

Title (107 chars): Cell cycle Start is coupled to entry into the yeast metabolic cycle across diverse strains and growth rates

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Running header (39 chars): Cell division and yeast metabolic cycle

Abbreviations: YMC, yeast metabolic cycle; CDC, cell division cycle; HOC, high oxygen consumption; LOC, low oxygen consumption

Abstract: (173 words)

Cells have evolved oscillators with different frequencies to coordinate periodic processes. Here, we studied the interaction of two oscillators, the cell division cycle (CDC) and yeast metabolic cycle (YMC), in budding yeast. Previous work suggested that the CDC and YMC interact to separate high oxygen consumption from DNA replication to prevent genetic damage. To test this hypothesis, we grew diverse strains in chemostat and measured DNA replication and oxygen consumption with high temporal resolution at different growth rates. Our data showed that high oxygen consumption is not strictly separated from DNA replication; rather, cell cycle Start is coupled with the initiation of high oxygen consumption and catabolism of storage carbohydrates. The logic of this YMC-CDC coupling may be to ensure that DNA replication and cell division occur only when sufficient cellular energy reserves have accumulated. Our results also uncovered a quantitative relationship between CDC period and YMC period across different strains. More generally, our approach shows how studies in genetically diverse strains efficiently identify robust phenotypes and steer the experimentalist away from strain-specific idiosyncrasies.

Highlight Summary: (309 characters)

The authors study the interaction of two oscillators (cell division cycle and yeast metabolic cycle) with different frequencies. They show that cell cycle Start is coupled with the initiation of high oxygen consumption and breakdown of storage carbohydrates across diverse strains and different growth rates.

INTRODUCTION

Biological oscillators such as the cell cycle, circadian clocks, and metabolic rhythms are ubiquitous across the domains of life. The cell division cycle has a period that depends on nutrients and temperature, and the cell cycle regulatory network is strongly conserved across eukaryotes (Morgan, 2007). On the other hand, circadian clocks have a fixed ~24-hour period and are robust to changes to temperature; yet, their regulatory networks have evolved independently in cyanobacteria, plants, and animals (Rosbash, 2009). These different biochemical oscillators co-exist in the same cellular environment, sharing and competing for resources. It is well established that even weak coupling between oscillators can lead to their synchronization (Pikovsky *et al.*, 2001). Thus, how do biochemical oscillators with different frequencies co-exist in the same cell? Are there mechanisms and regulatory principles that ensure functional harmony between oscillators?

Quantitative analysis of circadian clocks and the cell division cycle in cyanobacteria (Mori *et al.*, 1996; Mori and Johnson, 2001), algae (Moulager *et al.*, 2010; Miyagishima *et al.*, 2014), fungi (Hong *et al.*, 2014), and mammals (Nagoshi *et al.*, 2004; Bieler *et al.*, 2014; Feillet *et al.*, 2014) have shown that the circadian clock gates the cell division cycle, such that cell cycle events are halted at checkpoints during prohibitive circadian phases (Johnson, 2010). These oscillators appear to co-exist through asymmetric coupling, whereby a slower, rigid circadian clock dominantly regulates a faster, flexible cell cycle. A related phenomenon occurs in the budding yeast, *Saccharomyces cerevisiae*, where a well-studied metabolic cycle interacts with the cell cycle. In contrast to circadian clocks, the yeast metabolic cycle is shorter than 24 hours and shows a range of periods that depend both on the strain examined and growth conditions used.

The yeast metabolic cycle (YMC) was first observed as a synchronous oscillation in the pH and dissolved oxygen levels by a population of budding yeast in low-glucose chemostat conditions (Finn and Wilson, 1954; Kaspar von Meyenburg, 1969). The population oscillated between a low oxygen consumption (LOC) phase and a high oxygen consumption (HOC) phase. During LOC, cells build and accumulate storage carbohydrates, such as trehalose and glycogen (Küenzi and Fiechter, 1969). Upon reaching some threshold, a fraction of cells commit to HOC and catabolize their

storage carbohydrates to produce a pulse of energy for cell biosynthesis through aerobic fermentation (Futcher, 2006). These metabolically-committed yeasts produce and secrete two-carbon products of fermentation (e.g. ethanol, acetaldehyde) and sulphur assimilation (e.g. dihydrogen sulphide) (Sohn and Kuriyama, 2001; Wolf *et al.*, 2001), all of which can phase-shift the YMC of other cells. The population synchrony of the YMC in a chemostat is thought to arise from YMC-to-YMC coupling across cells via secreted metabolites (Robertson *et al.*, 2008).

However, within each cell, the YMC also interacts with the cell division cycle (CDC). The CDC and YMC periods range from 3 - 15 hours and 0.4 - 15 hours respectively in the literature, varying as a function of dilution rate which sets the population growth rate in a chemostat; see Table S1. The CDC and YMC periods do not appear to have a fixed relationship and, in fact, the ratio may change continuously with the dilution rate (Slavov and Botstein, 2011). Despite the seeming independence of these two oscillators, cell cycle events such as DNA replication occur once per YMC (Klevecz *et al.*, 2004; Tu *et al.*, 2005; Chen *et al.*, 2007; Slavov and Botstein, 2011). The flexible coupling of these two oscillators resembles the gating observed between the circadian clock and cell division cycle. Based on their work with the distiller's strain IFO 0233, Klevecz and colleagues first suggested that the observed gating between YMC and CDC might be functionally important for the temporal separation of CDC events (i.e. DNA replication) that are incompatible with YMC events (i.e. HOC and aerobic respiration, which could lead to oxidative damage of DNA) (Klevecz *et al.*, 2004).

This was confirmed by subsequent work with a different laboratory strain CEN.PK, which showed that DNA replication occurred towards the end of HOC as pO_2 levels rise (Tu *et al.*, 2005; Chen *et al.*, 2007). The authors classified the end of HOC (where yeast oxygen consumption is still high) as a 'reductive-building' phase, based on clustering of gene expression microarrays. However, later work with an S288C-derived laboratory strain (DBY12007) demonstrated that DNA replication could occur either in the middle of HOC or at the HOC/LOC boundary, depending on the underlying dilution rate (Slavov and Botstein, 2011). Thus, strict separation of DNA replication and HOC does not appear to be a consistent feature of the YMC. These discrepancies have led to a debate on the fundamental frequency of the ultradian YMC, the definition of relevant metabolic phases, and the nature of YMC-CDC

coupling (Lloyd, 2006; Murray, 2006; Tu *et al.*, 2006; Slavov and Botstein, 2011). This raises the important question: Which YMC and CDC coupling phenotypes (e.g. gating and timing of CDC and YMC events, ratio of CDC and YMC periods at different dilution rates) are universally shared across *all* budding yeasts, and which might be strain-specific idiosyncrasies?

Growth medium and chemostat settings differed between previous YMC experiments, which confounds interpretations of YMC phenotype because genotype (i.e. strains) and environment (i.e. chemostat settings) were uncontrolled; see Table S1.

Moreover, recent work on metabolic variation between a lab strain and a wild strain discovered genetic differences that affect sulfur assimilation, glucose uptake and ethanol secretion (Breunig *et al.*, 2014). These metabolic pathways are relevant to the YMC, which suggests that there could be profound differences between lab strains and wild isolates. To help untangle effects of genotype from those of environment, we first measured YMC-CDC coupling in lab strains CEN.PK, DBY12007 and two wild strains YJM128 (a clinical lung isolate) and YPS670 (an isolate from oak sap). These experiments were done in a chemostat environment with identical growth medium, oxygen flow, and pH. Growth rate is known to affect the timing of DNA replication with respect to HOC (Slavov and Botstein, 2011). Thus, we also measured YMC-CDC coupling in these different strains over a range of dilution rates.

