1/pCUP1-IME4 were collected after 2 h (premeiotic) or 4 h (meiotic prophase) in sporulation medium (SPO), which corresponds to 2 h after induction of meiosis by 50 μM CuSO4. This induction method and timing was used for all subsequent experiments.
the loci with 5′-extended transcripts had increased gene expression after the induction of IME1 (Figures 1I, S2B, and S2C; Table S3). The percentage was reduced to 2.2% for the 46 loci in which a URS1 motif is present and Ume6 binding is detected by ChIP-seq. In combination, this evidence suggests that Ime1-Ume6 directly regulates the vast majority of 5′-extended transcripts identified in this study (Table S3).

Transcript isoform diversity and the translational regulation of 5′-extended isoforms
LUTIs were originally identified based on an inverse correlation between transcript and protein levels (Chen et al., 2017; Cheng et al., 2018). Of the 380 meiotic LUTIs in Cheng et al. (2018), 32 (8.4%) of them were found to be expressed during the meiotic prophase time point collected in this study. The remaining 42 of our 74 candidate LUTIs were previously unidentified. While 10 of the false negatives were missed due to the lack of a quantifiable protein measurement, the other 32 were likely overlooked due to differences in the method of identification (Cheng et al., 2018).

Our pipeline detects all 5′-extended transcripts, regardless of their correlation with protein abundance. Such newly identified LUTI candidates are potentially very interesting because they may reveal biological insights into how frequently candidate LUTIs repress gene expression and under what circumstances this repression occurs.

It is difficult to answer these questions using traditional RNA-seq, because the transcript produced from the CDS-proximal promoter (PROX isoform) does not have any unique identifying sequence compared to the transcript produced from the CDS-distal promoter. By sequencing the most 5′ end of a transcript with TL-seq, this was no longer a limitation. We performed TL-seq and RNA-seq in parallel on samples collected during the premeiotic phase and meiotic prophase. RNA-seq and TL-seq measurements (in which the closest TSS to CDS was quantified) correlated well for most genes, demonstrating that TL-seq can quantitatively estimate transcript abundances (Spearman’s rank correlation coefficient, \( r = 0.771 \) [premeiotic], \( r = 0.732 \) [meiotic prophase]; Figures 2A and 2B). However, for genes with 5′-extended transcripts, this correlation was poor in meiotic prophase for both the PROX (\( r = 0.419 \)) and extended (\( r = 0.347 \)) transcripts; it improved when both transcripts were considered (\( r = 0.602 \)) (Figures 2B and S3A). When the fold change (FC) of expression between meiotic prophase and premeiotic stages was taken into account, there was no
correlation between TL-seq and RNA-seq for genes with 5'-extended transcripts \((p = 0.04; \text{Figure 2C})\). This suggests that TL-seq is a more robust method than RNA-seq to specifically quantify the 5'-extended and PROX isoforms.

With the ability to distinguish between the transcript isoforms, we set out to compare the correlation between CDS translation and transcript isoform abundance using a published dataset \(\text{(Cheng et al., 2018)}\). The meiotic prophase RNA-seq dataset from our study \(\text{(4-h)}\) best matched the 3-h time point from \(\text{Cheng et al., 2018}\), which used asynchronous meiotic cultures \((p = 0.833; \text{Figure S3B})\). This 3-h time point represents the peak period when recombination and synaptonemal complex formation occurs \(\text{(Cheng et al., 2018; Brar et al., 2012)}\). Gene set enrichment analysis \(\text{(GSEA)}\) from our RNA-seq dataset also revealed that recombination-related gene sets were highly enriched in the corresponding 4-h time point \(\text{(Table S4; Mootha et al., 2003; Subramanian et al., 2005)}\). The most enriched gene set was that of genes previously reported as highly induced during the process of recombination \(\text{(G, I, and K)}\). RNA blots performed with a probe specific for 3V5 and its linker. rRNA bands detected by methylene blue.

We next investigated how the LUTIs were translationally impaired. While 5’ leader-mediated translational repression can occur through various means \(\text{(Hinnebusch et al., 2016)}\), the abundance of uORFs in meiosis \(\text{(Brar et al., 2012)}\), as well as the uORF-dependent translational repression of \(\text{NDC80-LUTI}\) \(\text{(Chen et al., 2017)}\), led us to hypothesize that uORFs would play a dominant role in dampening CDS translation from LUTIs.

**Prevalence of uORFs in meiotic LUTIs**

We next investigated how the LUTIs were translationally impaired. While only a few extended transcripts showed a negative correlation \((p = -0.297; \text{Figure 2F})\). This analysis demonstrates that transcript isoform-specific measurements are more accurate in assessing translational status than RNA-seq and provides additional evidence that the 5'-extended transcripts do not productively translate the CDSs contained within them.

The number of ATG codons in the region between the distal TSS and the PROX TSS was used to determine the number of uORFs in candidate LUTIs. This analysis revealed that 72 of 74 candidate LUTIs had at least \(2\) uORF, and \(64\) had between \(4\) and \(17\) \(\text{(Figure 3A)}\). The two without a uORF \(\text{(PLM2 and YCL057C-A)}\) displayed an increase in TE \(\text{(translational efficiency)}\) ribosome footprints reads per kilobase of transcript, per million mapped reads \(\text{(RPKM)}\), indicating that they should not be categorized as LUTIs \(\text{(Table S5)}\).

When an extended transcript contained a single uORF, the effect on translational efficiency was variable. The TE changed very little for \(2\) genes \(\text{(ELO1 and COX16)}\), possibly because the LUTI was the minor isoform. The LUTI was the major isoform in the other \(2\) cases, and the TE decreased dramatically \(\text{(Table S5)}\), suggesting that a single uORF was sufficient to inhibit translation initiation at downstream AUGs. Based on these analyses, all of the 5'-extended transcripts containing at least \(1\) single ATG uORF will be referred to as LUTIs going forward.

We directly tested the ability of a single uORF to repress the translation of the downstream CDS at the \(\text{NDC80} \_\text{LUTI}\). Normalized normalizes \(9\) uORFs. When all \(9\) ATGs were mutated to ATCs \(\text{(J9AUG)}\), Ndc80 protein was translated from the LUTI \(\text{(Figures 3B and 3C; Chen et al., 2017)}\). Strains were constructed in which the ATG of either the first, fifth, or ninth uORF was the sole AGT-initiated uORF in the 5’ leader. Having the first uORF alone resulted in similar Ndc80 protein levels to what was observed in the J9AUG strain; however, LUTIs carrying only the fifth or the ninth uORF repressed Ndc80 translation just as efficiently as the wild-type \text{NDC80-LUTI} \(\text{(Figures 3B and 3C)}\).

The frequency of uORF translation in LUTIs going forward. We next set out to establish another metric to determine whether LUTIs are translationally repressed. As the uORF number in a 5’ leader increases, the likelihood of repression at the downstream CDS also increases \(\text{(Calvo et al., 2009; Chew et al., 2016; Johnstone et al., 2016)}\). We predicted that if translational repression occurs, then there would be more translation over the most 5’ uORFs and less translation over the uORFs closest to the CDS. Because uORFs are short and frequently overlapping, it can be difficult to accurately quantify their translation. Instead, we determined a threshold of at least 4 footprint counts within the first 6 codons of a uORF to call it as translated. With this metric, almost 80% of the first \(2\) uORFs in transcripts with at least \(4\) uORFs were classified as translated \(\text{(Figure 3D)}\), while 15% of the last \(2\) uORFs in those same transcripts were translated \(\text{(Figure 3D)}\). Thus, the ribosomes
frequently are caught up before scanning across all uORFs. This is consistent with the observation that ATG frequency was not higher in the 5′ extensions compared to the 500 bp upstream of genes not expressing LUTIs in meiotic prophase (Figure 5E). If LUTIs do play important and functional roles in mediating meiotic gene expression, then the lack of uORF selection would indicate that the natural frequency of ATGs in intergenic regions is sufficient to result in the necessary degree of translational inhibition. We conclude that uORFs are found in abundance in the 5′ leaders of most LUTIs.

Using three LUTIs (APL4, HSP60, and MSC6), we further tested the requirement of uORFs in translational repression. The ATG of all uORFs in the 5′ leaders of these LUTIs were mutated to ATC (∆uORF). Cells from wild-type and ∆uORF strains were synchronized in meiosis. Compared to cells with wild-type uORFs, the protein abundance was higher in the ∆uORF strains at the 4-h time point (Figures 3F, 3H, 3J, and S4A). This occurred even in the absence of any observed increase in the PROX transcript, suggesting that increased translation was occurring from the mutated LUTI transcript (Figures 3G, 3I, and 3K). Intriguingly, an increase in LUTI levels was evident in the ∆uORF mutants (Figures 3G, 3I, and 3K). However, this alone is not sufficient to alter protein abundance as observed in upf1Δ cells, which have higher LUTI levels due to deficiencies in nonsense-mediated mRNA decay, a degradation pathway for transcripts with premature translation-termination codons (Figures S4B–S4M). Our findings indicate that the presence of uORFs is a major driving factor behind the poor translational efficiency of LUTIs.

**Variable transcriptional repression by LUTIs**

To assess the degree of transcriptional repression, the abundances of LUTI and PROX transcripts were measured by T-Lseq. LUTI levels did not correlate with PROX transcript abundance before meiotic entry (∝ = −0.19; Figure 4A); however, a significant negative relationship developed in meiotic prophase (∝ = −0.359, p = 1.98 × 10<sup>−5</sup>), associating LUTI expression with a decrease in PROX isoform level (Figure 4B). In addition, the PROX transcript abundance in meiotic prophase was less than in the premeiotic stage for a large number of genes (Figures 4C and 4D).

