Transcriptional repression of CDC6 and SLD2 during meiosis is associated with production of short heterogeneous RNA isoforms

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Abstract
Execution of the meiotic and mitotic cell division programs requires distinct gene expression patterns. Unlike mitotic cells, meiotic cells reduce ploidy by following one round of DNA replication with two rounds of chromosome segregation (meiosis I and meiosis II). However, the mechanisms by which cells prevent DNA replication between meiosis I and meiosis II are not fully understood. Here, we show that transcriptional repression of two essential DNA replication genes, CDC6 and SLD2, is associated with production of shorter meiosis-specific RNAs containing the 3′ end of both genes. Despite the short CDC6 RNA coding for a short protein (Cdc6short), this protein is not essential for meiosis and it does not have either a positive or negative impact on DNA replication. Production of CDC6short mRNA does not require the upstream CDC6 promoter (P_{CDC6}) and is not a processed form of the full-length RNA. Instead, CDC6short depends on transcription initiation from within the ORF upon repression of P_{CDC6}. Finally, using CDC6 genes from related yeast, we show that repression of full-length CDC6 mRNA is evolutionarily conserved and that this repression is consistently associated with production of unique short CDC6 RNAs. Together, these data demonstrate that meiotic cells transcriptionally repress full-length CDC6 and SLD2, and that inactivation of P_{CDC6} results in heterogeneous transcription initiation from within the CDC6 ORF.

Keywords DNA replication · Meiosis · Cell cycle · Cdc6 · Sld2 · Mcm2-7

Introduction
Sexual reproduction requires the generation of haploid gametes from diploid cells. To produce these gametes, cells undergo meiosis, a specialized cell division program that is designed to reduce ploidy by half (Ohkura 2015). During meiosis, a single round of DNA replication (S-phase) is followed by two rounds of chromosome segregation (the meiotic divisions), meiosis I (MI) and meiosis II (MII). In contrast, ploidy is maintained during mitotic cell divisions which alternate between single rounds of DNA replication and chromosome segregation. A key requirement of the meiotic program is that DNA replication must be inhibited between the meiotic divisions. This inhibition targets multiple proteins acting at distinct steps during DNA replication initiation (Holt et al. 2007; Phizicky et al. 2018), but the mechanisms of inhibition are not fully understood.

Mitotic cells use oscillations of cyclin-dependent kinase (CDK) activity to temporally separate two essential steps of DNA replication, origin licensing, and origin firing (Bell and Labib 2016). The low-CDK activity during G1 is permissive for origin licensing, the process of loading the Mcm2-7 helicase onto origins of replication in an inactive state. Three additional proteins catalyze Mcm2-7 loading; Cdc6, Cdt1, and the origin recognition complex (ORC) (Remus et al. 2009; Evrin et al. 2009). At the G1-S transition, CDK bound to S-phase cyclins (S-CDK) oppositely regulates origin firing and origin licensing. First, S-CDK triggers origin firing by phosphorylating two essential proteins, Sld2 and Sld3. These modifications are required for helicase activation and replisome assembly, the molecular events that mediate origin firing (Masumoto et al. 2002; Tanaka et al. 2007; Zegerman and Diffley 2007). Second, S-CDK and CDK bound to M-phase cyclins (M-CDK) prevent origin licensing during S, G2, and

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M-phases. To prevent this step of DNA replication, these kinases phosphorylate and inhibit multiple helicase-loading proteins including Cdc6, which, when phosphorylated, is targeted for proteolytic degradation (Drury et al. 1997; Calzada et al. 2000; Drury et al. 2000; Perkins et al. 2001; Arias and Walter 2007). By using CDK activity to separate origin licensing and origin firing, cells ensure that DNA replication happens exactly once per cell cycle.

During meiosis, the link between DNA replication and oscillations of CDK-activity presents a unique problem. CDK activity has been shown to decrease upon exit from MI and then increase upon entry into MII (Choi et al. 1991; Iwabuchi et al. 2000; Carlile and Amon 2008), a time when DNA replication must remain inhibited. This oscillation of CDK activity is required for MII chromosome segregation (Buonomo et al. 2003; Marston et al. 2003; Fox et al. 2017). However, an oscillation of CDK activity is also sufficient for replication of the entire genome in mitotic cells (Broek et al. 1991; Dahmann et al. 1995). Thus, there must be meiosis-specific mechanisms that prevent DNA replication between MI and MII while CDK-activity oscillates.