RESULTS

Quantitative measurement and analysis of strain-specific and universal YMC phenotypes

All strains were first grown to saturation in batch growth and starved for at least six hours (Figure 1A). We then activated the chemostat feed pumps and added fresh medium at a constant dilution rate (D). The chemostat dilution rate sets the pace of glucose delivery and, hence, sets the average yeast growth rate and cell division time. Yeasts grew more slowly at lower dilution rates and exhibited a longer population doubling time. After an interval of transient arrhythmic behavior, the dissolved oxygen levels (pO_2) settled into a steady-state oscillation between HOC and LOC.

We confirmed that the measured pO_2 profile accurately tracks the steady state oxygen level and is determined by the yeast oxygen uptake rate; see Methods and Figure S1. Our measurements of gas transfer rates and pO_2 probe behavior indicated that the pO_2 measured in any given chemostat chamber reaches steady state after an abrupt change in oxygen uptake rate within 1-2 minutes. This is much shorter than the ~ 15 -30 minutes during which pO_2 moves from low to high values between the high oxygen consumption and low oxygen consumption phases of the YMC. Thus, the slow increase in pO_2 after a local minimum in pO_2 ('reductive-building' phase (Tu *et al.*, 2005)) corresponds to a high yeast oxygen uptake rate where the rate of oxygen consumption is slowly decreasing. Similarly, the slow drawdown of pO_2 after a local maximum in pO_2 ('oxidative phase' (Tu *et al.*, 2005)) is due to a steady increase in the yeast oxygen uptake rate rather than the slow depletion of dissolved oxygen after a rapid increase in yeast respiration. Our controls show that we can classify oxygen consumption *rate* (HOC or LOC) based on pO_2 levels (low or high). We empirically defined the boundaries of HOC and LOC at 65% of local peak-to-trough pO_2 levels because this threshold was (i) robust to slow drifts in pO_2 probe calibration, (ii) independent of magnitude of pO_2 , and (iii) entry into HOC was co-incident with the onset of aerobic fermentation and acidification of the growth medium; see Figure S1.

All tested strains were capable of synchronous metabolic cycles, including wild isolates from widely varying ecological contexts (Figure 1B). Thus, the YMC appears to be robust to any genetic variation in the sulfur assimilation, glucose uptake and ethanol secretion pathways across our strains. However, this variation does affect the quantitative features of the YMC. For example, each strain had a different pO_2 profile and YMC period ($\tau_{ymc} = 110 - 200$ minutes) at identical dilution rates. How does the pO_2 profile and τ_{ymc} of each strain change with growth rate?

To address this question, we first measured τ_{ymc} as a function of the dilution rate for CEN.PK (Figure 2). As glucose became more limiting at lower dilution rates, the cells exhibited a longer YMC and CDC. Strikingly, all of the increase in τ_{ymc} occurred in the LOC interval (τ_{loc}), whereas the HOC interval (τ_{hoc}) stayed constant or actually decreased with increasing cell cycle period (τ_{cdc}). To better quantify this relationship, we plotted τ_{ymc} , τ_{hoc} , and τ_{loc} as a function of τ_{cdc} across all strains (Figure 3A). These cross-strain data confirm that all of the increase in τ_{ymc} occurred in the LOC interval

(τ_{loc}). The LOC and HOC phases are associated with the anabolism and catabolism of storage carbohydrates, respectively. This suggests a simple model of how the YMC and CDC are related to metabolism: At lower dilution rates, cells spend more time in LOC accumulating storage carbohydrates because of the smaller glucose flux. Once these cells reach a metabolic threshold, they switch to HOC and catabolize their storage carbohydrates to provide biomass and energy for cell growth and division. On the other hand, cell division and the subsequent switch back to LOC occur after a similar amount of time regardless of the dilution rate.

Recent observations in the common lab strain background S288C showed that τ_{ymc} increases linearly with τ_{cdc} (Slavov and Botstein, 2011). To what extent does this quantitative relationship hold across multiple strains? Our analysis confirms that τ_{ymc} initially increases linearly, but that it slowly approaches a strain-specific maximum period at large τ_{cdc} (Figure 3A). This hyperbolic relationship was consistent across all strains, although each strain had a unique set of parameters (see Methods and Table S2). Growth rate is often the relevant variable in microbial physiology. Thus, we re-plotted our quantitative data in units of frequency, $f_{ymc} = 1/\tau_{ymc}$ and $f_{cdc} = 1/\tau_{cdc}$, because f_{cdc} is proportional to the average growth rate set by the dilution rate (D). The same data for each strain is now linear in frequency space with a unique slope and y-intercept (Figure 3B).

Extrapolation of the quantitative relationships in Figure 3A predicts a maximum growth rate at which yeasts spend all of their time in HOC, and where the YMC should disappear (i.e. vertical arrows, where $\tau_{loc}=0$). This maximum growth rate is strikingly close to the critical dilution rate D_c at which yeast transitions from pure oxidative metabolism to aerobic fermentation (Figure 3C). This critical point is also where genes involved in mitochondrial respiration and catabolism of gluconeogenic carbon sources are repressed by glucose, i.e. the Crabtree effect (Cortassa and Aon, 1998). This suggests that fully respiratory carbon metabolism and the absence of glucose repression may play a role in the emergence of synchronous YMC in chemostat conditions.

How do the frequencies of the YMC and CDC oscillators vary relative to one another as a function of growth rate? The ratio of frequencies of simple coupled oscillators might be expected to hop discontinuously from one simple rational number to

another (i.e. "Devil's staircase"), as the frequency of one oscillator changes relative to the other (Pikovsky *et al.*, 2001). Our data suggest that the ratio of YMC and CDC frequency (f_{ymc}/f_{cdc}) stays constant in wild strains, but varies continuously in laboratory strains as growth rate is adjusted (Figure 3B). The absence of discontinuous hopping indicates that the YMC and CDC oscillators across the population are not mode-locked in a simple and deterministic manner.

Cell division cycle and entry into HOC are tightly linked across different strains and growth rates

Budding yeast irreversibly commit to a cell division cycle after passing a commitment threshold known as Start (Hartwell *et al.*, 1974; Johnston *et al.*, 1977). This triggers the expression of hundreds of G1/S genes that are then followed by DNA replication. To quantify the timing of CDC relative to the YMC, we measured DNA replication of our different strains at high temporal resolution; see Figure 4. In agreement with previous studies, our cross-strain results showed that CDC events are gated such that all strains exhibit a single pulse of DNA replication per YMC. Moreover, only a fraction of cells commit to the CDC each YMC, i.e. 'one-to-some' coupling. This coupling suggests a complicated sub-population structure in the chemostat, where staggered cohorts enter the CDC each YMC. It also explains how the YMC and CDC appear to be one-to-one coupled in bulk assays, yet can have different periods.

In contrast to one-to-some coupling, the timing of DNA replication relative to HOC showed strain variability (Figure 4). For example, CEN.PK replicated its DNA in the middle of HOC, whereas DBY12007 replicated its DNA near entry into LOC. These cross-strain data reinforce that the HOC phase is not strictly separated from DNA replication (Slavov and Botstein, 2011). If not strict separation, might there be other timing relationships that are universal or invariant across strains?