To parse out how causative the relationship was between LUTI and PROX expression, five genes were selected for in-depth analysis. SWI4 and APL4 had very strongly repressed PROX transcripts in meiotic prophase, MSC6 and HSP60 had PROX transcripts present at similar levels in both time points, and CDC60 had an intermediate amount of PROX repression (Figure 4C). Transgenes carrying an epitope tag with either a normal LUTI promoter (wild type) or a deletion (∆LUTI) were constructed. No deletions affected PROX expression compared to wild type in the premeiotic phase, but in all instances, the LUTI deletion led to an increase in meiotic PROX transcript (Figures 4E, 4G, 4I, S5A, and S5B). Increases in protein levels for SWI4 and Cdc60 (Figure 4F and 4H), but not for Api4, Msc6 and Hsp60, were observed (Figures 4J, S5A, and S5B). Thus, even when LUTI-based repression is relieved and PROX transcript isoform abundance increases, additional layers of control can buffer the final protein output.

**The role of chromatin in LUTI-based transcriptional repression**

With the knowledge that LUTI expression can play a role in repressing transcription from downstream TSSs at some loci, but not others, we set out to understand what differentiates the two cases. In both yeast and human cells, LUTI expression leads to an increase in H3K36me3 over the proximal gene promoter (Chia et al., 2017; Hollerer et al., 2019). In yeast, NDC80<sup>LUTI</sup> expression also results in H3K4me2 enrichment (Chia et al., 2017). Both marks are necessary for the LUTI-based repression of NDC80 (Chia et al., 2017).

To determine the change in H3K36me3 and H3K4me2 patterns at LUTI-associated loci, we performed ChiP-seq. In premeiotic cells, the LUTI-containing genes displayed dis-enrichment of both modifications just upstream of the PROX TSS (Figures 5A and S5B, top panel, and Figures S6A–S6D). Interestingly, H3K36me3 was enriched in the meiotic prophase specifically over the promoters of those genes expressing LUTIs (Figure 5A, bottom panel). Strikingly, those genes whose PROX transcript is most repressed (FC < 0.25, n = 21) have the highest levels of H3K36me3 over their proximal TSS (Figure 5A; Table S6). In contrast, genes with LUTIs that do not experience a decrease in the PROX transcript (log2 FC > 0, n = 18) have only a minor increase in H3K36me3 levels (Figure 5A; Table S6). For H3K4me2, a moderate increase in the chromatin modification is observed over the PROX promoters of only those genes that are the most repressed in meiotic prophase (Figure 5B; Table S6). Thus, while increased H3K36me3 is a strong predictor for LUTI-based repression, H3K4me2 appears to be altered in a limited number of instances.

**Figure 4. Transcriptional repression by LUTIs**

(A) Scatterplot of LUTI abundance in cells from meiotic prophase versus PROX abundance in premeiotic cells (UB145584) (n = 3). Spearman’s correlation coefficient (∝) is shown. (B) The same as (A), except that the PROX transcript abundance is from the meiotic prophase time point. (C) Scatterplot of PROX abundance in meiotic prophase compared to premeiotic cells, the LUTI-containing genes displayed dis-enrichment of both modifications just upstream of the PROX TSS (Figures 5A and S5B, top panel, and Figures S6A–S6D). Interestingly, H3K36me3 was enriched in the meiotic prophase specifically over the promoters of those genes expressing LUTIs (Figure 5A, bottom panel). Strikingly, those genes whose PROX transcript is most repressed (FC < 0.25, n = 21) have the highest levels of H3K36me3 over their proximal TSS (Figure 5A; Table S6). In contrast, genes with LUTIs that do not experience a decrease in the PROX transcript (log2 FC > 0, n = 18) have only a minor increase in H3K36me3 levels (Figure 5A; Table S6). For H3K4me2, a moderate increase in the chromatin modification is observed over the PROX promoters of only those genes that are the most repressed in meiotic prophase (Figure 5B; Table S6). Thus, while increased H3K36me3 is a strong predictor for LUTI-based repression, H3K4me2 appears to be altered in a limited number of instances.
We next tracked genome-wide changes to nucleosome positions using micrococcal nuclease digestion with deep sequencing (MNase-seq) since the presence of nucleosomes can occlude the binding of TFs and machinery required for transcription initiation (Klemm et al., 2019; Venkatesh and Workman, 2015). In meiotic prophase, but not in the premeiotic stage, the nucleosome peaks near PROX promoters decreased while the signal in the valleys increased (Figure 5C; Chen et al., 2013). The effect was strongest for loci with the greatest degree of PROX transcript repression (Figure 5D). In that subset of genes, the nucleosome position was so disrupted that consensus nucleosome periodicity could not be identified (Figure 5D). Examining such a small subset of genes may make it hard to observe strong periodicity; however, in a similarly small subset of genes (n = 18), loci with non-repressive LUTIs, periodic nucleosome positioning was still present (Figure 5D). The complete lack of periodicity at loci with repressive LUTIs was most likely due to variability in the extent of repositioning at each locus (Figures S6E and S6F). We conclude that robust nucleosome repositioning occurs over the promoters of genes that are most subject to LUTI-based transcriptional repression.

The features defining LUTI-based transcriptional repression

We expected that other variables could play a role in LUTI-based transcriptional repression. Stemming from previous work showing that the distance between promoters with overlapping transcription affects the mechanism of transcriptional repression in cells undergoing carbon source shifts (Kim et al., 2016), we considered the importance of the distance between the PROX and the LUTI TSSs. The LUTI abundance and the length of the gene were also considered. To better assess how changes to the nucleosome over the PROX promoter may be associated with LUTI-based repression, the DANPOS2 toolkit was used to define changes to the +1 and –1 nucleosome positions and to assess “fuzziness,” defined as the frequency with which the nucleosome differs from the most observed position (Chen et al., 2013). We found that an increase in H3K36me3 (r = −0.492, p = 1.10 × 10⁻⁹), high LUTI levels (r = −0.456, p = 5.60 × 10⁻⁴), +1 nucleosome peak moving toward the nucleosome-depleted region (r = −0.423, p = 2.13 × 10⁻⁴), and an increase in +1 nucleosome fuzziness (r = −0.341, p = 3.41 × 10⁻³) displayed significant negative correlation with the log2 FC of PROX transcript abundance (Figures 6A and S7). However, changes at the –1 nucleosome, H3K4me2, and the distance between the TSSs had no significant correlation (Figures 6A and S7). These analyses helped distinguish a set of important factors involved in LUTI-based transcriptional repression.

Promoter strengths dictate LUTI-based transcriptional repression

Surprisingly, we also found that the longer the coding sequence of a gene, the more likely it was to have an abundant
LUTI, an increase in H3K36me3, greater +1 nucleosome fuzziness, and an increased likelihood of being repressed by a LUTI (Figure 6A). Coincidentally, in the set of genes with meiotic LUTIs, shorter genes had higher PROX transcript abundances than did longer genes (Figures 6B and 6C). Could it be that the promoters of strongly expressed genes are better able to continue transcribing their gene products even in the presence of LUTI mRNAs? We decided to investigate this possibility further.

If a more robustly transcribed PROX isoform is resistant to repression by LUTIs, then it follows that increasing LUTI expression or decreasing PROX expression could shift the balance in favor of LUTI-based repression. To test this hypothesis, we focused on HSP60, a gene whose PROX isoform is highly expressed both in the premeiotic stage and in meiotic prophase. We engineered a reporter construct in which the HSP60 regulatory region was upstream of a short-lived green fluorescent protein (ubiGFP) (Houser et al., 2012). To enable titratable expression of LUTI, a β-estradiol-inducible LexA/LexO system was used (Ottoz et al., 2014; Figure 7A). As the dose of β-estradiol increased, so did LUTI abundance and H3K36me3 over the HSP60 PROX promoter (Figures 7B and 7C). Importantly, as the expression of LUTI increased, the abundance of ubiGFP simultaneously decreased (Figures 7D and 7E). The dose-dependent decrease in ubiGFP levels was also confirmed during mitotic growth in rich media (Figures 7H and 7I). We conclude that changes to the strength of LUTI expression can lead to a decrease in protein abundance even at a strong PROX promoter.

Using mitotic cells, we further asked what would happen if both the LUTI and the PROX levels were altered. Using the same system as above, we generated a construct with a mutated Hsf1 binding site in the HSP60 PROX promoter (Figure 7F). Hsf1 is a TF that controls the expression of genes involved in heat shock response, including HSP60 (Sakurai and Ota, 2011). Mutating the Hsf1-binding site reduced the basal expression of ubiGFP, but did not change LUTI transcript levels (Figures 7G–7I). Upon treatment with β-estradiol, cells carrying the hsf1 mutant were highly susceptible to LUTI-based repression, decreasing to ~25% of its original level after 3 h of LUTI induction (Figures 7H and 7I). In contrast, the ubiGFP abundance reduced to only ~60% of its original level in cells carrying the wild-type HSP60 PROX promoter (Figures 7H and 7I). Thus, changes in the strength of the PROX promoter also affect LUTI-based repression.
DISCUSSION

Developmental regulation of LUTIs

We identified 72 LUTIs in meiotic prophase. Ume6 TF was enriched in the promoter of 61 of them, with its binding motif conserved in 33 LUTIs across the sensu stricto genus (Figures 1D–1H). We posit that the conservation of the regulatory binding site in such a large subset of LUTI promoters indicates that they may alter gene expression in a manner that is functionally important in meiosis. By using LUTIs, a single TF can both turn on and turn off gene expression in a coordinated and timely manner to facilitate the necessary shift in proteome synthesis. We predict that LUTIs will play important roles during other developmental processes.

uORF-mediated translational inhibition of LUTIs

uORFs are ubiquitously found in walks of life from yeast to humans. Features including initiation sequence context, distance between a uORF and CDS, and uORF number can affect the degree of translational repression (Calvo et al., 2009; Chew et al., 2016; Johnstone et al., 2016). What is clear is that not all uORFs are created equal. We confirm that a single uORF can lead to repression at the NDC80 locus, but that it depends on which uORF. The uORF closest to the LUTI TSS (uORF1) does not repress translation of the NDC80 CDS, despite evidence from ribosome profiling that UORF1 is well translated. uORFs 5 and 9 both lead to robust repression, even though no translation of uORF 9 is observed in meiotic cells by ribosome profiling (Figure 3C; Brar et al., 2012).
is consistent with the observation that greater distances between a uORF and CDS correlate with reduced translational repression (Chew et al., 2016; Johnstone et al., 2016). In the context of the most well-characterized case of uORF-mediated repression, GCN4, the distances between the 4 uORFs and the CDS start site matter greatly. That is because, upon amino acid starvation, the concentration of the ternary complex, a factor required for ribosome re-initiation, is decreased. Ultimately, the repressive uORF4 is skipped, due to extended ribosome scanning, and GCN4 is translated (Hinnebusch, 1997). It could be worthwhile to further study the role of the ternary complex or other factors in ensuring which uORFs are integral to the translational repression of LUTIs.