We recently demonstrated that meiotic cells inhibit both origin licensing and origin firing during the meiotic divisions, and that two critical targets of inhibition are Cdc6 and Sld2 (Phizicky et al. 2018). Both proteins are targeted for proteolytic degradation during meiosis, and CDC6 is also transcriptionally repressed. This co-inhibition of both proteins during meiosis is in contrast to the situation in mitotic cells wherein Cdc6 and Sld2 are inactivated during different stages of the cell cycle (Arias and Walter 2007; Bell and Labib 2016), and underscores the importance of preventing DNA replication during the meiotic divisions.

Consistent with the distinct requirements of meiosis and mitosis, meiotic cells activate genes required for meiosis and repress genes that promote mitotic cell division. In Saccharomyces cerevisiae, genome-wide studies have revealed that not only do meiotic and mitotic cells transcribe and translate different genes, but also that many genes expressed in both cell types make different RNA and protein isoforms (Chu et al. 1998; Lardenois et al. 2011; Brar et al. 2012; Zhou et al. 2017; Hollerer et al. 2017; Cheng et al. 2018). In particular, thousands of meiosis-specific short RNAs have been identified that are produced by a combination of antisense transcription of mitotically expressed open-reading frames (ORFs), intergenic transcription, and truncated RNA isoforms containing only part of an annotated ORF (Lardenois et al. 2011; Zhou et al. 2017; Hollerer et al. 2017). In addition to their specificity for meiosis, these short RNAs constitute >10% of the translational capacity of meiotic cells (Hollerer et al. 2017), and protein products have been observed in a limited number of cases (Zhou et al. 2017; Hollerer et al. 2017). However, the molecular mechanisms resulting in meiosis-specific expression of these short RNAs, as well as the function of their protein products, have not been elucidated.

Here, we show that repression of CDC6 and SLD2 mRNAs during meiosis is associated with production of heterogeneous short RNAs corresponding to the 3’ end of both genes. Although the short CDC6 RNA codes for a short protein (Cdc6\textsuperscript{short}), this protein is not essential for meiosis nor does it impact origin licensing. We found that production of Cdc6\textsuperscript{short} is dependent on one or more promoter(s) inside the ORF that are activated coincident with repression of P\textsubscript{CDC6}. Examination of CDC6 genes from related yeast species suggests that repression of full-length CDC6 mRNA and production of unique short CDC6 RNAs is an evolutionarily conserved feature of these genes. Together, these data show that full-length CDC6 and SLD2 are transcriptionally repressed during meiosis, and that the transcriptional inactivation of full-length CDC6 is associated with new sites of transcription initiation within the CDC6 ORF.

### Results

**Repression of CDC6 and SLD2 during meiosis coincides with expression of shortened isoforms**

To understand how meiotic cells repress expression of Cdc6 and Sld2 proteins during the meiotic divisions, we analyzed the protein and RNA expression of both genes using a previously described meiotic-synchronization system (Benjamin et al. 2003; Carlile and Amon 2008). In this system, meiotic cells undergo G1 and S-phases before accumulating at a G2-arrest. Release from this arrest allows cells to progress synchronously through both meiotic divisions (Fig. S1). While examining Cdc6 protein levels, we identified a short isoform of Cdc6 (Cdc6\textsuperscript{short}) that is 20 kDa smaller than the full-length protein and was expressed throughout the meiotic divisions (Fig. 1a, top). Cdc6\textsuperscript{short} protein expression coincided with expression of a shorter CDC6 mRNA isoform (Fig. 1a, bottom). Both the short RNA and short protein isoforms were anti-correlated with the corresponding full-length products. Cdc6\textsuperscript{short} protein was detected via a C-terminal tag indicating that Cdc6\textsuperscript{short} included the C-terminus of the ORF. Consistent with this conclusion, we detected the CDC6\textsuperscript{short} mRNA using a 3‘-end specific probe by northern blot (Fig. 1a). Neither the CDC6\textsuperscript{short} mRNA nor protein was detectable in asynchronous end fbot genes.