To address this question, we first analyzed DNA replication in a single strain (CEN.PK) at different dilution rates; see Figure 5. The YMC and CDC remained coupled one-to-some as we changed the dilution rate. We noticed that entry into HOC and onset of DNA replication was separated by a mostly invariant strain-specific time interval despite the longer periods of YMC, LOC, and CDC at lower dilution rates. Analysis of the separation between entry into HOC and DNA replication (Δ_s)

as a function of dilution rate across multiple chemostat runs supports a mostly invariant Δ_S model; see Table 1. Strikingly, this invariant Δ_S relationship extended to the other strains DBY12007 and YPS670 (see Figures S4-S5 and Table 1), although each value of Δ_S was strain-specific.

DISCUSSION

Our aim was to determine which behaviors of the YMC are universal. By studying multiple strains over a wide range of shared growth conditions, a number of key conclusions can be reached regarding the robustness of YMC-CDC phenotypes. First, all strains, including wild isolates, exhibited synchronous metabolic oscillations. Second, the increase in τ_{ymc} across all strains resulted from an increase in the length of LOC, whereas the length of HOC remained approximately constant or actually decreased (Figures 2-3). Third, all strains exhibited a single pulse of DNA replication per YMC where only a fraction of cells commit to division each YMC, i.e. one-to-some coupling (Figure 4). Last, all committed cells entered the cell cycle and replicated their DNA within a strain-specific interval (Δ_S) regardless of the extended length of YMC, LOC, and CDC at different growth rates (Figure 5 and Table I).

Population synchrony and structure of YMC-CDC coupling in a chemostat

The universal features of YMC/CDC coupling observed in this work point to a simplified model of YMC-CDC coupling (Figure 6) where cell cycle Start is intimately coupled to the catabolism of storage carbohydrates, as summarized in (Futcher, 2006). This connection between Start and metabolism is consistent with the known cell cycle literature, where smaller yeasts (i.e. daughter cells) spend more time in G1 accumulating biomass until they have reached a critical size and can commit to the CDC (Hartwell *et al.*, 1974). As the flux of glucose decreases at lower chemostat dilution rates, the yeasts spend more time in LOC and presumably G1 accumulating sufficient biomass and storage carbohydrates. This would explain the increase in YMC, LOC, and CDC periods at lower dilution rates (Figures 2-3). When a cell has accumulated enough biomass and storage carbohydrates, it is able to enter HOC and consume its storage carbs.

One-to-some coupling suggests that the population is likely composed of staggered cohorts, which may differ in their degree of storage carbohydrate and/or biomass

accumulation. Staggered cohorts could arise from and be maintained by the cell size asymmetry between mother and daughter cells (Hartwell and Unger, 1977) and/or secretion of different metabolites that advance or block different phases of the YMC and/or CDC (Young *et al.*, 2012). Each YMC, a fraction of cells (usually less than a third) pass Start, enter the cell division cycle, and replicate their DNA on schedule, i.e. one-to-some coupling (Figure 4). The temporal coordination of YMC events (i.e. burning storage carbohydrates and entering HOC) and CDC events (i.e. Start) would explain the mostly invariant Δ_S across dilution rates. A biological rationale for coordinating cell cycle Start and carbohydrate catabolism in this cohort of committed cells may be to ensure that sufficient energy is available to complete biomass accumulation, DNA replication, and cell division in a timely manner. More work will be required to elucidate the complex population structure of the YMC and the nature of the cohorts initiating cell division each YMC. For example, what fraction of mother versus daughter cells (each having different CDC periods at the same population doubling time) initiate cell division each YMC? Do all cells exhibit metabolic oscillations each YMC or is it only the cohort that enters the cell division cycle?

Strain-specific features

There were several notable YMC differences between strains. First, every strain had a distinct pO_2 waveform and τ_{YMC} when grown at the same dilution rate (see Figure 1). Second, the exact quantitative relationship between τ_{YMC} , T_{LOC} , T_{HOC} as a function of τ_{cdc} varied across strains (Figure 3 and Table S2). Third, the characteristic delay between entry into HOC and initiation of DNA replication (Δ_S) varied between strains (Table I) with wild strains having shorter delays than laboratory strains. Last, the overlap between HOC and DNA replication depended on the strain and the dilution rate (Figure 4 and Figure S6). Our results do not support the hypothesis that DNA replication is strictly segregated from aerobic respiration (Klevecz *et al.*, 2004; Tu *et al.*, 2005; Chen *et al.*, 2007). Strain-to-strain variation in the length of HOC at a given growth rate can lead to DNA replication occurring in early HOC, late HOC, or early LOC (Figure 4). Changes in the length of HOC at different dilution rates can also lead to DNA replication occurring in HOC or LOC (Figure S6).

Despite these strain differences, an unexpected regularity emerged when we examined the fraction of time spent in HOC (ϕ_{HOC}) as a function of growth rate (Figure S7). ϕ_{HOC} is as an approximately linear function of growth rate, such that all

strains spent similar fractions of time in HOC at identical growth rates. This is reminiscent of bacterial growth laws in which the RNA/protein ratio increases linearly with growth rate, as bacteria grow faster and put more energy and biomass into growth and ribosome biosynthesis (Schaechter *et al.*, 1958; Scott *et al.*, 2010). Bacteria achieve this by smoothly ramping gene expression and metabolism in response to changes in growth rate. The yeast metabolic cycle, in contrast, is associated with an oscillation in metabolic rate and expression levels of large fractions of the transcriptome. The early HOC phase (identified elsewhere as the 'oxidative phase') in particular is linked to a burst of expression of ribosome biosynthesis, amino acid biosynthesis, and RNA metabolism genes (Klevecz *et al.*, 2004; Tu *et al.*, 2005; Murray *et al.*, 2007). Thus, yeast appears to partition ribosome biosynthesis and carbon metabolism in time, using frequency modulation to smoothly tune the ratio of 'productive' periods to 'quiescent' periods with the growth rate.

Does the YMC occur outside of particular chemostat conditions?

Historically, the YMC has been studied in low-glucose chemostats because the budding yeast population self-synchronizes in these conditions. The synchrony is thought to arise from signaling between cells. The dispersal of secreted metabolites in a well-mixed bioreactor may help synchronize the oscillation across the population, allowing study at a bulk population level. However, mixing alone is insufficient to ensure synchrony because the synchronous YMC disappears at different chemostat dilution rates, oxygen flow, and nutrient conditions in different strains. Loss of population synchrony could arise because: (i) single cells no longer have an autonomous YMC oscillator, or (ii) YMC oscillators in cells have lost their ability to entrain to other YMC oscillators. One could distinguish between these two cases by looking for evidence of the YMC in conditions where population synchrony is lost.

Recent work suggests that the YMC may indeed occur in single cells in asynchronous populations. First, expression of genes known to oscillate across the YMC and CDC were measured in single cells using single-molecule RNA FISH (Silverman *et al.*, 2010). These yeast were grown in chemostat in nutrient-starved conditions, which did not exhibit synchronous YMC. The authors showed that different pairs of YMC and CDC transcripts, previously correlated and anti-correlated in populations undergoing

synchronous YMC oscillations in chemostat, continued to be correlated and anti-correlated in single cells from an asynchronous chemostat. Second, the degree of stress resistance exhibited by yeasts was unimodal in a synchronized population with all cells sensitive to heat shock during HOC and more resistant during LOC. However, stress resistance was bimodal in yeasts grown under similar but asynchronous conditions. The authors suggested that there was an asynchronous, biphasic mixture of cells in HOC and LOC – a large population dies rapidly, followed by a second population dying slowly during heat shock (Slavov et al., 2012).