The translation of uORFs and a corresponding decrease in TE is a good indication that uORFs are playing a functional role, but it does not indicate the extent of the translational regulation. TE measurements at loci with LUTIs are complicated by the presence of PROX transcripts. As an alternative to measuring TE, we measured the frequency of uORF translation. When ≥4 uORFs are present in a LUTI, translation occurs only 9.4% and 20.3% of the time for the penultimate and the last uORF, respectively (Figure 3D). The high correlation between PROX transcript abundance and footprints over the ORF (Figure 2E) is also a strong indication that LUTIs provide minor, if any, contribution to CDS translation. Future use of TL-seq and ribosome profiling will allow for the identification of other instances in which translational regulation is due to transcript isoform toggling rather than genuine translational regulation of a single mRNA isoform. Combined with a lack of uORF selection in the 5’ untranslated region (Hinnebusch, 1997). It could be worthwhile to further study the role of the ternary complex or other factors in ensuring which uORFs are integral to the translational repression of LUTIs.

A path toward predicting LUTI-based transcriptional repression

Previous work demonstrated that only ~50% of the meiotically expressed 5’-extended transcript isoforms have poorly translated CDSs (Brar et al., 2012). If almost all 5’-extended transcripts do not translate their CDSs, then why does LUTI expression not lead to translational repression more frequently? As it turns out, robust transcriptional repression of the canonical PROX transcript by its LUTI occurs far less frequently than does translational repression of the LUTI itself. In fact, only 21 of the 72 LUTIs identified here have a corresponding decrease in the canonical PROX transcript to ≤25% of the starting abundance. At sites where the canonical transcript appears unchanged following LUTI expression, LUTIs may still serve a purpose. For instance, LUTI expression could prevent an upsurge in canonical PROX transcript level, and changes to the chromatin landscape over the PROX promoter. Combined with a lack of uORF selection in the 5’ untranslated region (Hinnebusch, 1997). It could be worthwhile to further study the role of the ternary complex or other factors in ensuring which uORFs are integral to the translational repression of LUTIs.

Conclusions

In summary, we identified a group of LUTIs expressed in a coordinated manner during meiotic prophase in budding yeast. Almost all are severely translationally inhibited due to uORFs in their 5’ leaders. However, they do not ubiquitously lead to the repression of PROX promoters. Rather, an interplay between the strength of the promoters, the chromatin landscape, and possibly other yet to be discovered features interact to decide the transcriptional output. Future studies that integrate multiple datasets will help identify additional LUTIs and build predictive models to determine those that cause gene repression. In addition, it remains to be uncovered, in many cases, what the biological role of a given LUTI is and how its disruption affects cellular function. Lastly, in human cells, the presence of a LUTI at the MDM2 locus (Hollerer et al., 2019) opens the door for more studies into how significant LUTIs are in human development and disease.

Limitations of study

Here, we uncover examples whereby the upstream transcription of a LUTI, unexpectedly, does not alter the gene expression of a downstream promoter. Our analyses identified correlative relationships between PROX repression, LUTI expression, and changes to the chromatin landscape over the PROX promoter. A firm understanding of the causative interactions between such features is not addressed and is a key limitation. In addition, while we investigate the relationship between the LUTI and PROX promoters at the HSP60 locus in some depth, a more detailed exploration of LUTI and PROX promoter strength modulation under additional contexts has the potential to improve our understanding of the complex interplay that occurs when a single gene is controlled by the expression of transcripts from two distinct promoters. Lastly, we demonstrate that during early meiosis, LUTI expression occurs in a developmentally regulated fashion, but we have not yet established the biological need for such timely LUTI expression.
A.T., conceptualization, data curation, formal analysis, investigation, methodology, validation, visualization, and drafting of the manuscript. K.M., data curation, formal analysis, investigation, validation, manuscript editing, and visualization. V.J., data curation, formal analysis, investigation, methodology, validation, and visualization. M.C., TL-seq library preparations and manuscript editing. H.L., investigation of the NDC80 uORF analysis. F.J.v.W., supervision, project administration, and manuscript editing. E.U., conceptualization, formal analysis, methodology, visualization, supervision, project administration, funding acquisition, and drafting of the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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SUPPLEMENTAL INFORMATION

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### STAR★METHODS

#### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| H3K36me3            | abcam  | ab9050, RRID:AB_306966 |
| H3K4me2             | abcam  | ab32356, RRID:AB_7329224 |
| mouse α-V5          | Thermo Fisher | R960-25, RRID:AB_2556564 |
| α-GFP               | Takara | 632381, RRID:AB_2313808 |
| rabbit α-hexokinase (α-Hxk2) | US Biological | H2035-01, RRID:AB_2629457 |
| α-mouse antibody conjugated to IRDye 800CW | LI-COR Biosciences | 926-32212, RRID:AB_621847 |
| α-rabbit antibody conjugated to IRDye 680RD | LI-COR Biosciences | 926-68071, RRID:AB_10956166 |
| Chemicals, peptides, and recombinant proteins |        |            |
| Zinc-based alkaline fragmentation reagent | Ambion | AM8740 |
| recombinant shrimp alkaline phosphatase | NEW ENGLAND BIOLABS | M0371 |
| RNasin Plus         | Promega | N2611 |
| Cap-Clip acid pyrophosphatase | Tebu-Bio | C-CC15011H |
| T4 RNA ligase 1      | NEW ENGLAND BIOLABS | M0437M |
| SuperScript IV reverse transcriptase | Invitrogen | 18090010 |
| RNase H             | NEW ENGLAND BIOLABS | M0297 |
| RNase cocktail enzyme mix | Ambion | AM2286 |
| HighPrep PCR beads  | MagBio | AC-60050 |
| KAPA Hi-Fi hot start ready mix | Roche | KK2601 |
| KAPA single indexed adapters Set B | Roche | KK2802 |
| MyOne Streptavidin C1 Dynabeads | ThermoFisher Scientific | 65001 |
| oligo-dT DynaBeads   | ThermoFisher Scientific | 61002 |
| R9.4.1 flow cell     | Oxford Nanopore Technologies | FLO-MIN106 |
| Protein A Dynabeads  | Invitrogen | 10001D |
| RNase A              | Invitrogen | 12091021 |
| Proteinase K         | Roche | 3115879001 |
| AMPure XP beads      | Beckman Coulter | A63881 |
| ULTRAhyb Ultrasensitive Hybridization Buffer | ThermoFisher Scientific | AM8669 |
| Exonuclease III      | New England Biolabs | M0206S |
| Mnase                | Worthington | LS004797 |
| Superscript III kit  | ThermoFisher Scientific | 18080044 |
| Absolute Blue qPCR Mix | ThermoFisher Scientific | AB4162B |
| Critical commercial assays |        |            |
| Poly(A)Purist MAG kit | Ambion | AM1922 |
| KAPA Hyper Prep Kit | Roche | KK8504 |
| Direct RNA Sequencing Kit | Oxford Nanopore Technologies | SOK-RNA001 |
| ThruPLEX DNA-seq Kit | Takara | R400427 |
| NEXTflex™ Rapid Directional mRNA-Seq Kit | Bioo Scientific | NOVA-5138 |
| Prime-It II Random Primer Labeling Kit | Agilent Technologies, Inc | 300385 |

(Continued on next page)
| REAGENT or RESOURCE SOURCE IDENTIFIER | SOURCE |
|---------------------------------------|--------|
| Deposited data                        |        |
| TL-seq, RNA-seq, Direct RNA Nanopore Sequencing, ChIP-seq, MNase-seq | This study GEO: GSE140177 |
| RNA-seq, Ribosome Profiling           | Cheng et al. (2018) GEO: GSE108778 |
| Experimental models: organisms/strains |        |
| S. cerevisiae: UB3301: MATa/MATalpha UME6-3V5::His3MX/UME6-3V5::His3MX ime4::cup1::NAT/ime4::cup1::NAT | Brar-Ünal lab N/A |
| S. cerevisiae: UB6183: MATa/MATalpha ime4::cup1::NAT/ime4::cup1::NAT leu2::jAUG-NDC80-3V5::LEU2/ leu2::jAUG-NDC80-3V5::LEU2 ndc80::(−1000 and ORF):KanMX4 /ndc80::(−1000 and ORF):KanMX4 | Brar-Ünal lab N/A |
| S. cerevisiae: UB6190: MATa/MATalpha ime4::cup1::NAT/ime4::cup1::NAT ndc80::(−1000 and ORF):KanMX4 /ndc80::(−1000 and ORF):KanMX4 leu2::NDC80-3V5::LEU2/leu2::NDC80-3V5::LEU2 | Brar-Ünal lab N/A |
| S. cerevisiae: UB10579: MATa/MATalpha ime4::cup1::NAT/ime4::cup1::NAT leu2::pUB88-Δ8AUG-uORF1-revert-5'UTR-NDC80-3V5::LEU2/leu2::pUB88-Δ8AUG-uORF1-revert-5'UTR-NDC80-3V5::LEU2 ndc80::(−1000 and ORF):KanMX4/ndc80::(−1000 and ORF):KanMX4 | This study N/A |
| S. cerevisiae: UB10581: MATa/MATalpha ime4::cup1::NAT/ime4::cup1::NAT leu2::pUB88-Δ8AUG-uORF5-revert-5'UTR-NDC80-3V5::LEU2/leu2::pUB88-Δ8AUG-uORF5-revert-5'UTR-NDC80-3V5::LEU2 ndc80::(−1000 and ORF):KanMX4/ndc80::(−1000 and ORF):KanMX4 | This study N/A |
| S. cerevisiae: UB10583: MATa/MATalpha ime4::cup1::NAT/ime4::cup1::NAT leu2::pUB88-Δ8AUG-uORF9-revert-5'UTR-NDC80-3V5::LEU2/leu2::pUB88-Δ8AUG-uORF9-revert-5'UTR-NDC80-3V5::LEU2 ndc80::(−1000 and ORF):KanMX4/ndc80::(−1000 and ORF):KanMX4 | This study N/A |
| S. cerevisiae: UB14584: MATa/MATalpha pCUP-IME1::NAT/pCUP-IME1::NAT pCUP-IME4::NAT/pCUP-IME4::NAT amn1[BY4741 allele]unmarked/ amn1[BY4741 allele]unmarked | This study N/A |

