Circadian Output, Input, and Intracellular Oscillators: Insights into the Circadian Systems of Single Cells


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Circadian output comprises the business end of circadian systems in terms of adaptive significance. Work on Neurospora pioneered the molecular analysis of circadian output mechanisms, and insights from this model system continue to illuminate the pathways through which clocks control metabolism and overt rhythms. In Neurospora, virtually every strain examined in the context of rhythms bears the band allele that helps to clarify the overt rhythm in assexual development. Recent cloning of band showed it to be an allele of ras-1 and to affect a wide variety of signaling pathways yielding enhanced light responses and assexual development. These can be largely phenocopied by treatments that increase levels of intracellular reactive oxygen species. Although output is often unidirectional, analysis of the prd-4 gene provided an alternative paradigm in which output feeds back to affect input. prd-4 is an allele of checkpoint kinase-2 that bypasses the requirement for DNA damage to activate this kinase; FRQ is normally a substrate of activated Chk2, so in Chk2PRD-4, FRQ is precociously phosphorylated and the clock cycles more quickly. Finally, recent adaptation of luciferase to fully function in Neurospora now allows the core FRQ/WCC feedback loop to be followed in real time under conditions where it no longer controls the overt rhythm in development. This ability can be used to describe the hierarchical relationships among FRQ-Less Oscillators (FLOs) and to see which are connected to the circadian system. The nitrate reductase oscillator appears to be connected, but the oscillator controlling the long-period rhythm elicited upon choline starvation appears completely disconnected from the circadian system; it can be seen to run with a very long noncompensated 60–120-hour period length under conditions where the circadian FRQ/WCC oscillator continues to cycle with a fully compensated circadian 22-hour period.

INTRODUCTION

The innate ability to gauge time of day juxtaposed with the perception of environmental light represents one of the most widespread and closely coupled forms of cellular, tissue, and organismal regulation across a broad range of taxonomic groups. Changes in light fluence and wavelength are detected with the harvesting of photons by chromophore-binding photoreceptive molecules allowing immediate detection of important environmental changes. Biological rhythms provide organisms with the ability to anticipate environmental changes arising from the Earth’s rotation. The physiological and molecular mechanisms of daily biological rhythms, called circadian rhythms, have been the focus of study for more than a century, but the past quarter century has witnessed a wealth of information and one of the earliest model systems to be shown to possess a bona fide circadian system (Pittendrigh and Bruce 1959). Close to two decades ago, we predicted that daily clock control of gene expression might be a major aspect of output (Loros et al. 1989). This has proved not only to be true for Neurospora, but also to be universally the case for all circadian systems (see, e.g., Liu et al. 1995; Harmer et al. 2000; Ceriani et al. 2002; Duffield et al. 2002; Panda et al. 2002). We performed the first systematic screen, using a novel technique at the time (subtractive hybridization), of two different mRNA populations, one isolated from morning and one from evening, to identify genes whose mRNAs were rhythmically abundant (Loros et al. 1989). These

early genes were subsequently shown to be regulated at the level of transcriptional rate, suggesting that there would be cis-acting sequences in the promoter regions of these genes in addition to trans-acting factors that conferred circadian regulation (Loros and Dunlap 1991). Continuing studies using differential hybridization (Bell-Pedersen et al. 1996b), cDNA sequencing (Zhu et al. 2001), and cDNA and oligonucleotide microarrays (Correa et al. 2003; Nowroussian et al. 2003) have isolated several hundred more rhythmically regulated genes. Full-genome microarrays for the approximately 10,500 Neurospora genes have recently become available and are currently being used by more than one lab to examine clock regulation of gene expression at different points in the life cycle and under various growth conditions. In multicellular organisms, the identification of genes regulated by the clock was largely on a gene-by-gene basis until the advent of microarrays (Harmer et al. 2000; Duffield 2003) in addition to an innovative differential display protocol called ADDER used to isolate several cycling liver genes (Kornmann et al. 2001). These studies already have, and will continue to, dramatically improve our understanding of the global nature of circadian regulation on gene expression at the tissue and organ levels in plants and animals. Of great current interest is the mechanistic understanding of clock control of specific sets of genes at the level of individual cells, a more difficult problem to approach using in vivo samples. Adopting the same paradigms used in whole organisms, tissue cultures using individual cell types that display circadian properties (Balsalobre et al. 1998) have been recently leading the way (Duffield et al. 2002).