Yeasts can also exhibit synchronous metabolic oscillations outside of chemostat conditions. For example, oscillations were observed in batch growth upon diauxic shift to pure respiration on ethanol from aerobic fermentation (Mochan and Pye, 1973). More recently, the YMC was observed in batch-grown phosphate-starved yeast with ethanol as a sole carbon source. These yeast remained blocked in the G1 phase of the cell cycle due to phosphate starvation, which suggests that the YMC can operate independently of the CDC (Slavov et al., 2011). Last, respiratory oscillations have been observed in batch cultures of budding yeast (*Saccharomyces*), clinically significant yeasts (*Candida*), and fission yeast (*Pombe*) synchronized at similar cell cycle stages through centrifugation; see review by (Lloyd, 2008). Strikingly, these respiratory oscillations persisted during a subsequent cell cycle block, which shows that respiration can oscillate autonomously of the CDC.

Coordinated cross-talk between YMC and CDC oscillators

There is evidence to support a bi-directional link between the YMC and CDC where both of these two oscillations are autonomous but mutually coupled. For example, induction of a premature HOC phase via addition of ethanol to a culture in LOC also triggers cell cycle entry (Robertson *et al.*, 2008). Recent work has shown that increasing acetyl-coA levels during trehalose catabolism and glucose fermentation drives chromatin acetylation and increases the global expression of numerous genes, including the G1 cyclin *CLN3* (a key regulator of the cell cycle START) (Shi and Tu, 2013). Other work strongly suggests that the ATP/ADP ratio (which oscillates with the YMC and peaks in HOC) directly regulates global gene expression via ISWI and RSC during the YMC (Machne *et al.*, 2012; Amariei *et al.*, 2014). Thus, chromatin regulation may provide a generic mechanism by which the YMC can regulate the CDC.

A global survey of Cdk1 targets by the Morgan lab recently identified Nth1, a catabolic enzyme that degrades trehalose, as a likely target of Cdk1 (Holt *et al.*, 2009). Unpublished work shows that Cdk1 phosphorylation of Nth1p can trigger the liquidation of stores of trehalose (private communication, Jan Skotheim). We speculate that these mechanisms create a positive feedback loop (liquidation of storage carbohydrates can trigger cell cycle Start, and progress through cell cycle Start can trigger liquidation of storage carbs) that may be responsible for the strong coupling between the YMC and CDC. The bidirectional coupling may be a mechanism to ensure that these two oscillators with different average periods nonetheless maintain coordination between the production of energy and biomass and progress through the cell division cycle.

MATERIALS and METHODS:

Yeast strains

We used diploid prototrophs in our experiments. Yeast strain DBY12007 was generously provided by the lab of Dr. David Botstein (Silverman *et al.*, 2010). Haploid strains CEN.PK (MAT α and MAT a) were gifts from the lab of Dr. Benjamin Tu (Shi *et al.*, 2010). We mated these haploids to produce diploid CEN.PK. Wild isolates YJM128 and YPS670 were chosen for their breadth of ecological niches, and were gifts from the lab of Dr. Paul Magwene. Strain YJM128 is a clinical isolate from the lung of an HIV patient (McCusker *et al.*, 1994). YPS670 was originally isolated from oak sap exudate in the forests of Pennsylvania (Murphy *et al.*, 2006).

Growth medium

We used minimal defined (MD) growth medium, which we adapted from several sources (Saldanha *et al.*, 2004; Brauer *et al.*, 2005; Tu *et al.*, 2005). We first tested glucose concentrations between 0.1% and 2% mass/volume before choosing 0.25% for all subsequent experiments, due to the ability of all strains to oscillate at this concentration. MD medium was prepared by first autoclaving a base solution of 0.1 g/L CaCl₂•2H₂O, 0.5 g/L Mg₂(SO₄)₂ •7H₂O, 2 g/L K₂HPO₄, 5 g/L (NH₄)•2SO₄, and 70 μ L/L 96% H₂SO₄. After cooling, this autoclaved solution was supplemented with 1 mL/L 1000x vitamin solution, 100 μ L /L, 10,000x trace salt solution, 0.02 g/L solid FeSO₄•7H₂O, 100 μ L /L Sigma Antifoam 204, and 0.25% glucose mass/volume.

10,000x trace salt solution consisted of 10g ZnSO₄•7H₂O, 5g CuSO₄•5H₂O, 1g MnCl₂•4H₂O, and 100 mL filter sterilised H₂O. 1,000x vitamin solution consisted of 1 mg Biotin, 200 mg Calcium pantothenate, 1 mg Folic acid, 1000 mg Inositol, 200 mg Niacin, 100 mg P-aminobenzoic acid, 200 mg Pyridoxine HCL, 100 mg Riboflavin, 200 mg Thiamine HCL, and 500 mL H₂O, which was autoclaved for 40 minutes to dissolve vitamin powders, and immediately cooled to 4° C. All chemicals were purchased from Sigma Aldrich.

Continuous culture conditions

Continuous cultures were grown in a Multifors 6-vessel microbial system (Infors USA, Laurel, MD). All vessels had a 1000 mL working volume at 30° C. They were aerated with 1000 mL/minute of filtered room air and stirred at 550 rpm. Dissolved oxygen (pO₂) and pH were measured with OxyProbe and FermProbe (Broadley-James, Irvine CA). We recorded the pH, pO₂, and pump flow rates every minute using IRIS V5 and V6 monitoring software (Infors USA, Laurel, MD). The pH was clamped by the chemostat control computer via the automatic addition of sterile 1M phosphoric acid or 1M sodium hydroxide, as necessary. Medium pH was maintained at 4.0 ± 0.1 for all strains, except YPS670 which was kept at 5.0 ± 0.1 because its growth was significantly diminished at lower pH. Vessels were autoclaved containing 1000 mL of 1x base solution, and cooled to a working temperature of 30° C. To prevent precipitation, sterile trace salts, vitamins, and glucose were added after cooling to working temperature, just before yeast inoculation. We calibrated feed pumps by measuring the volume of liquid dispensed before autoclaving when run at 100% speed. The pO₂ sensors were calibrated by setting the probe to 100% at full oxygen saturation and 0% at full nitrogen gas in place of air. Below we verified that the measured pO₂ profile reflects actual dissolved oxygen levels as well as the yeast oxygen uptake rate due to very rapid oxygen mass transfer kinetics.

Measured pO₂ reflects the actual dissolved oxygen levels

We first measured the probe response time (τ_r) and the oxygen mass transfer coefficient (k) of our chemostat using the dynamic method (Garcia-Ochoa and Gomez, 2009). The probe and oxygen dynamics are described by:

$$\frac{dpO_{2m}}{dt} = \frac{(pO_2 - pO_{2m})}{\tau_r} \quad (\text{Eq. 1})$$

$$\frac{dpO_2}{dt} = k \cdot (pO_2^* - pO_2) \quad (\text{Eq. 2})$$

where pO_{2m} is the measured dissolved oxygen level, pO_2 is the actual dissolved oxygen level, and pO_2^* is the target steady-state level that the system approaches. We determined τ_r and k by experimentally switching gas-line input from nitrogen gas ($pO_2^* = 0\%$) to air ($pO_2^* = 100\%$). The dynamics of reaching the new steady-state is given by:

$$pO_{2m}(t) = 100\% \cdot \left[1 + \frac{\tau_r \cdot k \cdot e^{-\frac{-(t-t_0)}{\tau_r}} - e^{-k \cdot (t-t_0)}}{1 - \tau_r \cdot k} \right] \quad \text{for } t > t_0 \quad (\text{Eq. 3})$$

where t_0 is a variable time lag that arises from the manual switching of gas lines and adaptive temporal sampling of the recording software. The oxygen mass transfer coefficient (k) in a chemostat depends on the rate of air-flow, the impeller type, rotation speed, and the temperature. We fit Eq. (3) to experimental pO_{2m} from the same chemostat conditions used in our YMC experiments: air flow rate of 1000 mL / minute mixed with a Rushton impeller at 550 RPM at 30° C. Our best fit to multiple data sets gives a response time $\tau_r \approx 0.19$ mins and oxygen mass transfer coefficient $k \approx 1.2 \text{ mins}^{-1}$; see Figure S1A for one such fit.