(Continued on next page)
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| S. cerevisiae: UB18175: MATa/MATalpha int1::Hphmx/ir1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::SWI4-3V5-NDC80::LEU2/ leu2::SWI4-3V5-NDC80::LEU2 | This study | N/A |
| S. cerevisiae: UB18176: MATa/MATalpha int1::Hphmx/ir1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::pSWH4LUTI(1200 to –934)J–SWH4-3V5-NDC80::LEU2/ leu2::pSWH4LUTI(1200 to –934)J–SWH4-3V5-NDC80::LEU2 | This study | N/A |
| S. cerevisiae: UB18181: MATa/MATalpha int1::Hphmx/ir1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::pAPL4LUTI(415 to –800)J–APL4-3V5-NDC80::LEU2/ leu2::pAPL4LUTI(415 to –800)J–APL4-3V5-NDC80::LEU2 | This study | N/A |
| S. cerevisiae: UB18185: MATa/MATalpha int1::Hphmx/ir1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::CDC60-3V5-NDC80::LEU2/ leu2::CDC60-3V5-NDC80::LEU2 | This study | N/A |
| S. cerevisiae: UB18186: MATa/MATalpha int1::Hphmx/ir1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::pHSP60LUTI(1000 to –501)J–HSP60-3V5-NDC80::LEU2/ leu2::pHSP60LUTI(1000 to –501)J–HSP60-3V5-NDC80::LEU2 | This study | N/A |
| S. cerevisiae: UB18190: MATa/MATalpha int1::Hphmx/ir1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::pMSC6LUTI(800 to –335)J–MSC6-3V5-NDC80::LEU2/ leu2::pMSC6LUTI(800 to –335)J–MSC6-3V5-NDC80::LEU2 | This study | N/A |
| S. cerevisiae: UB18238: MATa/MATalpha int1::Hphmx/ir1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::MSC6-3V5-NDC80::LEU2/ leu2::MSC6-3V5-NDC80::LEU2 | This study | N/A |
| S. cerevisiae: UB18336: MATa/MATalpha int1::Hphmx/ir1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::HSP60-3V5-NDC80::LEU2/ leu2::HSP60-3V5-NDC80::LEU2 | This study | N/A |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| S. cerevisiae: UB18539: MATa/MATalpha | This study | N/A |
| irt1::cup1::Hphmx/irt1::cup1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::APL4-3V5-NDC80term::LEU2/leu2::APL4-3V5-NDC80term::LEU2 | | |
| S. cerevisiae: UB18838: MATa | This study | N/A |
| trp1::pGPD1-LexA-ER-HA-B112::TRP1 his3::8xLexO-HSP60leader-ubiGFP::HIS3 | | |
| S. cerevisiae: UB19257: MATa/MATalpha | This study | N/A |
| trp1::pGPD1-LexA-ER-HA-B112::TRP1/trp1::pGPD1-LexA-ER-HA-B112::TRP1 irt1::cup1::Hphmx/irt1::cup1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT his3::8xLexO-HSP60leader-ubiGFP::HIS3/ his3::8xLexO-HSP60leader-ubiGFP::HIS3 | | |
| S. cerevisiae: UB20485: MATa | This study | N/A |
| trp1::pGPD1-LexA-ER-HA-B112::TRP1 his3::8xLexO-hsf1-mut1-HSP60leader-ubiGFP::HIS3 | | |
| S. cerevisiae: UB20649: MATa/MATalpha | This study | N/A |
| trp1::CUP1-3HA-IME1::KANMX/ trp1::CUP1-3HA-IME1::KANMX ime4::pCUP1-3HA-IME4::KANMX/ ime4::pCUP1-3HA-IME4::KANMX Ume6-3V5::His3MX/Ume6-3V5::His3MX | | |
| S. cerevisiae: UB22629: MATa/MATalpha | This study | N/A |
| trp1::CUP1-3HA-IME1::KANMX/ trp1::CUP1-3HA-IME1::KANMX pCUP-IME4::NAT/ pCUP-IME4::NAT ume6::UME6([T99N]-3V5::KanMX/ ume6::UME6([T99N]-3V5::KanMX | | |
| S. cerevisiae: UB26122: MATa/MATalpha | This study | N/A |
| trp1::cup1::Hphmx/irt1::cup1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::APL4uORFdel-3V5-NDC80term::LEU2/leu2::APL4uORFdel-3V5-NDC80term::LEU2 | | |
| S. cerevisiae: UB26123: MATa/MATalpha | This study | N/A |
| trp1::cup1::Hphmx/irt1::cup1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::HSP60uORFdel-3V5-NDC80term::LEU2/leu2::HSP60uORFdel-3V5-NDC80term::LEU2 | | |
| S. cerevisiae: UB26124: MATa/MATalpha | This study | N/A |
| trp1::cup1::Hphmx/irt1::cup1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::MSC6uORFdel-3V5-NDC80term::LEU2/leu2::MSC6uORFdel-3V5-NDC80term::LEU2 | | |
| S. cerevisiae: UB26705: MATa/MATalpha | This study | N/A |
| trp1::cup1::Hphmx/irt1::cup1::Hphmx ime4::cup1::NAT/ime4::cup1::NAT leu2::HSP60-3V5-NDC80term::LEU2/ leu2::HSP60-3V5-NDC80term::LEU2 upf1::NAT/upf1::NAT | | |

(Continued on next page)
| REAGENT or RESOURCE SOURCE | IDENTIFIER |
|-----------------------------|------------|
| **S. cerevisiae: UB26707: MATa/MATalpha** | This study N/A |
| **S. cerevisiae: UB26708: MATa/MATalpha** | This study N/A |

**Oligonucleotides**

| 3V5 probe primers | See Table S7 N/A |
| ACT1 qPCR primers | See Table S7 N/A |
| PFY1 qPCR primers | See Table S7 N/A |
| HSP60/UBI qPCR primers | See Table S7 N/A |
| APL4 qPCR primers | See Table S7 N/A |
| HSP60 qPCR primers | See Table S7 N/A |
| MSC6 qPCR primers | See Table S7 N/A |

**Software and algorithms**

| cutadapt, v2.3 | Martin, 2011 | https://cutadapt.readthedocs.io/en/stable/ |
| STAR, v2.5.3a | Dobin et al., 2013 | https://bioconductor.org/packages/release/bioc/html/STAR.html |
| BSgenome, v1.50.0 | Pagè, 2018 | https://bioconductor.org/packages/release/bioc/html/BSgenome.html |
| CAGEr, v1.24.0 | Haberle et al., 2015 | https://bioconductor.org/packages/release/bioc/html/CAGEr.html |
| DESeq2, v1.22.2 | Love et al., 2014 | https://bioconductor.org/packages/release/bioc/html/DESeq2.html |
| MinKNOW, v1.10.23 | Oxford Nanopore Technologies | https://nanoporetech.com/ |
| Albacore, v2.1.10 | Oxford Nanopore Technologies | https://nanoporetech.com/ |
| minimap2, v2.9.720 | Li, 2018 | https://github.com/lh3/minimap2 |
| Meme | Bailey and Elkan, 1994 | http://meme-suite.org/tools/meme |
| bowtie2, v2.3.4.4 | Langmead and Salzberg, 2012 | http://bowtie-bio.sourceforge.net/bowtie2/index.shtml |
| macs2, v2.1.1.20160309 | Zhang et al., 2008 | https://github.com/macs3-project/MACS |
| bedGraphToBigWig, v4 | Kent et al., 2010 | https://bioconda.github.io/recipes/ucsc-bedgraph2bigwig/README.html |
| deeptools2, v3.0.1 | Ramirez et al., 2016 | https://deeptools.readthedocs.io/en/develop/ |
| phastCons | Siepel et al., 2005 | http://compgen.cshl.edu/phast/phastCons-HOWTO.html |
| salmon, v0.13.1 | Patro et al., 2017 | https://combine-lab.github.io/salmon/ |
| matplotlib | Hunter, 2007 | https://matplotlib.org/ |
| Image Studio Lite | LI-COR | https://www.licor.com/bio/image-studio-lite/ |
| Graphpad Prism | GraphPad Software | https://www.graphpad.com/scientific-software/prism/ |
| bedtools | Quinlan and Hall, 2010 | https://bedtools.readthedocs.io/en/latest/ |
| GSEA, v4.1.0 | Subramanian et al., 2005 and Mootha et al., 2003 | https://www.gsea-msigdb.org/gsea/index.jsp |
| DANPOS2, v2.2.2 | Chen et al., 2013 | https://sites.google.com/site/danposdoc/ |
**RESOURCE AVAILABILITY**

**Lead contact**
The Lead Contact, Elçin Ünal (elcin@berkeley.edu), can provide further information and fulfill resource requests.

**Materials availability**
All yeast strains generated in this study are available and can be requested by contacting the Lead Contact.