THE CIRCADIAN CLOCK AND THE CIRCadian SYSTEM

The haploid filamentous fungus Neurospora crassa grows as incompletely septate, highly branched mycelia capable of fusing with separate but compatible strains, thereby conning both cytoplasm and nuclei in a common cytoplasmic compartment. Neurospora is a superb organism for genetics and biochemistry with a well-defined genetic history, allowing the maintenance of lethal mutations, gene-dosage analysis, and complementation. The genome is about three times the size of yeast at 4.3\(x10^7\) base pairs, encoding about 10,500 highly nonredundant genes expressed at different stages of the sexual and asexual life cycle. The clock in Neurospora is now monitored in a number of different ways. Historically and still of great utility is the monitoring of asexual conidiospore (conidia) development as the fungus grows on medium placed in hollow glass culture tubes called race tubes (Fig. 1) (Ryan et al. 1943). One consequence of growing in the enclosed environment of the race tube is that CO\(_2\) levels become elevated, resulting in suppression of condensation and masking of the periodic formation of conidia within the tube. A mutation called band (bd) (Sargent et al. 1966) overcomes this repression and has been used in Neurospora laboratory stocks in rhythm studies for the last 40 years. The bd locus has recently been cloned and found, as expected, not to alter function of the clock’s central mechanism, and also not completely surprisingly, it was found to have global effects on the regulation of clock output (Belden et al. 2007). For molecular and biochemical analysis, Neurospora is grown in liquid culture (Nakashima 1981; Perlman et al. 1981; Loros et al. 1989; Loros and Dunlap 1991; Aronson et al. 1994b; Garceau et al. 1997). Mycelial disks in liquid culture (Shi et al. 2007) maintain endogenous rhythmicity and phase, even upon transfer to solid medium. The ability to use luciferase (Morgan et al. 2003) to follow either in vivo transcriptional or translational fusions of rhythmically expressed genes has recently become a valuable tool to monitor clock progress by the complete codon optimization of firefly luciferase (Gooch et al. 2007).

An Overview of the Circadian Oscillator

Genetic, molecular, and biochemical analyses of the Neurospora clock have led to truly extraordinary advances in our understanding of much of the organism’s circadian properties, including the generation and sustainability of rhythmicity, phase resetting by light and temperature, and the means by which the clock controls metabolism and behavior. The following is an extremely brief overview of the molecular underpinnings: In Neurospora, as in all eukaryotes studied to date, a critical part is played by an autoregulatory, molecular feedback loop between the FREQUENCY protein(s) (FRQ); the White Collar transcription factors (WC-1 and WC-2) (Aronson et al. 1994ab; Crosthwaite et al. 1997; Dunlap 1999; Lee et al. 2003); and at least one other associated protein, FRH (Cheng et al. 2005). Expression at the frq locus is activated by the heterodimeric White Collar Complex (WCC) (Crosthwaite et al. 1995; Ballario et al. 1996, 1998; Linden and Macino 1997; Linden et al. 1997; Talora et al. 1999; Collett et al. 2001, 2002; Denault et al. 2001; Cheng et al. 2002). Long and short forms of FRQ (Garceau et al. 1997), produced from temperature-regulated alternative splicing (Colot et al. 2005; Diernfellner et al. 2005), then feed back to block activation of frq expression (Aronson et al. 1994b; Garceau et al. 1997; Merrow et al. 1997; Froehlich et al. 2003), resulting in rhythmic waves of both frq mRNA and FRQ protein, highly regulated by both phosphorylation (Garceau et al. 1997; Liu et al. 2000; Schafmeier et al. 2006) and ubiquitination (He et al. 2003, 2005) over the course of the 24-hour day. The environment signals to the clock, most notably via light and temperature, result in phase resetting such that the organism is in synchrony with the external world (Crosthwaite et al. 1995; Liu et al. 1998; Kramer et al. 2003; Price-Lloyd et al. 2005; Hunt et al. 2007) (for more a detailed overview, see Dunlap et al., this volume). Finally, the FRQ/WCC feedback oscillator can signal temporal information to the cell via daily changes in transcript abundance of pertinent output genes (Loros et al. 1989). A major regulatory means to this end is through cis-acting sequences in promoters that confer daily changes of transcriptional rates (see, e.g., Loros and Dunlap 1991; Bell-Pedersen et al. 1996a).