By manipulating Equations 1 and 2, we can derive the time-varying steady state level $pO_2^*(t)$ from $pO_{2m}(t)$ (and associated first-derivative $pO_{2m}'(t)$ and second-derivative $pO_{2m}''(t)$):

$$pO_2^*(t) = pO_{2m}(t) + \tau_r \cdot pO_{2m}'(t) + \left[\frac{\tau_r \cdot pO_{2m}''(t) + pO_{2m}'(t)}{k} \right] \quad (\text{Eq. 4})$$

The value of $pO_2^*(t)$ represents what $pO_{2m}(t)$ would be if the chemostat had an infinite oxygen mass transfer coefficient and zero probe response time. We used Eq. 4 and our best-fit k and τ_r parameters to convert measured $pO_{2m}(t)$ into $pO_2^*(t)$. The measured dissolved oxygen profile is nearly identical to steady state oxygen levels; see Figure S1B. Thus, we will use pO_2^* , pO_2 , pO_{2m} interchangeably throughout our manuscript.

Levels of pO_2 are determined by the yeast oxygen uptake rate

It is important to clarify the relationship between high and low dissolved oxygen (pO_2)

and the *rate* of oxygen consumption by yeast in a bioreactor. Briefly, the dissolved oxygen level (pO_2) is a balance between the mass action oxygen transfer *rate* (k) from gas to liquid and the yeast oxygen uptake *rate* (OUR). Mathematically, the rate of change of pO_2 is given by Eq. (2):

$$\frac{dpO_2}{dt} = k \cdot (pO_2^* - pO_2) - OUR \quad (\text{Eq. 5})$$

where pO_2^* is the saturation level of dissolved oxygen (100%) that our chemostat would approach in the absence of any yeast respiration. The general solution to Eq. (5) is:

$$pO_2(t) = \underbrace{pO_2(0)}_{\text{Initial}} \cdot e^{-kt} + \underbrace{\left[pO_2^* - \frac{OUR}{k} \right]}_{\text{Steady-state}} \cdot (1 - e^{-kt}) \quad (\text{Eq. 6})$$

The bioreactor mass transfer coefficient (k) determines the timescale of how quickly an initial pO_2 reaches its new steady-state after a change in yeast OUR. Our calibration measurements showed that steady-state is reached quickly ($k \approx 1.2 \text{ mins}^{-1}$). When the yeast OUR changes on a timescale that is slower than k , Eq. (6) is well-described by the steady-state solution:

$$pO_2(t) = pO_2^* - \frac{OUR(t)}{k} \quad (\text{Eq. 7})$$

If yeast OUR is high, then pO_2 is low (or 0%, the absolute minimum). Conversely, if yeast OUR is low, then pO_2 is high (or 100%, the absolute maximum). Our control experiments indicate that dissolved oxygen levels (pO_2) accurately reflect the yeast oxygen uptake *rate* in our chemostat.

Automated data analysis

Each chemostat run lasted from 1 to 3 weeks. During each run, we scanned through different dilution rates. We allowed enough time to reach steady-state after each change in the dilution rate. There were 16, 12, 9, and 10 chemostat runs and we collected a total of 52, 50, 16, 30 YMC time-series over a range of different dilution rates for strains CEN.PK, DBY12007, YJM128 (lung), and YPS670 (oak), respectively. We discarded those parts of the time-series that had not yet reached steady state by removing the first 1000 minutes after initiation of continuous growth, and the first

250 minutes after each subsequent change in dilution rate. We applied a MATLAB-based peak-detection algorithm to these steady-state pO₂ time-series to identify local maxima and minima. We included a minimum peak size filter and a smaller local-peak-exclusion size filter to ensure that only one local maximum and minimum per cycle were identified. Filter parameters had to be adjusted manually for each experiment to ensure that exactly one maximum and minimum were identified per cycle. Entry into HOC was defined as the time at which pO₂ drops to 65% of the local pO₂ maximum to the next local minimum, whereas entry into LOC was defined as the time at which the pO₂ rises to 65% of the way from a local minimum to the next local maximum. Our definition of entry into HOC was co-incident with the onset of aerobic fermentation and acidification of the medium; see Figure S1C. For every time-series at a fixed dilution rate, we recorded the time of each entry into HOC, minimum pO₂, exit from HOC, maximum pO₂, and the distribution of lengths of each YMC, HOC, and LOC. We discarded a time-series if the coefficient of variation of τ_{ymc} , τ_{hoc} , or τ_{loc} was greater than 0.2. This stringent filter flagged and removed 19 YMC time-series with stability problems (less than 9% of the data).

Quantitative relationship between YMC and CDC

The total YMC period (τ_{ymc}) was split into an HOC interval (τ_{hoc}) and LOC interval (τ_{loc}). Each strain dataset $\{\tau\}$ contained average $\{\tau_{ymc}, \tau_{hoc}, \tau_{loc}\}$ for each YMC time-series at different dilution rates. For each strain, the $\{\tau\}$ at different dilution rates (τ_{cdc}) were best fit using nonlinear least squares fitting (nlinfit in MATLAB). We fit these data in Figure S2 to a linear model.

$$\tau = m \cdot \tau_{cdc} + b \quad (\text{Eq. 8})$$

where b is the y -intercept, m is the relative slope. We also fit these data in Figure S2 to a hyperbolic model (i.e. Michaelis-Menten).

$$\tau = \tau_{max} \cdot \frac{\tau_{cdc}}{\tau_{cdc} + \kappa} \quad (\text{Eq. 9})$$

where τ_{max} is the maximum YMC and κ is the τ_{cdc} at which τ is half-maximum. The hyperbolic model becomes a linear model when the time variables τ are inverted to frequencies (f), such that

$$f = \frac{1}{\tau_{max}} \cdot (1 + \kappa \cdot f_{cdc}) \quad (\text{Eq. 10})$$

where $f=1/\tau$ and $f_{cdc}=1/\tau_{cdc}$.

In the end, all data were best fit by a mixed model, where τ_{ymc} is hyperbolic, τ_{hoc} is linear, and $\tau_{loc}=\tau_{ymc}-\tau_{hoc}$; see Figures S2 and the best-fit residuals of each model to different data sets in Figure S3. The best-fit parameters of the mixed model to different data sets are listed in Table S2.

Cell cycle analysis

Samples were taken from stably oscillating chambers at 10 or 15 minute time intervals. These samples were immediately spun down, washed in distilled water, and cells were fixed in 70% ethanol at 4° C in 96-well plates. Our DNA staining protocol was modified from (Haase and Reed, 2002) to handle reduced fluid volumes in 96-well plates: cells were first washed with 200 μ L distilled water, resuspended in 100 μ L RNase A solution (Qiagen) for 2-6 hours, resuspended in 100 μ L pepsin solution for 10-15 minutes, and then stored in 200 μ L storage buffer. Before the final staining step, 30 μ L of each sample was sonicated to separate cell clumps on medium setting (strains CEN.PK, DBY12007, YPS670) or high setting (YJM128) for 45 seconds in 3x8 strips of 96-well PCR plates floating in a Bioruptor water-bath sonicator (Diagenode, Denville, NJ). The sonicated cells were immediately added to 96-well plates containing 200 μ L per well of 1 μ M SYTOX Green (LifeTechnologies). The fluorescence of these stained cells was measured with a MACSQuant VYB (Miltenyi Biotech, Bergisch-Gladbach, Germany) using laser excitation at 488 nm and a 525/50 bandpass filter. We normalized the fluorescence distributions across all time samples by (i) shifting the fluorescence by a linear amount so that the maximum of all G1 peaks were aligned, and (ii) multiplying all frequencies with a scaling factor, such that each time-point had the same rescaled cell count.