**Data and code availability**
Data generated in this study are available at NCBI GEO under the accession ID GSE140177 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE140177). The custom code used for the analysis is available in the following code repository: https://github.com/atresen/LUTI_key_features.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Strain construction and cloning**
All strains for the meiotic experiments were derived from the SK1 background and contained the previously published pCUP1-IME1/ pCUP1-IME4 meiotic synchronization system (Benchowitz et al., 2013). The genotypes of all the strains used in this study are listed in the Key resources table. For LUTI-regulated genes of interest (SWI4, APL4, CDC60, HSP60, and MSC6), the wild-type gene with between 800 and 1200 bp upstream of the CDS, depending on the length of the 5’-extension, and a C-terminal 3V5 epitope tag with the NDC80 terminator were cloned into a LEU2 single integration vector by Gibson Assembly (Gibson et al., 2009). LUTI promoter deletion strains were similarly constructed for each gene with the cloned upstream region only containing the sequence downstream of the LUTI TSS as determined by TL-seq. For genes in which uORF deletions were made (APL4, HSP60, MSC6) gBlocks with the ATG > ATC mutations were cloned by Gibson Assembly into the 3V5-tagged LEU2 single integration vectors described above. In all strains, the WT copies of these genes remained untouched. To construct the LexO-HSP60-ubiGFP reporter construct, a three-piece Gibson Assembly was performed in which a HIS3 single integration plasmid carrying ubiGFP was engineered to accept fragments carrying the 8xLexO-pCYC1 and the HSP60 leader sequences. The hsf1 mutation was generated in the above plasmid by Q5 site-directed mutagenesis (E0552s, New England Biolabs). All single integration plasmids were digested with PmeI before transformation and the correct integrations were confirmed by PCR. All strains and plasmids used in this study are available upon request.

**pCUP1-IME1/pCUP1-IME4 sporulation**
For genome-wide cell collections, cells were prepared to progress synchronously through meiosis as described in Chia and van Werven (2016). Briefly, liquid YPD (1% yeast extract, 2% peptone, 2% dextrose, tryptophan (96 mg/L), uracil (24 mg/L), and adenine (12 mg/L) cultures were started and grown for ~6 hours at 30°C until they reached an OD_{600} between 0.5 and 2.0. They were then diluted to an OD_{600} of 0.05 in reduced YPD (1% yeast extract, 2% peptone, 1% dextrose, uracil (24 mg/L), and adenine (12 mg/L)) and allowed to grow for 16–18 hours at 30°C until they reached an OD_{600} > 6. Cells were transferred to supplemented sporulation media or SPO (1% potassium acetate at pH 7.0 supplemented with adenine and uracil to 40 mg/L and histidine, leucine, and tryptophan to 20 mg/L, and 0.02% raffinose) with a final OD_{600} of 2.5 for 2 hours at 30°C before inducing pCUP1-IME1 and pCUP1-IME4 with 50 μM CuSO4. In all other meiotic experiments, cells were prepared as in Carlile and Amon (2008) but with 2% potassium acetate and supplements as above. Briefly, after 24 hours of growth in YPD at RT, saturated cells (OD_{600} > 10) were diluted to an OD_{600} of 0.2-0.3 and inoculated in BYTA (1% yeast extract, 2% bacto tryptone, 1% potassium acetate, and 1.02% potassium phthalate) for 16-18 hours of growth at 30°C (ideally to an OD_{600} of > 5). Enough cells to give a final OD_{600} of 1.85 were transferred into SPO with 2% acetate at 30°C. After 2 hours in SPO, IME1 and IME4 were induced with 50 μM CuSO4. In meiotic experiments with the LexA-ER-AD inducible system, cells were induced with either 2 or 5 nM of β-estradiol.

**Mitotic cell collection**
Exponentially growing cells from W303 background were back diluted to an OD_{600} of 0.2 and then treated with either 2 or 10 nM of β-estradiol. Cells were collected before induction as well as 1.5 and 3 hours after induction.
METHOD DETAILS

RNA extraction for TL-seq, nanopore sequencing and RNA-seq

At the indicated time points, 50 OD_{600} units of cells were collected by vacuum filtration and snap frozen in liquid nitrogen. Cells were thawed on ice and resuspended in 400 µl TES buffer (10 mM Tris pH 7.5, 10 mM EDTA, 0.5% SDS). An equal volume of Acid Phenol:Chloroform:Isoamyl alcohol (125:24:1; pH 4.7) was added to cells, and they were incubated at 65°C for 45 minutes in a Thermomixer (Eppendorf) shaking at 1400 RPM. The aqueous phase was transferred to a second tube of acid phenol. Samples were incubated at RT for 5 minutes while shaking at 1400 RPM in a Thermomixer. A final extraction with chloroform was performed. The aqueous phase was vortexed with chloroform for 30 s, separated by centrifugation, and then precipitated in isopropanol and sodium acetate for overnight at –20°C. Pellets were washed with 80% ethanol and resuspended in DEPC water for 10 min at 37°C. Total RNA was quantified using the Qubit RNA BR Assay Kit (Q10211, ThermoFisher Scientific).

Transcript leader sequencing (TL-seq)

The 5’ end sequencing approach was performed as in (Wu et al., 2018). At least 5 µg of mRNA was purified from total RNA using the Poly(A) Pursit MAG kit (AM1922, Ambion). mRNAs were fragmented for 3 minutes at 70°C using a Zinc-based alkaline fragmentation reagent (AM8740, Ambion). RNAs were cleaned up using RNasey MinElute Cleanup Kits (74204, QIAGEN) to enrich for 200-300 nucleotide fragments. These fragments were dephosphorylated with 30 units of recombinant shrimp alkaline phosphatase (M0371, New England Biolabs) for 1 hour at 37°C with RNasin Plus (N2611, Promega). The RNA was extracted with Acid Phenol:Chloroform:Isoamyl alcohol (125:24:1, pH 4.7) and precipitated in ethanol with 0.3 M sodium acetate and 1 µl linear acrylamide (AM9520, Ambion). RNA was then subjected to a decapping reaction with 2 units of Cap-Clip acid pyrophosphatase (CC150111H, Tebu-Bio) and with RNasin Plus. RNA was then again extracted using Acid Phenol:Chloroform:Isoamyl alcohol (125:24:1) and precipitated in ethanol. Some RNA from a premeiotic time point was set apart without the decapping reaction as a non-decapping control. Subsequently, the RNA was mixed with 10 µM of custom 5’ adaptor and the ligation reaction was done using T4 RNA ligase 1 (M0437M, New England Biolabs) and RNasin Plus. The ligation reaction was cleaned up with the RNasey MinElute Cleanup Kit (74204, QIAGEN) and RNAs were mixed with 2.5 µM random hexamers (N8080127, ThermoFisher Scientific) and RNasin Plus, denatured at 65°C for 5 minutes and cooled on ice. Reverse transcription reactions were carried out using SuperScript IV reverse transcriptase (18090010, Invitrogen). The RNA templates were degraded by incubating reactions with 5 units of RNase H (M0297, New England Biolabs) and 1.0 µL of RNase cocktail enzyme mix (AM2286, Ambion). DNA products were purified using 1.8x volume of HighPrep PCR beads (AC-60050, MagBio). Purified products were subjected to second strand synthesis using 0.3 µM of second strand biotinylated primer and the KAPA Hi-Fi hot start ready mix (KK2601, Roche). The second strand reaction was carried out at 95°C for 3 minutes, 98°C for 15 s, 50°C for 2 minutes, 65°C for 15 minutes and held at 4°C. Double stranded product (dsDNA) was purified with 1.8x volume HighPrep PCR beads and concentration was quantified using the Qubit dsDNA HS assay kit (Q32851, Invitrogen). 25 ng of dsDNA was then used as input for the KAPA Hyper Prep Kit (KX8504, Roche) and ligated to KAPA single indexed adapters Set B (KK8702, Roche). Samples were processed according to manufacturer’s instructions with one exception: just prior to the library amplification step, samples were bound to MyOne Streptavidin C1 Dynabeads (65001, ThermoFisher Scientific) to capture biotinylated dsDNA. Library amplification over 14 PCR cycles was done on the biotinylated dsDNA fraction bound to the beads. Amplified libraries were quantified by Qubit, and adaptor-dimers were removed by electrophoresis of the libraries on Novex 6% TBE gels (EC62655BOX, Invitrogen) at 120 V for 1 hour, and excising the smear above ~150 bp. Gel slices containing libraries were shredded by centrifugation at 13000 g for 3 minutes. Gel shreds were re-suspended in 500 µL crush and soak buffer (500 mM NaCl, 1.0 mM EDTA and 0.05% v/v SDS) and incubated at 65°C for 2 hours on a thermomixer (1400 rpm for 15 s, rest for 45 s). Subsequently, the buffer was transferred into a Costar SpinX column (8161, Coming Incorporated) with two 1 cm glass pre-filters (1823010, Whatman). Columns were centrifuged at 13000 g for 1 minute. DNA libraries in the flowthrough were precipitated at –20°C overnight in ethanol with 0.3 M sodium acetate and 1 µl linear acrylamide (AM9520, Ambion). Purified libraries were further quantified and inspected on a Tapestation (Agilent Technologies, Inc). The libraries were sent for 100 bp SE sequencing on an Illumina HiSeq 4000 at the Vincent J. Coates Genomics Sequencing Laboratory.

polyA selection for Nanopore sequencing and RNA-seq

PolyA selection was performed on 100 µg of RNA using 150 µL of oligo-dT DynaBeads (61002, ThermoFisher Scientific). RNA was denatured at 80°C for 2 minutes in binding buffer (10 mM Tris-HCl pH 7.5, 0.5 M LiCl, 3.35 mM EDTA) before being placed on ice. At RT the oligo-dT beads were added to the sample and together they were incubated at room temperature for 5 minutes. Beads were washed 2x in Buffer B (10 mM Tris-HCl pH 7.5, 0.15 M LiCl, 1 mM EDTA). PolyA-selected RNA was eluted from the beads by heating at 80°C for 2 minutes in 10 mM Tris 7.0. RNA was quantified with a Qubit using the RNA HS assay kit (Q32852, ThermoFisher Scientific).

Nanopore sequencing

500 ng of polyA-selected RNA was used as directed in the Direct RNA Sequencing Kit (SQK-RNA001, Oxford Nanopore Technologies). The library was loaded onto a miniON (MIN–101B, Oxford Nanopore Technologies) with an R9.4.1 flow cell (FLO-MIN106, Oxford Nanopore Technologies). MinKNOW (v1.10.23, Oxford Nanopore Technologies) was run without live base calling for 48 hours.
Bases were called from fast5 files with the Albacore script read_fast5_basecaller.py (v2.1.10, Oxford Nanopore Technologies). 491,142 reads were sequenced. Reads were aligned to the genome with minimap2 (v2.2-r1720; Li, 2018) using options --ax splice --k14 --uf. Bam files were visualized directly in IGV.