Noncircadian Oscillators

Above, we alluded to the classic “input-oscillator-output” paradigm (Eskin 1979), but it is clear that organisms
contain a circadian system, not just an oscillator mechanism with linear inputs and outputs. Part of the system will doubtlessly include other oscillators (see, e.g., Dunlap and Loros 2004). As a field, regardless of organism, we are focusing more closely on individual cells, and as we do, examples of noncanonical rhythms are appearing more frequently. Oscillations can occur in any cellular pathway that has negative feedback regulation and a lag in at least one step, a common biological occurrence. Some of these oscillators will be unrelated and unconnected to the clock, and some may be part of the circadian system, either by having a necessary role in the clock oscillatory mechanism or by feeding into the clock, somewhere in the input or core oscillator, to alter properties of the clock. In addition, some will be “slave” oscillators (see below), controlled by the clock at some step within its own feedback to confer the properties of circadian period length and phase stability. Identification and elucidation of these alternative oscillators are major topics of interest to chronobiologists and are dealt with in depth below.

OUTPUT FROM THE CLOCK IS THE NATURE OF TEMPORAL INFORMATION

The utilitarian and therefore most important aspect of all clocks, often referred to as output, is the ability to invoke “time of dayness” onto the organism, such that it can predict daily changes in the environment to regulate its own changing metabolic needs over the course of the diurnal cycle. Most outputs may occur in a linear fashion and be distinct from the clock mechanism.

Neurospora has been developed as a paradigmatic system for understanding the physiological changes governed by the clock, including development of the hypothesis that changes in transcription would be a universal means by which the clock mechanism could regulate clock output. Testing this hypothesis led to the first genome-wide screens, using subtractive hybridization methodologies, for the daily changes in gene expression alluded to above and to coining the term “clock-controlled gene” (ccg) (Loros et al. 1989). The identity of hundreds of ccgs are currently known through a combination of differential hybridization, cDNA and oligomer microarrays, and individual gene examination (Bell-Pedersen 2000; Correa et al. 2003; for review, see Loros et al. 2003; Nowrousian et al. 2003).

Isolation of the initial ccgs led to a first working model for defining output at the molecular level of gene expression. First, the endogenous rhythm of expression of a ccg would persist under constant conditions, reflecting control exerted by the clock, as opposed to external factors such as changes in light intensity. Second, a defining test was that the period length of the molecular rhythm would reflect the genotype of the strain. For example, in Neurospora, the first ccgs were shown to have changes in mRNA abundance with 22-hour period lengths in wild
type and 29-hour period lengths when examined in the long-period strain fraq. The third initial criterion was that the inactivation of the ccg would have no effect on the clock, demonstrating that the gene represented a true molecular output in the circadian system as a whole. We now know that this third criterion is not universally true and although it may be true that most ccgs in any cell, tissue, or whole organism can be considered linear outputs, distinct from the mechanistic core of the oscillator mechanism, there exist important examples where an output can influence aspects of oscillator function, leading to coupled loops within the system.

With the early isolation of ccgs in several organisms, the question of ccg function emerged: What was the clock used for in different organisms and were there overlaps among organisms? It had been suggested that, as a general rule, clocks controlled “nonhousekeeping” processes that would be distinctly organism-dependent. This idea may have arisen due to the spectrum of output processes that had been studied before molecular analysis and that included such diverse biological functions as locomotor activity and behavior, leaf movement, asexual development, cardiovascular function, electrolyte balance, and timing of cell division. An observation surfacing from early clock genetics was that organisms were viable in the absence of operational clocks, leading to the thought that clock components, and possibly their regulatory outputs, would be found to be nonessential. We now know that virtually all aspects of biological function are controlled by the clock in Neurospora (Fig. 2) as well as in other systems examined (see, e.g., Liu et al. 1995; Harmer et al. 2000; Ceriani et al. 2002; Duffield et al. 2002; Panda et al. 2002; Wijnen and Young 2006). For example, the glycolytic gene encoding glyceraldehyde-3-phosphate dehydrogenase (GAPDH), the rate-limiting and first energy-harvesting enzyme in glycolysis, was identified early as a ccg in Neurospora (Bell-Pedersen et al. 1996b; Shinozaka et al. 1998) and subsequently found in the dinoflagellate Gonyaulax (Fagan et al. 1999), chick retina (Bailey et al. 2004), mouse hepatocytes (Temme et al. 2000), and elsewhere (see, e.g., Iwasaki et al. 2004; Kamphuis et al. 2005).