Similar to CDC6, we found that full-length SLD2 mRNA was reduced during the meiotic divisions, and that this decrease correlated with expression of a short SLD2 RNA species (Fig. 1b). We did not, however, detect a short Sld2 protein isoform (Fig. S1b). To determine which end of the full-length RNA was included in SLD2\textsuperscript{short}, we performed northern blots...
using probes specific for the 5′- and 3′-end of the gene. Although both probes detected full-length SLD2 mRNA, only the 3′-probe detected the SLD2short RNA (Fig. 1c). Like CDC6short, SLD2short RNA was not detectable in mitotically dividing cells (Fig. 1c). Together, these data show that both CDC6 and SLD2 produce a shortened RNA isoform during meiosis while the full-length mRNA of both genes is repressed.

Cdc6short is produced using an internal start codon and contains the C-terminal portion of CDC6

We focused our studies on CDC6short because the production of a short protein product in addition to the short mRNA suggested a potential function. To elucidate any potential functions of Cdc6short, we wanted to identify which portion
of Cdc6 was expressed in this isofrom. To determine the expressed region, we performed 5′ RACE on CDC6 mRNAs from different cell-cycle stages (Fig. 2a). Full-length CDC6 was detectable in asynchronous mitotic and pre-meiotic G1 cells, and the 5′ end of these products mapped primarily to the previously-defined start site of transcription (nucleotide −38, Fig. 2a) (Pelechano et al. 2013). cDNA from mitotic cells was present almost entirely as the full-length band, consistent with our northern blots (Figs. 1a and 2a). In contrast, during both meiosis I and II, multiple shorter versions of CDC6 cDNA were identified but none of the full-length product (Fig. 2a). The 5′ ends were located throughout the ORF, ranging from nucleotides +335 to +1005, and none of the clones contained splice-site junctions. The robust detection of full-length CDC6 cDNA from asynchronous mitotic cells shows that these shorter products are not due to RNA 5′ cDNA showed similar 5′-end heterogeneity during meiosis and a lack of splice-site junctions (Fig. S2a). Along with our northern blots, these data demonstrate that the CDC6short and SLD2short RNAAs have many different sub-species, but that all of these forms are missing part of the 5′ end.

Due to the large number of 5′ ends detected for CDC6short, it was unclear which AUG was used as the translation start codon for the Cdc6short protein. To address this question, we integrated a second copy of CDC6 at the LEU2 locus with mutations in potential translation start codons (ATG (M) → GTG (V)). Unlike the full-length protein which relied entirely on Met1, Cdc6short depended mostly on Met202, but also used Met225 at low efficiency (Fig. 2b). None of the other ATG codons tested prevented either full-length Cdc6 or Cdc6short protein expression (Figs. 2b, S2b). In combination with the 5′ RACE results, these data lead us to conclude that Cdc6short contains the Cdc6 sequence from Met202 to the C-terminus.

To determine whether Cdc6short is essential for meiosis, we replaced the endogenous copy of CDC6 with the Met-202/225-Val mutant. This mutant prevented expression of Cdc6short protein but not CDC6short mRNA (Fig. 2c). Eliminating Cdc6short protein did not prevent completion of meiosis I or meiosis II (Fig. 2c) nor did it cause decreased viability of the resulting spores (Fig. 2d). From these data, we conclude that Cdc6short protein is not essential for meiosis or spore formation.