**Chromatin immunoprecipitation (ChIP)**

For Ume6-3V5 ChIP, 300 OD600 units of stationary phase cells in BYTA were collected. In histone modification experiments, 112.5 OD600 units of cells were collected during both premeiotic stage and meiotic prophase. In all instances, cells were fixed with 1% formaldehyde. The formaldehyde was quenched with 125 mM glycine, cells were pelleted, washed with PBS, and then lysed by Bead-beater (Mini-Beadbeater-96, Biospec Products) with zirconia beads for 4 × 5 minutes in FA Buffer (50 mM HEPES pH 7.5, 150 mM NaCl, 1 mM EDTA, 1% Triton, 0.1% sodium deoxycholate, 0.1% SDS supplemented with cOmplete protease inhibitor tablets (11873580001, Roche). Note that for the Ume6-3V5 ChIPs, due to the number of cells collected, lysates were prepared in 3 separate tubes. They were processed separately until after the IP. Lysates were collected and centrifuged for 3 minutes at 2000 g. The supernatants were transferred to fresh tubes and centrifuged for 15 minutes at 20,000 g. The supernatant was discarded, and the pellet of chromatin was resuspended in 1 mL FA Buffer. Samples were sonicated: 30 s on, 30 s off, for 5 minutes with a Bioruptor Pico (Diagenode) to an average fragment size of ~200 bp. The supernatant from a 1 minute 20,000 g centrifugation was carried forward to the IP.

From the isolated chromatin, 30 μL were set aside as input. For Ume6-3V5 ChIPs, to each of 3, 600 μL chromatin aliquots, 1 μL of a mouse α-V5 antibody (R960-25, Thermo Fisher) was added. For histone modification ChIPs, 3 μL of an antibody specific to either H3K36me3 (ab9050, abcam) or H3K4me2 (ab32356, abcam) was added to the chromatin. Samples were incubated for 2 hours with nutation. Protein A Dynabeads (10001D, Invitrogen) were blocked with 0.1% BSA in FA Buffer for at least 2 hours at 4°C. They were washed 2 x with FA Buffer, resuspended with FA Buffer to their original volume, and 10-20 μL (10 for V5 and 20 for histone modification IP) of the resuspended beads were added to each tube of chromatin. The chromatin-bead mixture was nutated at 4°C overnight. The IP was washed 6 times: 2x FA Buffer, 2x Buffer 1 (FA Buffer, 260 mM NaCl), and 2x Buffer 2 (10 mM Tris pH 8.0, 250 mM LiCl, 0.5% NP-40, 0.5% sodium deoxycholate, 1 mM EDTA). Between each wash, samples were nutated for 5 minutes at 4°C. To IP and input samples, 150 μL or 120 μL of TE (10 mM Tris pH 8.0, 1 mM EDTA) with 1% SDS was added, respectively. The precipitate was eluted from the beads by shaking at 1200 RPM in a Thermomixer (ThermoFisher Scientific) at 65°C. Eluates were treated with ~0.33 mg/mL RNase A (12091021, Invitrogen) for 30 minutes at 37°C and then with ~1.2 mg/mL Proteinase K (3115879001, Roche) overnight at 65°C. Samples were cleaned up with QiaQuick PCR Purification Kit (28106, Qiagen) and eluted in EB. DNA was quantified by Qubit with the dsDNA HS Assay Kit (Q32854, Invitrogen). Libraries were prepared as instructed by the ThruPLEX DNA-seq Kit (R400427, Takara). For input and H3K36me3 IP samples, 15 ng of starting material was used with 7 rounds of PCR. For all other IP samples, 0.5-1 ng of starting material was used with 11 rounds of PCR. AMPure XP beads (A63881, Beckman Coulter) were used to select fragments between 200-500 bp. Samples were submitted for 50 bp SE sequencing by the Vincent J. Coates Genomics Sequencing Laboratory with a HiSeq4000.

**RNA-seq**

The RNA-seq libraries were prepared using the NEXTflex™ Rapid Directional mRNA-Seq Kit (NOVA-5138, Bio Scientific) according to manufacturer’s instructions. 100 ng of polyA selected mRNA was used for all libraries. Libraries were quantified using the Agilent 4200 TapeStation (Agilent Technologies, Inc.). AMPure XP beads (A63881, Beckman Coulter) were used to select fragments between 200-500 bp. Samples were submitted for 150 bp SE sequencing by the Vincent J. Coates Genomics Sequencing Laboratory with a HiSeq4000.

**Immunoblotting**

Immunoblotting was performed as previously described in Chen et al. (2017). To track tagged proteins of interest, membranes were incubated with either a mouse α-V5 antibody (R960-25, Thermo Fisher) or a mouse α-GFP antibody (632381, Takara) to a dilution of 1:2000 in Odyssey Blocking Buffer (PBS) (LI-COR Biosciences) with 0.01% Tween. Both V5 and GFP immunoblots were also incubated with a rabbit α-hexokinase (α-Hxk2) antibody (H2035, US Biological) diluted between 1:15,000-1:20,000. Secondary antibodies included a α-mouse antibody conjugated to IRDye 800CW (926-32212, LI-COR Biosciences) and a α-rabbit antibody conjugated to IRDye 680RD (926-68071, LI-COR Biosciences). They were each diluted to 1:15,000 in Odyssey Blocking Buffer (PBS) with 0.01% Tween. An Odyssey system (LI-COR Biosciences) was used to image all blots, and Image Studio Lite (LI-COR Biosciences) was used for all quantification.

**RNA extraction for blotting and RT-qPCR**

At the indicated time points, between 4 and 5 OD600 units of cells were collected by centrifugation and snap frozen in liquid nitrogen. Cells were thawed on ice and resuspended in TES (10 mM Tris pH 7.5, 10 mM EDTA, 0.5% SDS). An equal volume of Acid Phenol-Chloroform:Isoamyl alcohol (125:24:1; pH 4.7) was added to cells, and they were incubated at 65°C for 45 minutes in a Thermomixer C (Eppendorf) shaking at 1400 RPM. The aqueous phase was transferred to a second tube with chloroform. The aqueous phase was...
vortexed with the chloroform for 30 s, separated by centrifugation, and then precipitated in isopropanol and sodium acetate overnight at −20 °C. Pellets were washed with 80% ethanol and resuspended in DEPC water for 10 min at 37 °C. Total RNA was quantified by Nanodrop.

**RNA blotting**

RNA blot analysis protocol was performed as described previously (Koster et al., 2014) with minor modifications. 8 μg of total RNA was denatured in a glyoxal/DM SO mix (1M deionized glyoxal, 50% v/v DMSO, 10 mM sodium phosphate (NaPi) buffer pH 6.5–6.8) at 70 °C for 10 minutes. Denatured samples were mixed with loading buffer (10% v/v glycerol, 2 mM NaPi buffer pH 6.5–6.8, 0.4% w/v bromophenol blue) and separated on an agarose gel (1.1% w/v agarose, 0.01 M NaPi buffer) for 3 hr at 100 V. The gels were then soaked for 25 minutes in denaturation buffer (0.05 N NaOH, 0.15 M NaCl) followed by 20 minutes in neutralization buffer (0.1 M Tris-HCl pH 7.5, 0.15 M NaCl). RNA was transferred to nitrocellulose membrane for 1 hour via vacuum transfer as described in Stratagene’s Membranes Instruction Manual (Stratagene) and crosslinked using a Stratalinker UV Crosslinker (Stratagene). RNA bands were visualized using methylene blue staining. The membranes were blocked in ULTRAhyb Ultrasensitive Hybridization Buffer (AM8669, Thermofisher Scientific) for 3 hours before overnight hybridization. Membranes were washed twice in Low Stringency Buffer (2x SSC, 0.1% SDS) and three times in High Stringency Buffer (0.1X SSC, 0.1% SDS). All hybridization and wash steps were done at 42 °C. Radioactive probes were synthesized using a Prime-It II Random Primer Labeling Kit (300385, Agilent Technologies, Inc).

**Micrococcal nuclease sequencing (MNase-seq)**

The protocol was adapted from Basic Protocol 1 in Rodriguez et al. (2014) with the following changes. In premeiotic stage and meiotic prophase, 112.5 OD600 units of cells were fixed in 1% formaldehyde with light shaking at RT for 15 minutes. Crosslinking was quenched by 125 mM of glycine for 5 minutes at RT. Cells were pelleted and washed twice with ice cold milliQ water. Cells were spheroplasted in 20 mL of Spheroplast Solution (1 M Sorbitol, 50 mM Tris pH 7.5, 10 mM β-ME) with 100 μL of 10 mg/mL zymolase until they appeared non-refractive and shadow-like after ~20-30 minutes. Spheroplasted cells were resuspended in 2 mL MNase Digestion Buffer (1 M Sorbitol, 50 mM NaCl, 10 mM Tris pH 7.5, 5 mM MgCl2, 1 mM CaCl2, 0.075% NP-40, 0.5 mM spermidine, 1 mM β-ME) if collected during premeiotic stage and 4 mL if collected during meiotic prophase, after completion of S-phase. Digestions were performed with 600 U of spheroplasts, 30 units of Exonuclease III (M0206S, New England Biolabs), and either 10, 20, or 40 units of MNase (LS004797, Worthington). Crosslinks were reversed, protein was degraded by Proteinase K (3115879001, Roche), and a phenol/chloroform/soamyl alcohol DNA extraction, ethanol precipitation, RNase A (12091021, Invitrogen) treatment, and phosphatase treatment were performed as described previously (Rodriguez et al., 2014). Size selection was performed by running samples on a 1.8% LMT agarose gel at 80 V for 40 minutes at room temperature and gel extracting the mononucleosome band with a Monarch Gel Extraction Kit (T102OS, New England Biolabs). Note that of the samples digested with 10, 20, and 40 units of MNase, only the samples with a ratio of mononucleosomes to dinucleosomes closest to 80/20 were size selected and carried forward for library preparation. Gel extracted samples were quantified by Qubit with the dsDNA HS Assay Kit (Q32854, Invitrogen). Libraries were prepared with 50 ng starting material as instructed by the ThruPLEX DNA-seq kit (R400427, Takara). Amplification was performed with 5 rounds of PCR. AMPure XP beads (A63881, Beckman Coulter) were used to select fragments between 150-500 bp. Samples were submitted for 100 bp PE sequencing to the Vincent J. Coates Genomics Sequencing Laboratory with a HiSeq4000.