The clock regulates a broad variety of cellular functions in the Neurospora cell. RNAs were harvested over two circadian cycles and probed with oligomeric microarrays representing the approximately 10,500 genes from Neurospora. The results from two biologically independent sets of RNAs found 64% of the genes passed detection criteria with approximately 10% of those classified as clock-controlled genes (ccgs). About half of these displayed functional hits in the Functional Catalogue (http://mips.gsf.de/projects/functat) (C.-H. Chen et al., unpubl.).

**Figure 2.** The clock regulates a broad variety of cellular functions in the Neurospora cell. RNAs were harvested over two circadian cycles and probed with oligomeric microarrays representing the approximately 10,500 genes from Neurospora. The results from two biologically independent sets of RNAs found 64% of the genes passed detection criteria with approximately 10% of those classified as clock-controlled genes (ccgs). About half of these displayed functional hits in the Functional Catalogue (http://mips.gsf.de/projects/functat) (C.-H. Chen et al., unpubl.).

**THE SPECTRUM OF CCGS WITHIN A SINGLE CELL TYPE MAY REFLECT GROWTH/EXTERNAL CONDITIONS**

Certainly, GAPDH does not cycle in all cells nor under all conditions (Okamura et al. 1999; Kobayashi et al. 2004). An interesting feature of GAPDH, in addition to its role in glycolysis, is its ability to alter a cell’s redox state through the reduction of NAD+ to NADH. It may be that the clock is used to control GAPDH in specific cells or during specific oxidative/reduction conditions. In Neurospora, the spectrum of ccgs certainly reflects growth conditions. Early microarray studies identifying ccgs from high glucose (Nowrouzian et al. 2003) versus lower glucose (Correa et al. 2003) cultures found largely nonoverlapping sets of cycling genes that ranged between 5% and 20%, respectively, of all genes on the array.

A lesson about the plasticity of circadian gene regulation within a single cell type and within the organism can be deduced from the identity of the bd gene, an allele commonly used in the background of laboratory stocks of Neurospora used for rhythms research. In the beginning years of Neurospora clock research, bd was identified as a spontaneous mutation. It was shown not to alter the canonical properties of the clock, but to significantly enhance the conidiation rhythm when strains were grown on race tubes (Sargent et al. 1966). Since this time, virtually all circadian research has been performed with bd-containing strains (for review, see Dunlap and Loros 2005; Liu and Bell-Pedersen 2006), although bd was known to display enhanced effects on both light and clock-regulated gene expression (Arpaia et al. 1993, 1995). A completed genome and single-nucleotide polymorphism (SNP) analysis fostered the identification of bd as a T791I point mutation in ras-l (Belden et al. 2007). RAS is a small, conserved, membrane-attached G protein, activated by exchange of GDP with GTP, which is modulated by guanine nucleotide exchange factors. It is a key player in numerous cellular signaling cascades, including those that link extracellular signals to gene expression (see, e.g., Mitin et al. 2005). The ras-l* hypermorph is somewhat more active in GDP/GTP exchange than the wild type (Belden et al. 2007). Not unexpectedly, genes involved in asexual development were found to be up-regulated in this altered function mutant. In addition, photo-induced expression of the wc-l gene is also increased in the ras-l* background. Importantly, in a ras-l* background, the C6 zinc cluster transcription factor fluffy (fl), which is both necessary and sufficient for asexual development (Bailey-Shrode and Ebbole 2004) and is itself rhythmically expressed (Correa and Bell-Pedersen 2002), is expressed at significantly enhanced levels both after light exposure and rhythmically in the dark. This enhanced expression is the cause of the increased visibility of rhythmic conidiation in ras-l* containing strains (Fig. 3) (Belden et al. 2007).

While investigating the source of the ras-l* signal that results in changes in expression of light and clock-regulated genes, we found that altering the amount of reactive oxygen species (ROS) could also influence the banding of wild-type strains in race tubes. ROS are increasingly rec-
Some outputs feed back to modulate clock function, resulting in additional loops

Most ccgs, when deleted from the system through gene inactivation, show no effect on clock function. However, some circadianly regulated outputs can change the workings of the clock. Two recent examples of different ways this can happen in Neurospora are detailed below. The first output feeds back to alter the way light information is received by the clock, and the second output feeds back to alter the biochemical nature of a clock component.