Cdc6short cannot participate in or inhibit origin licensing

We hypothesized that Cdc6short could act as a dominant-negative protein isoform to inhibit origin licensing during the meiotic divisions. To determine whether Cdc6short impacted origin licensing, we purified Cdc6 and Cdc6short (Fig. 3a) and tested their function using a reconstituted Mmc2-7 loading assay (Remus et al. 2009; Evrin et al. 2009). In this assay, origin-containing DNA attached to a magnetic bead was incubated with the four proteins required for Mmc2-7 loading: Cdc6, Cdt1, ORC, and Mmc2-7. After this incubation, the DNA-beads were washed and the proteins that remained associated with DNA were detected (diagram in Fig. 3b). Although wild-type (WT) Cdc6 efficiently loaded Mmc2-7 complexes onto origin DNA, Cdc6short did not facilitate this event (Fig. 3c). To test whether Cdc6short acted as a dominant-negative, we pre-incubated Cdc6short with ORC, Mmc2-7, Cdt1, and origin-containing DNA before adding WT Cdc6. Even when present in 16-fold excess compared to the WT Cdc6, Cdc6short did not inhibit Mmc2-7 loading (Fig. 3d). To determine whether Cdc6short could form initial physical interactions with helicase-loading proteins, we repeated the same experiment in the presence of ATPγS, a slowly hydrolyzable ATP-analog (diagram in Fig. 3b). ATPγS stalls Mmc2-7 loading at an intermediate stage of assembly that contains one copy each of ORC, Cdt1, Cdc6, and Mmc2-7 bound to the origin DNA (the OCCM complex) (Randell et al. 2006; Sun et al. 2013; Ticau et al. 2015; Zhai et al. 2017; Ticau et al. 2017; Yuan et al. 2017). If Cdc6short could interact with these proteins, we would expect them to associate with DNA in a Cdc6short-dependent manner. Unlike WT Cdc6, Cdc6short...
Cdc6 short does not form the OCCM complex in ATPγS, suggesting that it cannot physically interact with at least one of these proteins (Fig. S3a). Finally, to test whether Cdc6 short acts as a dominant-negative inhibitor in vivo, we overexpressed this protein in mitotically dividing cells. Consistent with our in vitro data, Cdc6 short can catalyze Mcm2-7 loading, whereas Cdc6 short cannot. Titrations of Cdc6 short (lanes 5–7) are shown, and a reaction lacking Cdc6 (lane 1) shows that detection of Mcm2-7 complexes depends on the helicase-loading reaction. Cdc6 short does not have a dominant-negative effect on Mcm2-7 loading in vitro. Top: flowchart of experiment. Bottom: pre-incubation of Cdc6 short at the indicated concentration with the other helicase-loading proteins and origin DNA does not prevent WT Cdc6 from promoting Mcm2-7 loading (lanes 2–6). A reaction lacking Cdc6 (lane 1) shows that detection of Mcm2-7 complexes depends on the helicase-loading reaction. Overexpression of Cdc6 short-3V5 does not cause growth defects in mitotic cells. Strain containing P GAL-Cdc6 short-3V5 overexpresses Cdc6 short-3V5 on YP-Gal plates but not YPD plates. The control strain does not overexpress Cdc6 short-3V5 on either medium.

Production of Cdc6 short during meiosis does not require activity from the upstream CDC6 promoter

Given the apparent lack of function for Cdc6 short protein, we asked why cells produce CDC6 short mRNA during meiosis. Perhaps Cdc6 short expression is associated with repression of full-length CDC6 and is not important for making a protein product. We hypothesized two possible mechanisms that would cause the observed anti-correlation between full-length CDC6 and Cdc6 short levels (Fig. 1a): (1) the full-length mRNA is cleaved or partially degraded from the 5′-end to make CDC6 short mRNA, which is then translated by cap-independent mechanisms to produce Cdc6 short protein; (2) the upstream CDC6 promoter (P CDC6 ) is mutually exclusive with a cryptic promoter within the CDC6 ORF. A key distinction between these models is that the first model would require P CDC6 for Cdc6 short production but the second would not.

To address these two possibilities, we inserted a second copy of the CDC6 gene with multiple promoter alterations and examined the production of Cdc6 isoforms from this locus using the 3V5 tag (Fig. 4a). We found that shorter Cdc6 isoforms are produced under any condition during which P CDC6 is inactivated, supporting a model in which expression of the full-length and short CDC6 RNAs depends on mutually...
exclusive promoters (Fig. 4b–d). Deletion of nucleotides −550 to −1 strongly reduced $P_{CD6}$ activity resulting in dramatic reduction of full-length Cdc6 protein in asynchronous mitotic cells. Interestingly, these cells produced an intermediate length Cdc6 isoform ($Cdc6_{middle}$, discussed further below) (Fig. 4b). During meiosis, the promoter deletion did not affect
full-length Cdc6 production, but did increase levels of Cdc6 middle and Cdc6 short (Fig. 4c, panel II). It is noteworthy that the decreased production of full-length Cdc6 using this deletion in mitotic cells is similar to the amount produced by meiotic cells (Fig. 4c). These data suggest that meiotic cells produce less Cdc6 than mitotic cells due to reduced activity of P CDC6 during meiosis. The basal Cdc6 expression in the P cdc6 deletion construct is likely to be the result of a promoter in the LEU2-integrating vector that is juxtaposed with the CDC6 ORF upon deletion of P CDC6.