**Reverse transcription-quantitative polymerase chain reaction (RT-qPCR)**

To 5 μg of isolated total RNA in 1x DNase I reaction buffer, 1 unit of DNase I was added. The reaction was incubated at 37 °C for 30 minutes. The DNase-treated RNA was extracted by Acid Phenol:Chloroform:isoamyl alcohol DNA extraction, ethanol precipitation, RNase A (12091021, Invitrogen) treatment, and phosphatase treatment were performed as described previously (Rodriguez et al., 2014). Size selection was performed by running samples on a 1.8% LMT agarose gel at 80 V for 40 minutes at room temperature and gel extracting the mononucleosome band with a Monarch Gel Extraction Kit (T102OS, New England Biolabs). Note that of the samples digested with 10, 20, and 40 units of MNase, only the samples with a ratio of mononucleosomes to dinucleosomes closest to 80/20 were size selected and carried forward for library preparation. Gel extracted samples were quantified by Qubit with the dsDNA HS Assay Kit (Q32854, Invitrogen). Libraries were prepared with 50 ng starting material as instructed by the ThruPLEX DNA-seq kit (R400427, Takara). Amplification was performed with 5 rounds of PCR. AMPure XP beads (A63881, Beckman Coulter) were used to select fragments between 150-500 bp. Samples were submitted for 100 bp PE sequencing to the Vincent J. Coates Genomics Sequencing Laboratory with a HiSeq4000.

**Quantitative and Statistical Analysis**

**TL-seq analysis**

From the sequencing reads, the 3’ Illumina adaptor (AGATCGGAAGAGC) was trimmed using cutadapt with the minimum-length option set to 20 bp (v2.3; Martin, 2011). From the 3’ trimmed output, the 5’ Illumina adaptor (CAGTCTGAGCAATACC) was trimmed from reads by cutadapt. To select for reads with the most 5’ end of a transcript, only reads in which the 5’ adaptor was recognized and then trimmed were carried forward. Reads were aligned by STAR (v2.5.3a; Dobin et al., 2013) using indices generated from an SK1 genome assembled by combined PacBio and Illumina sequencing (Yue et al., 2017). A custom SK1 genome was forged with BSgenome (v1.50.0; Pagé, 2018) using the above assembly (Yue et al., 2017). Bam files were imported into CAGEr and the CAGEr pipeline was applied to define TSSs and quantify transcript abundances as follows (v1.24.0; Haberle et al., 2015). Reads at TSSs were
Bigwig files for viewing in IGV were generated by macs2 bdgcmp with option –B –q 0.001–keep-dup ''all''–call-summits –nomodel –extsize 147 sample from macs2, all libraries were down-sampled to 2 million reads (v2.1.1.20160309; Zhang et al., 2008). Macs2 callpeak was 

Motif discovery
Meme was applied to the sequences 300 bp up and 300 bp downstream of each LUTI TSS with options –w 10 –dna –revcomp. (Bailey and Elkan, 1994). A motif was considered significant in an individual sequence if it had a combined match p value < 0.05.

Ume6 ChIP-seq analysis
Reads were aligned to the SK1 genome with bowtie2 (v2.3.4.3; Langmead and Salzberg, 2012; Langmead et al., 2019). Using random samples from macs2, all libraries were down-sampled to 2 million reads (v2.1.1.20160309, Zhang et al., 2008). Macs2 callpeak was used to call peaks in IP samples over input samples with options –B –q 0.001–keep-dup “all”–call-summits –nomodel–extsize 147. Bigwig files for viewing in IGV were generated by macs2 bdgcmp with option –m FE followed by bedGraphToBigWig (v4; Kent et al., 2010). Heatmaps and metagene plots centered around TSSs as defined by TL-seq were constructed with deepTools2 (v3.0.1; Ramirez et al., 2016). A 5’-extended or canonical target promoter was considered to be enriched by Ume6 if, in at least 2 of 3 ChIP replicates, a peak was called (log2 fold-change > 2 over input) within 300 bp of the transcript’s TSS.

Conservation of URS1 binding sites
The 5’-extended and canonical Ume6 targets enriched with both Ume6 and a URS1 binding site in their promoters were analyzed for degree of conservation. Meme was run on the sequences ± 300 bp from their TSSs with options –w 10 –dna –revcomp was run to identify the location of the URS1 binding sites with regard to the TSS. Using a custom python script, the midpoint of each URS1 binding motif was determined relative to the CDS associated with either the LUTI or canonical target. Subsequently, the chromosome locations of the URS1 midpoint in the sacCer3 S288C reference genome were found. To assess conservation of the regions around URS1 motifs at 5’-extended and canonical targets, phastCons (Siepel et al., 2005) was performed with options –target-coverage 0.025 -expected-length 12 –rho 0.4. The tree phylogeny model and the genome alignments of S. cerevisiae, S. paradoxus, S. mikatae, S. kudriavzevii, S. bayanus, S. castelli, and S. kluveri were from Siepel et al. (2005). However, the alignments of S. castelli and S. kluveri were excluded in all analyses here because these yeast species have lost the IME1 and UME6 genes. Metagene plots and heatmaps of conservation were generated with deepTools2 (v3.0.1; Ramirez et al., 2016).

Gene set enrichment analysis (GSEA)
Normalized counts for each gene were calculated by DESeq2 using default options (v1.22.2; Love et al., 2014). The Saccharomyces cerevisiae collection of gene sets (updated from SGD on 2020-12-28) was downloaded from the Gene Ontology Consortium (Ashburner et al., 2000; Carbon et al., 2021). The collection was supplemented with a gene set including the 41 genes identified in Figure 2A of Brar et al. (2012) as being expressed during recombination/synaptonemal complex formation. GSEA was performed using the desktop app version 4.1.0 with the Collapse/Remap to gene symbols setting set to “No_Collapse” and the permutation type defined as gene_set. All other parameters were default settings.
uORF analysis
ATGs were counted and the codon frequency was determined with a custom python script. For genes with LUTIs, the counts and codon frequencies were determined for the region between the PROX TSS and the LUTI TSS. For all other genes, sequences from the 500 bp upstream of the TSS were used.
LUTIs with > 4 uORFs were analyzed to determine which of the uORFs were translated. Footprints were quantified for the first 6 codons of each uORF using the tools in Brar et al. (2012) and Ingolia et al. (2009). The ribosome footprinting data were taken from the 3 h time point in Cheng et al. (2018). Any uORFs with at least 4 footprint reads found across the first 6 codons of the gene were considered to be translated.

Chromatin modification analysis
Reads were aligned to the SK1 PacBio genome assembly with bowtie2 (v2.3.4.3; Langmead and Salzberg, 2012). Macs2 callpeak was used to call peaks in IP samples over input samples with options -B -q 0.01 -nomodel -extsize 147. Bigwig files for viewing in IGV and for further quantification were generated by macs2 bdgcmp with option -m FE followed by bedGraphToBigWig (v4; Kent et al., 2010; Zhang et al., 2008). To quantify the change in H3K36me3 and H3K4me2 enrichment over the promoters of PROX transcripts, fold enrichment scores were extracted from regions 50 bp upstream and 500 bp downstream of the PROX TSS with bedtools (Quinlan and Hall, 2010). With custom python scripts, the scores from each bp of the upstream regions and each bp of the downstream regions were summed for each gene. The ratio of the upstream and the downstream region enrichments were quantified and the change in the score from premeiotic stage to meiotic prophase was determined. Ultimately, the mean of the FC was calculated from samples in triplicate. Heatmaps and metagene plots were prepared with deeptools2 (Ramirez et al., 2016).

RNA-seq analysis
Quantification of RNA as transcripts per million was done using salmon in the mapping-based mode with mapping validation (v0.13.1; Patro et al., 2017). FC quantification was performed by DESeq2 using default options (v1.22.2; Love et al., 2014). Scatterplots were generated with matplotlib (Hunter, 2007). For all scatterplots in Figures 2 and S3 with RNA-seq data, the spearman correlation coefficient was calculated and is displayed on the plot. Heatmaps were generated with pheatmap.

MNase-seq analysis
Reads were aligned to the SK1 genome with bowtie2 (v2.3.4.3; Langmead and Salzberg, 2012). To select for only fragments between 130 and 170 bps, alignmentSieve from deeptools2 was performed (Ramirez et al., 2016). BigWig files were generated by bamCoverage with options --MNase --bs 1 --normalizeUsing CPM (Ramirez et al., 2016). DANPOS (v2.2.2) was run to determine various aspects of nucleosome location and occupancy and fuzziness (Chen et al., 2013). A custom python script was used to assign the locations of +1 and −1 nucleosome with respect to PROX TSSs.

Clustered heatmap
Spearman’s correlation coefficients were calculated between all pairs of features in 6A. The coefficients are displayed on the scatterplots in Figure S7. P values of significant features are indicated in the text.
Supplemental information

Integrated genomic analysis reveals key features of long undecoded transcript isoform-based gene repression

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Supplemental Figure Legends:

Figure S1. TL-seq and direct RNA sequencing for a LUTI gene and a meiosis-specific gene, Related to Figure 1

A. Genome browser views of TL-seq and direct RNA sequencing using Nanopore at the NDC80 locus. The TL-seq and Direct RNA sequencing were performed in cells with the pCUP-IME1/pCUP-IME4 meiotic induction system (UB14584). The premeiotic time point was taken after 2 hours in SPO, and the meiotic prophase time point was taken after 4 hours in SPO, which corresponds to 2 hours after 50 µM CuSO₄ addition. B. As in (A) but for SPO11.