A Molecular Output Can Feed Back to Input

Research on Neurospora has had a leading role in understanding the molecular basis of how light resets the clock, a process called entrainment (Crosthwaite et al. 1995), and that includes gating (Heintzen et al. 2001). Much of the central mechanism of light entrainment in Neurospora is conserved in mammals (Shigeyoshi et al. 1997). Gating refers to the condition in which the clock regulates its own input such that it responds to an identical stimulus (e.g., a specific amount and wavelength of light) in nonidentical ways at different times of the circadian day. In gating, output and input become mechanistically merged. The WC-1 protein is the primary blue light photoreceptor for Neurospora (Froehlich et al. 2002; He et al. 2002). A covalently bound FAD chromophore in the WC-1 light-oxygen-voltage (LOV) domain absorbs photons, leading to rapid and strong activation of frq transcription by the WCC in response to light input and to clock resetting (Crosthwaite et al. 1997). The strength of this frq-induction signal is modulated by another LOV domain containing photoreceptive protein called VIVID (VVD) that gates the light information coming into the clock at different circadian times (Heintzen et al. 2001). The expression of vvd is clock-controlled, making vvd both a ccg and a clock input (Fig. 4). VVD is also responsible for the ability of Neurospora to sense changes in light intensity (Schwerdtfeger and Linden 2003; Schwerdtfeger et al. 2003), a process termed photoadaptation, and it additionally has a role in temperature compensation of phase (Heintzen and Liu 2007; Hunt et al. 2007). In response to light, a transient cysteine-flavin adduct forms in the LOV domain that promotes breaking of the hydrogen bonds holding an amino-terminal helix; this conformational change results in signaling (Zolowskii et al. 2007). In natural photoperiods of light and dark, the rhythmic output gene VVD effectively modulates the WCC transcriptional response to light on the frq promoter, permitting the clock to accurately keep time during
A Molecular Output Can Directly Affect Clock Mechanisms

In the case of VVD, a gene deletion results in a largely normal clock, albeit with defects in phase control, where it is clear that the oscillator mechanism itself is not greatly perturbed. A recent and surprising example of a clock-regulated output with the ability to conditionally feed into the oscillator came through the cloning of a genetic mutation in 2006 (Elvin et al. 2005). Rhythmic control of outputs that feed back onto LOV-domain-containing inputs or oscillator components (Kim et al. 2007) may turn out to be a functionally conserved mechanism to modulate clock properties, as LOV-domain-containing photoreceptors are widely found.

Figure 4. Coupled feedback loops within Neurospora may be part of the molecular mechanism of the clock or operate to connect output to input. Molecular clock components necessary for circadian rhythmicity, FRQ (Frequency), FRH (FRQ-interacting Helicase), WC-1 (White Collar-1), and WC-2 (White Collar-2), are shown in simplified coupled feedback relationships, acting through the vvd promoter. The vvd gene, a ceg, is clock-regulated at the level of mRNA abundance. The blue light photoreceptor VVD modulates the light information coming into the clock at different times of day, resulting in circadian gating.

the day until the dusk reset (Elvin et al. 2005). Rhythmic control of outputs that feed back onto LOV-domain-containing inputs or oscillator components (Kim et al. 2007) may turn out to be a functionally conserved mechanism to modulate clock properties, as LOV-domain-containing photoreceptors are widely found.