To eliminate any transcription elongation originating from within or before P CDC6, we made constructs with transcription terminators or similarly sized control sequences inserted between nucleotides +48 and +49 of the Cdc6 short ORF (Fig. 4a). These sequences were flanked by stop codons to prevent translation of full-length Cdc6, but these insertions do not affect the Cdc6 short ORF. We found that both the CYC1 and TEF terminator sequences were sufficient to induce Cdc6 short accumulation during meiosis.

**Evolutionarily-related CDC6 genes repress full-length CDC6 expression but produce unique short RNAs**

The strong repression of full-length CDC6 transcription during meiosis suggests that repressing CDC6 expression is critical to prevent origin licensing (Fig. 1a) (Phizicky et al. 2018). To determine whether the mechanism to repress full-length CDC6 expression is conserved in other organisms, we cloned the CDC6 gene from four yeast species (Saccharomyces bayanus, Saccharomyces kudriavzevii, Saccharomyces mikatae, and Saccharomyces paradoxus) into plasmids and tested for RNA and protein expression. These species are estimated to be between 5 and 20 million years diverged from S. cerevisiae (Kellis et al. 2003; Scannell et al. 2011). Each gene tested included the entire CDC6 ORF, ~550 nucleotides upstream of the coding region, and a 3V5 tag at the C-terminus for detection of the protein and RNA produced. The primary Cdc6 short start site (Met202) is conserved in all of these species.

We analyzed expression of CDC6 from each species during mitosis and meiosis. Each of the genes expressed full-length Cdc6 in mitotic cells at similar levels to S. cerevisiae (Fig. 5a). During meiosis, S. bayanus CDC6 showed complete repression of full-length CDC6 mRNA and expressed CDC6 short protein and mRNA to similar levels as the S. cerevisiae CDC6 gene (Fig. 5b, c). Interestingly, although the genes from both S. kudriavzevii and S. mikatae also repressed full-length CDC6 during MI and MII, only a small amount of Cdc6 short protein was detected (Fig. 5b, c). Instead, both of these genes showed meiosis-specific enrichments of much shorter RNA isoforms than those observed from S. cerevisiae. In contrast to the other CDC6 genes, S. paradoxus CDC6 did not produce detectable Cdc6 short protein and had only limited enrichment of short RNA isoforms during meiosis, despite it having decreased expression of full-length CDC6 during MI (Fig. 5b, c). Together, these data suggest that transcriptional repression of full-length CDC6 during meiosis is associated with aberrant transcription initiation from within the CDC6 ORF, and that these features are evolutionarily conserved in related yeast species.

**Discussion**

Preventing DNA replication between MI and MII is critical for reducing ploidy during meiosis, but the mechanisms that cells use to repress specific steps of replication initiation have not been fully elucidated. In this study, we found that two essential DNA replication genes, CDC6 and SLD2, are transcriptionally repressed during the meiotic divisions. For each gene, the full-length mRNA is lost, expression of shorter RNAs that display a high degree of 5'-end heterogeneity increases. The CDC6 short mRNA encodes a short form of Cdc6, but this protein does not have an obvious function during origin licensing. We provide evidence that the short RNAs are from internal promoters rather than processing of the full-length mRNA. Finally, we demonstrate that meiotic repression of full-length CDC6 and production of shorter RNA isoforms is evolutionarily conserved in genes from related yeast species. These findings provide additional insight into how DNA replication is inhibited during MI and MII and suggest that production of CDC6 short is a byproduct of intragenic transcription upon repression of the full-length mRNA. In addition, our findings suggest that a subset of novel meiotic RNA and protein isoforms are a byproduct of inhibition of the associated full-length genes.