Dean-Jett-Fox and other cell cycle parsing algorithms were unable to reliably identify the fraction of S-phase cells across all time points due to the width of G1 and G2/M distributions, and asymmetrical G1 fluorescence peaks at slow growth rates. For each data set, we estimated two fluorescence thresholds that (i) separated G1 cells from S/G2/M (i.e. G1 threshold) and (ii) separated G2/M cells from G1/S (i.e. G2/M threshold). We determined G1 and G2/M threshold by first averaging the

fluorescence distribution across all time points, finding the local minimum between the G1 and G2 peaks, and choosing the G1 threshold such that it included 95% of cells with a DNA content below this local minimum. We chose the G2/M threshold such that it included 95% of the cells with DNA content above this local minimum. The fraction of cells above G1 threshold is the S/G2/M fraction and the fraction between G1 and G2/M threshold is the S fraction.

For each 8-hour time course, we also created an "average" YMC-CDC data set across a yeast metabolic cycle. The start ($t=0$ mins) of each yeast metabolic cycle was defined as entry into HOC. We partitioned the yeast metabolic cycle into 1-minute bins, and averaged DNA content, acidification, S/G2/M and S fraction across a window of 10 or 15 minutes centered on each bin. For each averaged YMC-CDC data set, we calculated Δ as the time at which S/G2/M fraction reached 50% peak-to-trough and Δ_S as the time at which S fraction reached 100% peak.

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AUTHOR CONTRIBUTIONS:

AJB conceived the study, built strains, designed and performed experiments, and analysed the data. MA built strains and performed experiments. NEB conceived the study and supervised the project. AJB and NEB wrote the manuscript. All authors edited and approved the final manuscript.

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FIGURE LEGENDS:

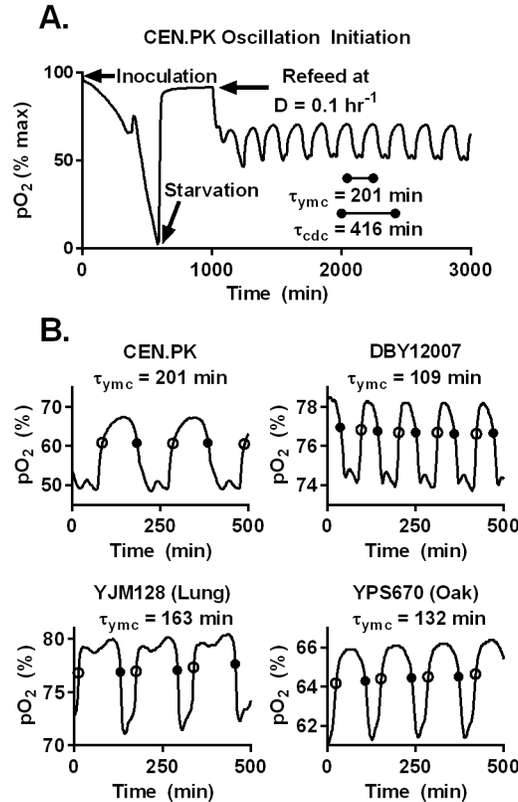


Figure 1: Measurement and analysis of the yeast metabolic cycle across different strains. (A) Representative dissolved oxygen trace (pO_2) after inoculation of chemostat with strain CEN.PK. Logarithmically growing yeasts first aerobically fermented the available glucose to produce ethanol, followed by a diauxic shift to pure respiration on ethanol. After a period of yeast starvation, we started the flow of fresh medium into the chemostat at a constant dilution rate ($D=0.1 \text{ hr}^{-1}$). The population began to exhibit clear oscillations in pO_2 after re-feeding. The period of the YMC (τ_{ymc}) is the elapsed time from peak-to-peak in pO_2 signal. At steady-state, the average CDC period (τ_{cdc}) must be equal to the inverse of the dilution rate, or $\tau_{cdc} = \ln(2)/D$. (B) Sample pO_2 traces of different yeasts at the same dilution rate. We used lab strains CEN.PK, DBY12007 and wild-isolates YJM128 (lung), YPS670 (oak). We developed an automated analysis pipeline to extract the YMC period, timing of entry into high oxygen consumption (HOC, solid circle), and timing of entry into low oxygen consumption (LOC, open circle) across different strains and dilution rates; see Methods.

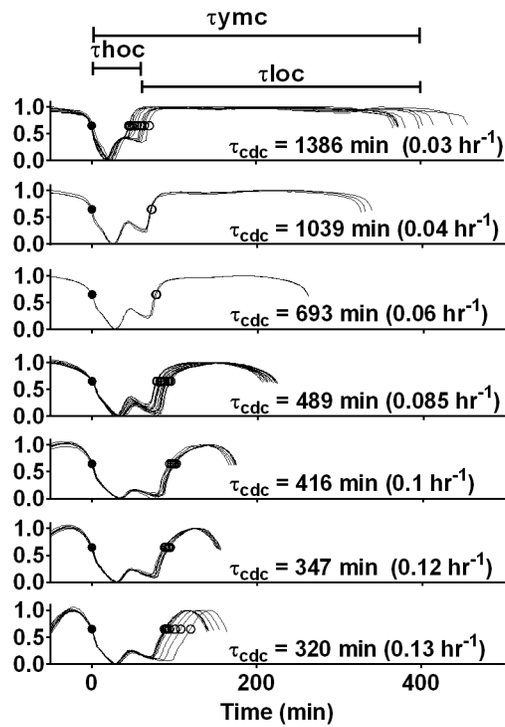


Figure 2: Increase in YMC period at slower dilution rates occurs in the LOC. The YMC of strain CEN.PK was analyzed over a range of dilutions rates ($D = 0.03$ to 0.13 hr^{-1}). All YMC oscillations in pO_2 from a single chemostat run were overlaid according to their entry into HOC (solid circle, $t=0$ mins), their amplitudes were normalized, and arranged vertically by their dilution rate. As τ_{cdc} increased, both τ_{ymc} and τ_{loc} increased whereas τ_{hoc} slowly decreased.