ALTERNATIVE OSCILLATORS AND DISSECTION OF THE HIERARCHICAL ORGANIZATION OF THE CIRCADIAN SYSTEM

Oscillatory behavior is a major theme in living systems. Oscillations can naturally occur in any cellular pathway where there is negative feedback regulation with a lag. The vast majority of these oscillations do not meet the criteria for being a circadian rhythm. Much of the biochemistry of cellular metabolism can be described as feedback loops; many have short periods on the order of seconds and minutes, like feedbacks in the glycolytic oscillator, but many are also known with longer periods in the range of several hours. Often, such cycles are invisible because biochemical assays performed on a collection of cells, whether in a tissue or a dish, will report arrhythmicity if the cells are not in synchrony. Recall, for instance, that circadian rhythms in tissue culture were not found for many species that DNA damage results in cell cycle arrest (see, e.g., Harrison and Haber 2006) and that mutagens, although never for Neurospora. It is clearly established in many species that DNA damage results in cell cycle arrest (see, e.g., Harrison and Haber 2006) and that mutagens, through DNA damage, can affect the cell cycle. Although it was unknown if DNA damage could reset the circadian clock, γ-irradiation had been found to induced several important clock genes (Fu et al. 2002; Lee 2005), suggesting that this might be the case. Exposure to the radiomimetic drug methylmethane sulfonate (MMS) results in double-stranded DNA breaks and was found to reset the clock in wild type but not in the prd-4 Δ strain. When MMS was given at different times in the circadian cycle, the resulting phase-response curve (PRC) (Fig. 5) showed strong advance resetting during the subjective day in wild type but not in the prd-4 Δ strain (Pregueiro et al. 2006).

What appears to be happening is this: Whenever Chk2 becomes activated by DNA damage, one of its normal substrates is FRQ. Phosphorylation of FRQ promotes its turnover. In Chk2PRD-4, the mutation has resulted in a kinase with enhanced binding to FRQ, thereby partially bypassing the requirement for DNA damage to activate Chk2 as regards FRQ phosphorylation. As a result, in Chk2PRD-4, FRQ is always precociously phosphorylated and the circadian period length is shortened. Chk2PRD-4 is an example of a protein kinase not normally involved in operation of the clock but that regulates a clock protein in response to environmental damage to the cell.
tained oscillation in nuclear levels of the transcription factor p53, and its negative regulator Mdm2 that is seen upon γ-irradiation (Lev Bar-Or et al. 2000; Geva-Zatorsky et al. 2006). Other examples of infradian rhythms include the redox rhythmicity in yeast that has been modeled as a possible evolutionary origin of circadian rhythms (Tu and McKnight 2006).

Colin Pittendrigh and Victor Bruce (1959) first hypothesized a type of oscillation that might come under control of the circadian pacemaker, calling it a slave oscillator. They noted that “any feedback loop in the organism is a potential slave oscillator and if the circadian pacemaker can make input to the loop, the slave will assume a circadian period and become a part of the temporal program that the pacemaker drives” (Pittendrigh 1981). Are there actually slave oscillators in Neurospora or other organisms? In many organisms, regulatory relationships among factors involved in nitrate assimilation can give rise to a feedback loop, such that nitrate reductase activity is rhythmic, first shown in Gonyaulax (Ramalho et al. 1995). Possibly, the best-described putative slave oscillator is the nitrate reductase (the NIT3 protein) rhythm in Neurospora (see, e.g., Lillo et al. 2001), the metabolic activity that regulates the conversion of assimilated nitrate to ammonia and then to glutamine in a negative feedback loop. NIT3 activity is rhythmic not only in wild-type strains, but also in frq-null and probably wc-1-null strains, and interestingly, even in constant light when the FRQ/WCC is not rhythmic (Fig. 6) (Christensen et al. 2004). Preliminary work examining nitrogen reductase activity at different temperatures gives some indication that the period length of the rhythm in a wild-type strain shows temperature compensation that may then be lost in the absence of FRQ/WCC. This would suggest that coupling to the clock allows the nitrogen reductase rhythm to display circadian characteristics of period length, compensation, and phase control in clock wild-type strains.

**FRQ-LESS OSCILLATORS AND OTHER NONCIRCADIAN FEEDBACK LOOPS**

Ancillary oscillators exist in organisms without functional circadian clocks and may display some circadian properties, but do they represent part of the working clock mechanism? Genetic, molecular, and biochemical approaches aimed at understanding FRQ/WCC regulatory loops have been highly successful in providing major insights into such canonical clock oscillator features as the production and maintenance of rhythmicity, light and temperature resetting, and phase control with respect to the physical diurnal cycle. Nevertheless, within any organism, there is a complex circadian system that is thought to encompass other feedbacks that result in oscillatory behavior. Some of these will be feedback loops with functions distinct from the clock mechanism. Some of these will be clock-regulated, and some will be examples of clock-regulated output that feeds back to some aspect of the clock, either directly to mechanism or to input as discussed above for Chk2^{PRD-4} and VVD. In Neurospora, oscillators that are unmasked when the FRQ/WCC feedback loop is eliminated have been referred to as FRQ-less oscillators or FLOs (Iwasaki and Dunlap 2000). To date, several FLOs have been identified, although their importance in terms of clock function has yet to be established. Among the fungi, a number of noncircadian rhythms in development have been described (for review, see Büning 1973) in otherwise wild-type strains. An early example is the clock strain (Sussman et al. 1964, 1965), identified as producing a rhythm in growth when cultured in race tubes, but later shown to not be circadian (Feldman and Hoyle 1974).