How is the CDC6 short mRNA produced? The accumulation of CDC6 short mRNA upon loss of the full-length mRNA (and vice versa) suggests that activation of one or more promoters inside the ORF is mutually exclusive with activation of the upstream promoter, P CDC6. Our finding that P CDC6 inactivation results in mitotic cells producing shorter Cdc6 isoforms (Fig. 4) suggests that transcription from P CDC6 represses the internal promoter(s). Consistent with this hypothesis, transcription elongation results in repressive chromatin marks being placed within gene bodies, and the lack of these marks allows transcription initiation from within ORFs (Carrozza
et al. 2005; Kouzarides 2007; Wagner and Carpenter 2012; Smolle et al. 2013; Nerie et al. 2017). Further support for this model stems from the observation that the distribution of CDC6short 5′ ends spans > 600 nucleotides (Fig. 2a). The elimination of repressive marks within a transcribed region would be consistent with the activation of multiple promoters within the CDC6 ORF. Although less likely, it is also possible that activation of the internal promoter(s) causes repression of PCDC6, thereby limiting full-length CDC6 expression. Our results are not consistent with CDC6short being produced as a result of partial decay or processing of full-length CDC6 mRNA. These models would predict that Cdc6 short would depend on PCDC6 activity, which it does not (Fig. 4). In addition, it is unlikely that partially degraded transcripts would have the necessary requirements (e.g., a 5′ cap) for translational initiation that the short transcripts direct. Finally, we detected no evidence of splice junctions in our analysis of the CDC6short RNAs.

Despite observing many 5′ ends for CDC6short mRNA, Met202 is by far the most highly used translation start codon during meiosis in unperturbed meiotic cells (Fig. 2). There are multiple reasons for this preference. First, we note that protein products produced from out-of-frame start codons would not be detectable by our C-terminal tag. Second, we detected very few RNAs with 5′-ends that are far enough upstream to allow translation initiation from either of the two upstream, in-frame start codons, Met89 and Met120 (Fig. 2a). Third, the sequence context of Met202 is more favorable for translation initiation than that of nearby, downstream AUGs due to its enrichment of upstream adenosines (Hinnebusch 2011; Dvir et al. 2013). Of the eleven nucleotides upstream of each start codon, Met202 has nine adenosines whereas Met225, Met261, and Met267 have two, two, and four adenosines, respectively. Yeast 5′-UTRs that are adenosine-rich, especially in the −1 to −10 region relative to the start codon, promote efficient translation initiation, likely by keeping the 5′-UTR...
unstructured and thus accessible to the ribosome (Hinnebusch 2011; Dvir et al. 2013). Thus, it is likely a combination of transcriptional and translational control that selects Met202 as the start codon for Cdc6\textsuperscript{short}.

Although our data are consistent with P\text{CDC6} and an internal promoter(s) being mutually-exclusive, this model may not be predictive under all conditions. Full-length CDC6 is transcriptionally repressed in mitotic cells at least from late S-phase until the end of M-phase (Zwerschke et al. 1994; Piatti et al. 1995). A simple mutually exclusive promoter model would predict that the shorter CDC6 mRNA isoforms would be observed at these stages of mitosis, yet we did not detect short CDC6 RNA or protein isoforms in asynchronous mitotic cells (Fig. 1a). Short CDC6 isoforms may not have been detected in mitotic cells either because P\text{CDC6} is not as fully repressed as it is in meiotic cells, or because P\text{CDC6} must be shut off for a longer time to allow chromatin changes that promote intragenic transcription initiation. Considering that Cdc6\textsuperscript{short} protein is observed in mitotic cells containing transcription terminators after P\text{CDC6} (Fig. 4b), mitotic cells most likely have all the factors required for internal initiation.