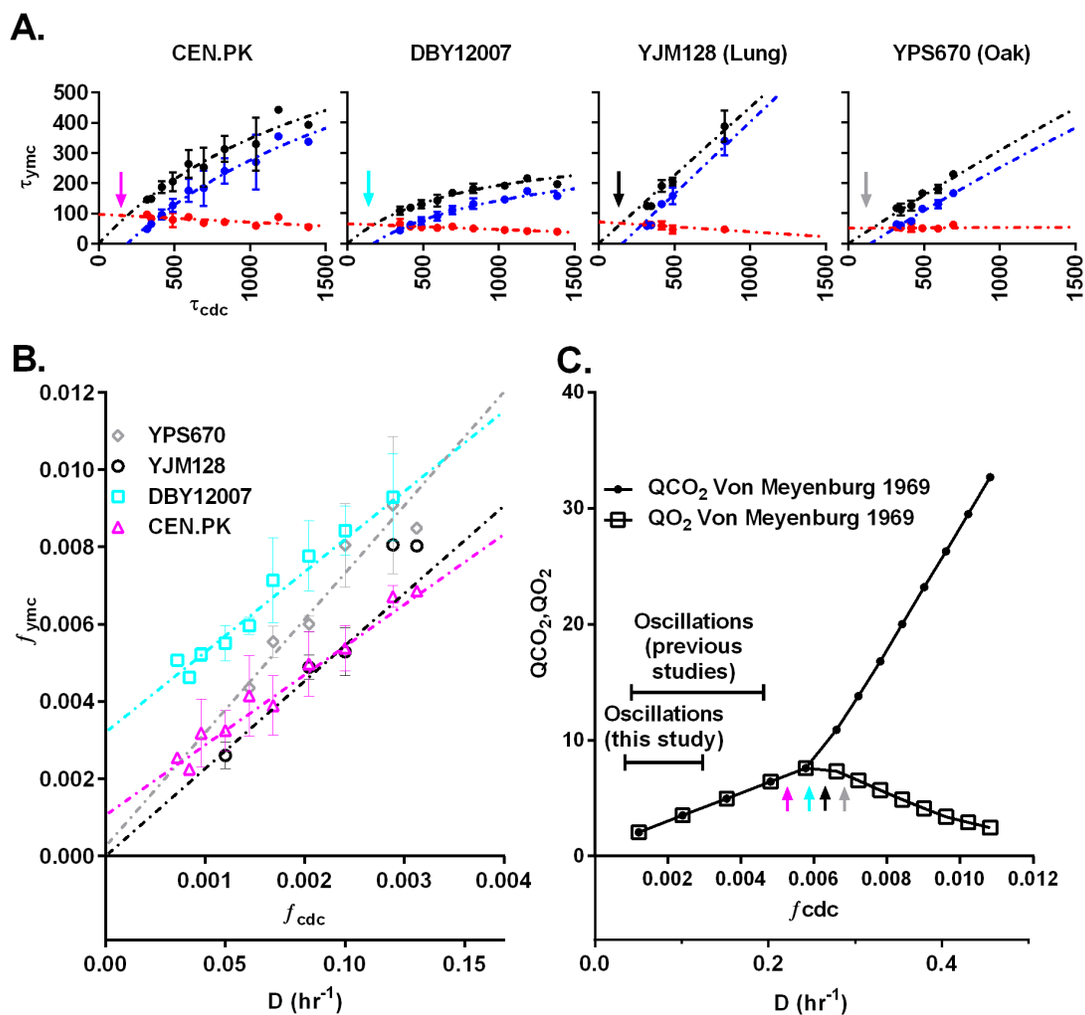


Figure 3: Quantitative relationship between the yeast metabolic cycle and cell division cycle. (A) Plots of YMC period (τ_{ymc}) (black dots), time spent in HOC (τ_{hoc}) (red dots), and time spent in LOC (τ_{loc}) (blue dots) of CEN.PK as a function of CDC period (τ_{cdc}) for CEN.PK, DBY12007, YJM128 (lung) and YPS670 (oak). For clarity, we plot the mean and standard deviation of time-series averages at identical dilution rates, which were measured during separate chemostat runs. We used nonlinear regression to best fit a mixed model to each data set; see Methods and Figures S2-S3. (B) Plot of YMC frequency ($f_{ymc}=1/\tau_{ymc}$) and CDC frequency ($f_{cdc}=1/\tau_{cdc}$) for all strains. (C) Metabolic profile of oxygen consumption (QO_2) and carbon dioxide production (QCO_2) of yeast strain LBGH1022 at different dilution rates from previous literature (Kaspar von Meyenburg, 1969). There is a transition from oxidative respiration (QCO_2 is equal to QO_2) to aerobic fermentation (QCO_2 is greater than QO_2) at a critical dilution rate, $D_c \approx 0.24 \text{ hr}^{-1}$, which corresponds to the dilution rates at which our tested strains are extrapolated to spend all their time in HOC phase (colored arrows) and not oscillate.

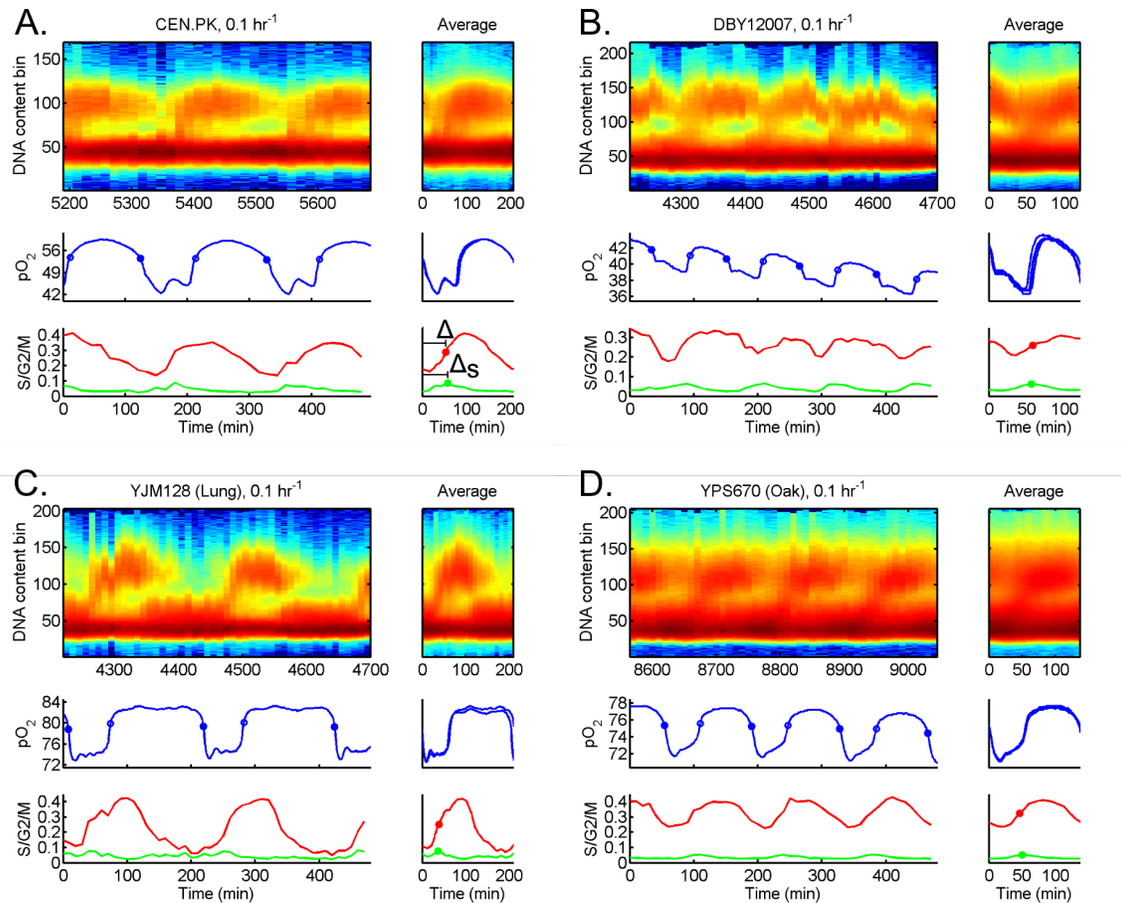


Figure 4: Timing of DNA replication relative to HOC across strains. Strains (A) CEN.PK, (B) DBY12007, (C) YJM128, (D) YPS670 were cultured in chemostat at the same dilution rate ($D=0.1 \text{ hr}^{-1}$). We extracted samples every 10 minutes and measured CDC events (DNA content) and YMC events (pO_2); see Methods. For each strain, the raw DNA content, pO_2 , S/G2/M fraction, and S fraction are plotted over several YMC (left). The average over each YMC is plotted to the right of each full data set, where $t=0$ corresponds to entry into HOC. We plot the 50% mid-point of S/G2/M fraction (solid red circle) and 100% peak of S fraction (solid green circle). The time of DNA replication (50% of S/G2/M fraction) after entry into HOC is defined as Δ , where as time of DNA replication (100% of S fraction) is Δ_s .