Figure S2. Ume6 enrichment for the genes without 5’ extensions and RNA-seq browser from UME6 and ume6 T99N strains, Related to Figure 1

A. Heatmap of Ume6 enrichment over input for all of the genes without 5’-extensions (n=5298). A representative image from one of three replicates is shown. B-C. Genome browser views of RNA-seq at the (B) NDC80 and the (C) APL4 locus. RNA-seq was performed on strains expressing either WT Ume6 (UB20649) or Ume6 with a T99N mutation (UB22629). The strains also harbor the pCUP1-IME1/pCUP1-IME4 meiotic induction system and were induced with 50 µM CuSO₄ at 2 hours in sporulation medium (SPO). Cells were collected after 2 or 4 hours in SPO.

Figure S3. Contribution of LUTI and PROX isoforms to total RNA abundance and assessment of RNA-seq similarity between this study and Cheng et al. 2018, Related to Figure 2

A. Scatterplots comparing RNA-seq and TL-seq. Cells harboring the pCUP1-IME1/pCUP1-IME4 meiotic induction system and were induced with 50 µM CuSO₄ at 2 hours in sporulation medium (SPO). Analysis was performed with data from cells collected during meiotic prophase (4 hours in SPO). In all plots the teal data points display RNA-seq vs TL-seq transcripts per million for genes without 5’-extensions. All orange points indicate loci with 5’-extended transcripts. TL-seq values for genes with 5’-extended transcripts were calculated either from PROX-specific TPM (Left), LUTI-specific TPM (Middle), or the addition of both PROX- and LUTI-specific TPM (Right). The Spearman’s rank correlation coefficient (p) was calculated and is displayed on the upper left corner for both teal and orange points separately. B. Scatterplots comparing RNA-seq performed in this study to RNA-seq performed in Cheng et al. 2018. In this study, cells with the pCUP1-IME1/pCUP1-IME4 meiotic induction system (UB14584) were collected for RNA-seq (in duplicate) after 4 hours in sporulation medium (SPO), which corresponds to 2 hours after induction of meiosis by 50 µM CuSO₄. In Cheng et al., a prototrophic strain was used. Cells were collected after 1.5 (left), 3 (middle) or 4.5 (right) hours in SPO. The Spearman’s rank correlation coefficient (p) was calculated and is displayed on the upper left corner. C. GSEA enrichment plot for the SC/Recombination gene set defined in Brar et al. (2012). Analysis was performed with RNA-seq (duplicates) collected from cells harboring the pCUP1-IME1/pCUP1-IME4
meiotic induction system (UB14584). Enrichment was determined by comparing gene expression at 4 hours in SPO (meiotic prophase) to 2 hours (premeiotic).

**Figure S4. The effect of uORF deletion or upf1Δ on LUTI abundance and protein level, Related to Figure 3**

A. Cells with either wild-type uORFs or uORFs with ATGs mutated to ATC and a 3V5 tagged APL4 (WT: UB18539, uORFΔ: UB26122), HSP60 (WT: UB18335, uORFΔ:UB26123), or MSC6 (WT: UB18238, uORFΔ:UB26124) were induced to undergo meiosis with 50 µM CuSO₄ after 2 hours in SPO using the pCUP1-IME1/pCUP1-IME4 meiotic induction system. Immunoblot was performed against the 3V5 epitope and Hxk2 as a loading control. One of two replicates is displayed. B-M. Wild-type or upf1Δ strains harboring 3V5 tagged APL4 (WT: UB18539, upf1Δ: UB26707), HSP60 (WT: UB18335, upf1Δ:UB26705), or MSC6 (WT: UB18238, uORFΔ:UB26708) were induced to undergo meiosis with 50 µM CuSO₄ after 2 hours in SPO using the pCUP1-IME1/pCUP1-IME4 meiotic induction system. (B-D) RNA blots whereby transcripts were detected using a probe directed against the 3V5 epitope and its linker sequence. rRNA was stained by methylene blue as a loading control. (E-G) qPCR quantification of LUTI transcripts using the matched RNA from B-D. Quantification was performed in reference to the levels of the meiotic housekeeping gene PFY1. (H-J) Immunoblot with α-3V5 and α-Hxk2 antibodies. (K-M) Quantification of immunoblots in H-J. Abundance of the 3V5 tagged protein was quantified by first normalizing to the Hxk2 loading control and then to the first time point (0 h).

**Figure S5. Single gene analysis of LUTI deletion, Related to Figure 4**

A-B. LUTI deletion analysis of additional genes. Cells with the pCUP1-IME1/pCUP1-IME4 meiotic induction system were induced to enter meiosis with 50 µM CuSO₄ after 2 hours in SPO. A. Experiments were performed with a MSC6-3V5 tagged strain harboring either wild type (UB18238) or LUTI deletion (UB18190). RNA blots and Immunoblots were performed on samples collected between 0-6 hours in SPO. RNA blots were performed with a probe specific for 3V5 and its linker. Methylene blue detection of rRNA bands was the loading control. Immunoblots were performed with a α-V5 antibody to recognize the 3V5-tagged proteins. Hxk2 was used as a loading control. The blots represent one of two replicates. Quantification of the immunoblot is displayed on the right. 3V5 signal was quantified relative to Hxk2 and then relative to 0 hr. Graphs represent one of two replicates B. Same as A, but for APL4-3V5 tagged strains with either wild type (UB18539) or LUTI deletion (UB18181).

**Figure S6. Alterations to chromatin over the PROX promoter upon LUTI expression, Related to Figure 5**

A. Metagene analysis of H3K36me3 ChIP-seq for LUTI and non-LUTI genes in both the premeiotic and meiotic prophase stages. The ChIP was performed in cells with the pCUP1-IME1/pCUP1-IME4 meiotic induction system (UB14584). The premeiotic time point was taken after 2 hours in SPO, and the meiotic prophase time point was taken
after 4 hours in SPO, which corresponds to 2 hours after 50 µM CuSO₄ addition. The images are from one of three replicates. **B.** Same as (A) but for H3K4me2. **C.** Heatmap of H3K36me3 ChIP-seq in premeiotic and meiotic prophase. The ChIP was performed in cells with the pCUP1-IME1/pCUP1-IME4 meiotic induction system (UB14584). The premeiotic time point was taken after 2 hours in SPO, and the early meiotic time point was taken after 4 hours in SPO, which corresponds to 2 hours after 50 µM CuSO₄ addition. The plot is centered around the PROX TSS. Top: LUTIs, bottom: canonical transcripts. The images are from one of three replicates. **D.** Same as (C) but for H3K4me2. **E-F.** Genome browser views of MNase-seq and TL-seq. The TL-seq and MNase DNA digestion was performed in cells with the pCUP-IME1/pCUP-IME4 meiotic induction system (UB14584). The premeiotic time point was taken after 2 hours in SPO, and the meiotic prophase time point was taken after 4 hours in SPO, which corresponds to 2 hours after 50 µM CuSO₄ addition. Three candidate genes, RPO41 (E), APL4 (F) and SWI4 (G), all of which are robustly transcriptionally repressed are shown. The green box denotes the approximate location of the PROX promoter.

**Figure S7.** Reduction in PROX transcript abundance shows significant correlation with a select number of criteria, Related to Figure 6

Scatterplots of the relationships tested in Figure 6A. Spearman’s correlation coefficients (ρ) are displayed on the upper left corner.

**Supplemental Table Legends:**

**Table S5.** Translational efficiency (TE) fold-change between meiotic prophase and premeiotic stage for LUTIs with < 4 ATG codons in the region between the LUTI and the PROX TSSs, Related to Figure 3

**Table S7.** Primers used in this study, Related to STAR Methods
Figure S1, Tresenrider et al.
Figure S2, Tresenrider et al.
Figure S3, Tresenrider et al.

ρ = 0.732
ρ = 0.419
ρ = 0.347
ρ = 0.602

Enrichment plot: Brar et al. SC and Recombination Cluster

Enrichment profile

Hits

Ranking metric scores

Zero cross at 2651

'1' (positively correlated)

'0' (negatively correlated)
Figure S5, Tresenrider et al.
Figure S7, Tresenrider et al.
### Table S5. LUTI candidates with fewer than 4 uORFs

| Gene     | uORF number | TE fold-change |
|----------|-------------|----------------|
| PLM2     | 0           | 4.894          |
| YCL057C-A| 0           | 3.411          |
| ELO1     | 1           | 1.078          |
| COX16    | 1           | 0.995          |
| ULP2     | 1           | 0.057          |
| MNE1     | 1           | 0.005          |
| ITR2     | 2           | 1.906          |
| YNL305C  | 2           | 0.193          |
| VPS30    | 3           | 0.236          |
| APL4     | 3           | 0.193          |
| Primer Name       | Oligonucleotide sequence from 5’ to 3’                      |
|------------------|------------------------------------------------------------|
| 3V5_probe_F      | CTAGTGGATCCAGGTAACCTAT                                     |
| 3V5_probe_R      | TAATACGACTCACTATAGGCGAGTCCTAATAGAGGATTAGG                |
| ACT1_F           | GTACCACCATGGTCCCCAGGTATT                                  |
| ACT1_R           | AGATGGACCACCTTTCGTCGT                                     |
| PFY1_F           | ACGGTTGGTGGATAATGAGC                                     |
| PFY1_R           | ACCTGACATGATGTGCTAGG                                      |
| HSP60_F          | TGGTTGCGTCTTCAATGAC                                       |
| UBI_R            | GGTCAAAAGTCTTGAGCAAATCTG                                  |
| APL4_qPCR_F      | GCTACAGGAAGGTGGAAGAG                                       |
| APL4_qPCR_R      | CTGCACCTAAGCTACCTAC                                       |
| HSP60_qPCR_F     | CGCCAAGACTAGTTGTGCAA                                       |
| HSP60_qPCR_R     | CAGATACAGATGGGCGGACTA                                     |
| MSC6_qPCR_F      | AGGTTGCTCCGGCACTAAGT                                      |
| MSC6_qPCR_R      | GTGCAGAGCCAACTTTAAGCA                                     |