There are likely to be other transcriptional or translational regulatory mechanisms that affect which Cdc6 protein isoforms are produced. The detection of a distinct isoform, Cdc6\textsuperscript{middle}, from constructs either lacking P\text{CDC6} or those containing internal transcription terminators suggests that different Cdc6 protein isoforms are expressed as a function of the mechanism preventing transcription elongation into the ORF. These methods of inhibiting transcription elongation into the CDC6 ORF are different than the regulated transcriptional shutoff of full-length CDC6 observed during meiosis, during which Cdc6\textsuperscript{middle} is not detected above background levels (Figs. 1a and 4c). It is possible that different methods of inactivating P\text{CDC6} cause distinct chromatin changes, thereby permitting transcription initiation from different parts of the ORF. There is also clear evidence of cell-cycle-dependent regulation of Cdc6 isoforms. All of our constructs, either with WT CDC6 or the promoter perturbations, produce different ratios of Cdc6:Cdc6\textsuperscript{middle}:Cdc6\textsuperscript{short} in mitotic and meiotic cells (Fig. 4).

Determining the mechanisms that direct expression of meiosis-specific RNAs will be important for assessing their function, and should allow perturbation of many RNAs at the same time. We note that neither CDC6\textsuperscript{short} nor SLD2\textsuperscript{short} depended on the meiotic transcription factor Ndt80, as Ndt80 is not expressed from 0 to 6 hours during which both short RNAs are detectable (Fig. 1). The lack of dependence on Ndt80 for CDC6\textsuperscript{short} and SLD2\textsuperscript{short} is in contrast to many other meiosis-specific RNA isoforms, suggesting that different mechanisms are used to produce meiosis-specific transcripts (Zhou et al. 2017; Cheng et al. 2018). Finally, it is also clear that the transcriptional regulation of CDC6 and SLD2 activity is but one of many ways that cells ensure that these proteins are eliminated at the appropriate times in the cell cycle (Drury et al. 2000; Masamoto et al. 2002; Zegerman and Diffley 2007; Reusswig et al. 2016; Phizicky et al. 2018).

**Methods**

**Yeast strains and plasmids**

All yeast strains and their genotypes are described in Table S1. All strains used for in vivo experiments were derived from the SK1 background with the following genotype: ho::LYS2/ho::LYS2, lys2/lys2, ura3/ura3, leu2::hisG/leu2::hisG, his3:hisG/his3:hisG, trp1::hisG/trp1::hisG. CDC6 genes from *S. bayanus*, *S. kudriavzevii*, *S. mikatae*, and *S. paradoxus* were cloned out of the isolates given in the strain list. Plasmids used for protein expression are described in Table S2.

**Meiotic time courses**

All strains put through meiotic time courses contained P\text{GPD–GAL4–ER}, P\text{GAL–NDT80} for controlled block-release from meiotic G2 phase. Meiotic time courses were done as described in (Berchowitz et al. 2013). Briefly, cells were diluted to an OD\textsubscript{600}=0.25 in BYTA medium from saturated YPD cultures. BYTA cultures were grown for 18–20 h while shaking at 30 °C. After growth in BYTA, cells were washed with water and resuspended to OD\textsubscript{600}=1.9 in sporulation medium. The 0-hour sample was taken immediately after transfer to sporulation medium. Cultures were shaken at 30 °C for 6 h to allow cells to accumulate at the NDT80 block. To release cells from the block, 1-mM β-estradiol (5-mM stock in ethanol [Sigma, E2758]) was added to the culture medium. Protein, RNA, and immunofluorescence samples were harvested in parallel at the indicated time points.

**Immunofluorescence**

Tubulin immunofluorescence was performed as previously described (Berchowitz et al. 2013). One hundred cells were counted per time point for both spindle and nuclei scoring. Metaphase I and metaphase II cells were defined as having, respectively, one or two short, thick, bipolar spindle(s); anaphase I and anaphase II cells as having, respectively, one or two long, bipolar spindle(s); nuclei were counted as the number of distinct DNA masses.

**Immunoblots and RNA isolation**

Immunoblots and RNA isolation were done as previously described (Phizicky et al. 2018).
Northern blots

Equal amounts of total RNA (between 12 and 16 μg) were loaded in each lane of a formaldehyde gel and transferred to a nylon membrane (GE HyBond N+, catalog number RPN3050B). rRNA was detected by methylene blue and used as a loading control, and specific RNAs were detected using gene-specific probes. $^{32}$P-labeled probes were made by Klenow extension (GE Healthcare, RPN1605) of gene-specific PCR products containing the indicated regions. Primers for Klenow extensions were random hexamers. Templates: The CDC6 template was specific to the 3’ half of the ORF (nucleotides +923 to +1488). The three SLD2 templates used covered either the entire ORF (nucleotides +1 to +1359), or were specific to the 5’ half (nucleotides +1 to +547) or the 3’ half (nucleotides +805 to +1349) of the ORF.