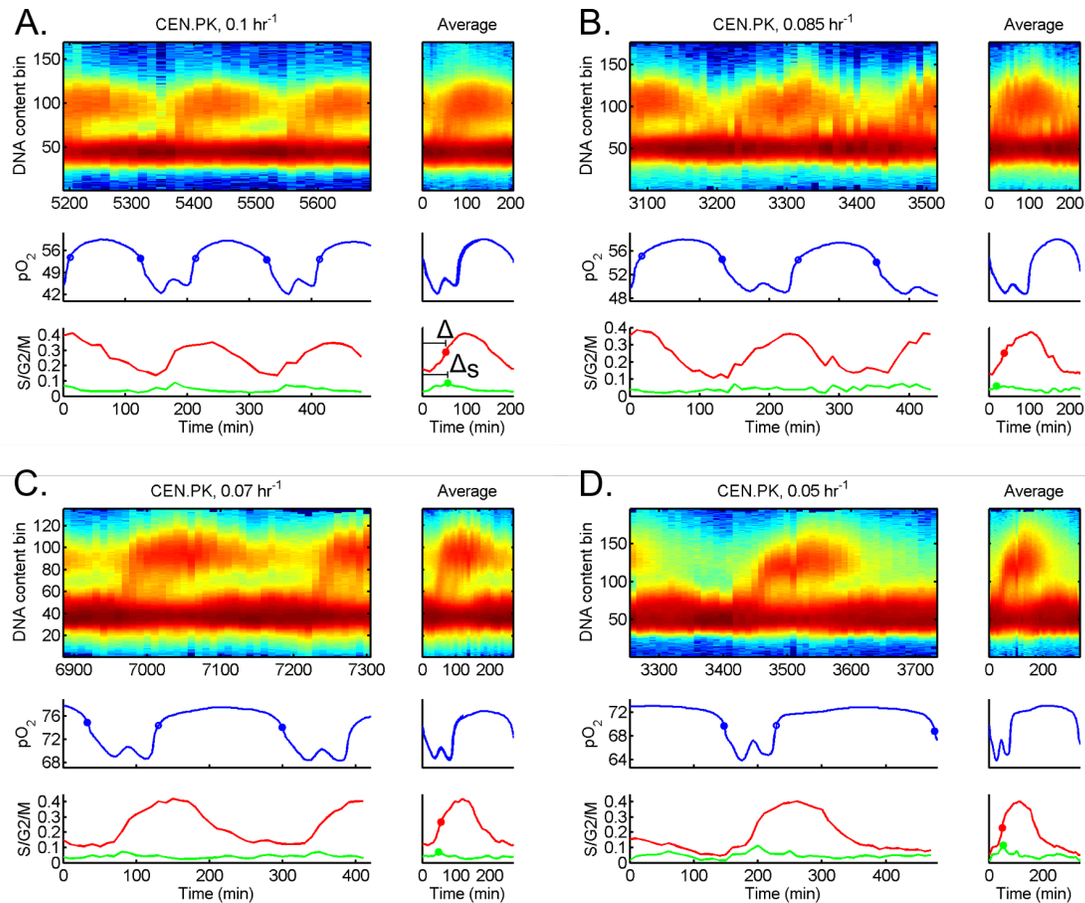


Figure 5: Timing of DNA replication relative to HOC in strain CEN.PK across growth rates. Additional cell cycle analysis for strain CEN.PK at different dilution rates: (A) 0.1 hr^{-1} , (B) 0.085 hr^{-1} , (C) 0.07 hr^{-1} , and (D) 0.05 hr^{-1} . The raw DNA content, pO_2 , S/G2/M fraction, and S fraction are plotted over several YMC (left). The average over each YMC is plotted to the right of each full data set, where $t=0$ corresponds to entry into HOC. We plot the 50% mid-point of S/G2/M fraction (solid red circle) and 100% peak of S fraction (solid green circle). The time of DNA replication (50% of S/G2/M fraction) after entry into HOC is defined as Δ , where as time of DNA replication (100% of S fraction) is Δ_s .

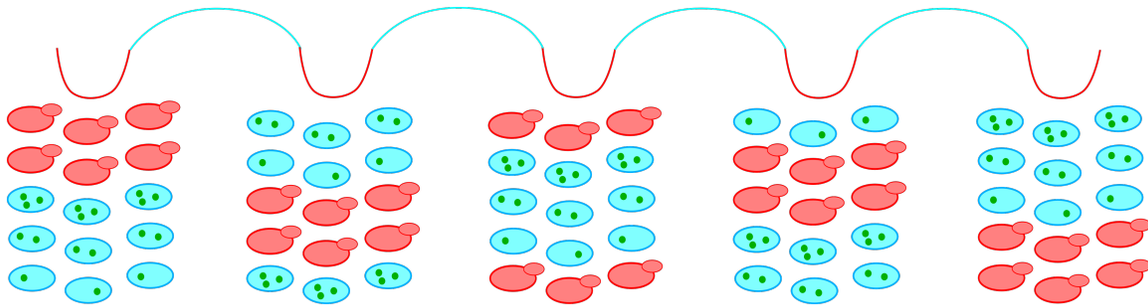


Figure 6: A simplified model of YMC-CDC coupling. The dissolved oxygen pO_2 trace indicates LOC (blue line) and HOC (red line) over multiple YMCs. At the beginning of each YMC, a fraction of the population (red, budded cells) commits to catabolizing storage carbohydrates, entering HOC, and starting the cell division cycle. These "committed" yeasts secrete metabolites, which trigger other "susceptible" yeasts with sufficient storage carbohydrates to catabolize their storage carbohydrates (Robertson *et al.*, 2008). Such autocatalytic signaling through secreted metabolites causes an avalanche of susceptible yeasts to synchronously enter HOC and commit to the CDC. However, it is clear that not all yeasts commit to cell division cycle each YMC. The rest of the yeast population in LOC (blue cells) is either refractory to metabolic signals because the cells have not accumulated sufficient energy reserves (green dots) to commit to YMC and/or has not accumulated sufficient biomass to initiate cell cycle. Start upon entry to HOC. Over the coming YMC, these "refractory" yeasts in LOC continue to build up their reserves of storage carbohydrates and biomass, such that a new fraction of yeasts will be ready to spontaneously initiate and trigger other susceptible yeasts to commit to the next YMC. The yeast population in a low-glucose chemostat thus self-organizes into multiple staggered cohorts, such that only one cohort synchronously enters the CDC each YMC (i.e. one-to-some coupling). Cells likely migrate between cohorts over time due to cell-to-cell variability in both the YMC and CDC.

TABLE LEGENDS:

D (hr ⁻¹)	0.1	0.085	0.07	0.05
CEN.PK	56.5 ± 0.7	53.7 ± 32.6	48.0	53.5 ± 0.7
DBY12007	58.0 ± 2.0	59.0 ± 5.7	59.0	54.0
YPS670 (oak)	44.8 ± 8.5	58.0	44.0	
YJM128 (lung)	33.3 ± 3.8			

Table I: Timing of entry into HOC and DNA replication (Δ_S) across strains and growth rates. The mean and standard deviation of Δ_S interval (mins) of various strains at different growth rates. Mean and standard deviation are for biological replicates of the same strains at identical growth rate on different days. YJM128 was unstable at lower dilution rates.