5′ race

cDNA of CDC6 or SLD2 was made from 4 μg of total RNA using Superscript IV Reverse Transcriptase (Life Technologies, catalog number 18090010) and gene-specific primers according to manufacturer’s instructions. After cDNA synthesis and purification, a poly-deoxycytosine tail was added to the 3′ end of the cDNA using Terminal Transferase with a 19:1 ratio of dCTP:ddCTP (to limit the length of the tail). Poly-dC-tailed cDNA was then amplified by PCR using a poly-G primer and the same gene-specific primer as was used for the RT-step. A second PCR was performed using the resulting PCR product as a template, and a different gene-specific primer was used to increase the specificity of the product. This second-round PCR product was digested with SacI and SalI restriction enzymes, sites for which were encoded on the primers in the PCR reaction. The resulting products were run on an agarose gel (shown). Sections of this gel were purified and ligated into a similarly digested vector to prevent ligation reaction bias for short products. Ligation reactions were transformed and DNA from individual colonies was miniprepped. The entire ligated region was sequenced, and 5′-ends were called based on the last nucleotide that was successfully aligned before the poly-Cytosine region was reached.

Helicase-loading assays

Helicase-loading assays were done as described in (Kang et al. 2014). Briefly, 50-nM ORC, 150-nM Mcm2–7/Cdt1, and the indicated concentration of Cdc6 or Cdc6$^{\text{short}}$ were combined in a 40-μL reaction with 25-nM bead-bound 1.3-kB DNA containing the ARS1 origin. After combining the proteins and DNA, reactions were incubated at 25 °C for 30 min while shaking at 1250 rpm (Thermo-mixer, Eppendorf). Beads were then washed three times in buffer. For helicase-loading reactions, the three washes had buffer containing 300-mM KGlut, 500-mM NaCl, and 300-mM KGlut respectively. Proteins were eluted from DNA with DNase, run on SDS–PAGE gels, and detected with Krypton fluorescent stain (Fisher, PI-46629). To assay OCCM formation, ATP was replaced with 5-mM ATP$^\gamma$S, and beads were washed three times with buffer containing 300-mM KGlut.

Protein purifications

Cdc6 (plasmid pDP69; MBP-FLAG-Cdc6-6xHIS) and Cdc6$^{\text{short}}$ (pDP70; MBP-FLAG-Cdc6$^{\text{short}}$-6xHIS), containing CDC6 amino acids 202-513 were purified from Escherichia coli Rosetta 2 cells grown to log phase and induced overnight with 0.5-mM IPTG and shake at 18 °C for 20 h. Cells were resuspended in Buffer G (50-mM potassium phosphate [pH = 7.6], 5-mM magnesium chloride, 1% triton, and 1-mM DTT) with 400-mM potassium acetate (KAc), 2-mM ATP, 1-mM PMSE, and 1× Complete Protease Inhibitors (Roche), and then flash frozen in liquid nitrogen. Cells were thawed and lysed with lysozyme and sonication. After centrifugation at 12,000 rpm, 10-mM imidazole was added to the supernatant which was flowed over Nickel resin equilibrated in the same buffer. The resin was washed with 20 bed volumes of buffer. Protein was eluted using Buffer G, 400-mM KAc, 2-mM ATP, and 350-mM imidazole. Peak fractions were pooled and incubated with equilibrated amylose resin, which was then washed with 20 bed volumes of Buffer G with 400-mM KAc. Resin was then washed with five bed volumes of Buffer G, 400-mM KAc, and 15% glycerol, and protein was eluted using the same buffer with the addition of 10-mM amyllose. Peak fractions were pooled, aliquoted, flash frozen, and stored at −80 °C. ORC (ySDORC) and Mcm2–7–Cdt1 (yST144) were purified as previously described (Kang et al. 2014).

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