More than two decades ago, a FLO rhythm was found in a clock-defective strain, frq^{-} (Loros 1984; Loros and Feldman 1986), that makes a truncated and nonfunctional
version of the FRQ protein (Aronson et al. 1994a). Although the period length of the FLO-driven rhythm in conidial development can range from 12 to 34 hours, it can be manipulated to occur within the circadian range because the period is highly sensitive to changes in both temperature and nutrition. Several other FLOs have since been described (see, e.g., Mattern and Brody 1979; Lakin-Thomas 1998; Merrow et al. 1999; Correa et al. 2003; Granshaw et al. 2003; Christensen et al. 2004; dePaula et al. 2006; Lombardi et al. 2007; for review, see Dunlap et al. 2004; Vitalini et al. 2006), but none display the complete set of formal circadian properties nor have they been shown to affect the operation of the FRQ/WCC feedback loop. However, FLOs may exhibit some circadian properties (see, e.g., dePaula et al. 2006). The extent to which a particular FLO might be a manifestation of the circadian mechanism revealed by the removal of some clock components is an appealing idea, particularly because the appearance of noncircadian and ultradian rhythmic behavior has been found following genetic lesion of clock genes in other systems (see, e.g., Dowse et al. 1987; Hamblen-Coyle et al. 1989; Liu et al. 2007; Storch et al. 2007).

The *Drosophila* circadian system has been modeled to contain a master oscillator that drives or entrains slave oscillators directly involved with outputs (Pittendrigh and Bruce 1959), a model that may be pertinent to *Neurospora* (Iwasaki and Dunlap 2000; Merrow et al. 2001; Dunlap and Loros 2006) and other systems. Pittendrigh speculated that the use of several slave oscillators by the core clock would allow some aspects of the system to be open to evolutionary adjustment without altering other component parts and that the slaves, because they would normally be entrained by the master, need not have all circadian properties (Pittendrigh 1981). A reasonable interpretation of FLOs is that they represent a set of Pittendrighian slave oscillators coupled to the FRQ/WCC master oscillator; when the master oscillator is removed, the slaves run on their own in various noncircadian or partially circadian modes (Loros and Dunlap 2001; Dunlap and Loros 2004, 2005, 2006). A major and untested caveat to this model is that most FLO components have not been molecularly identified and therefore cannot be compromised or deleted to test for effects on the FRQ/WCC oscillator, so their involvement in the clock system is to date unanswered. Figure 7 summarizes the currently understood interrelationships among oscillators.

The most extensively studied FLO is the asexual...
ANCILLARY OSCILLATORS CAN AFFECT THE SAME OUTPUTS AS THE CLOCK AND CAN MASK CIRCADIAN CONTROL

Work begun in the late 1970s described conidial banding rhythms in fungal strains with defects in lipid metabolism. One of these was called cel, a fatty acid chain elongation mutant that had defective circadian properties in period length and temperature compensation of the conidiation rhythm (see, e.g., Mattern and Brody 1979). Another mutant, chol-1, a morphological strain reparable by addition of choline, showed large changes in linear growth under limiting choline, manifesting as a rhythm in conidiation with extraordinarily long periods sometimes exceeding 100 hours (Lakin-Thomas 1996, 1998). Both the choline-starvation-induced and the cel rhythms, plus a recent mutation called ult (Lombardi et al. 2007), were shown to be largely independent of the FRQ/WCC oscillator. An interpretation has been that the FRQ/WCC feedback is dispensable in determining conidiation period length and therefore that this feedback loop is not required for circadian rhythmicity (Lakin-Thomas 1998; Lakin-Thomas and Brody 2000), but instead provides input into the system (Lakin-Thomas 1998, 2006b). Another explanation is that perhaps lipid manipulations can change the coupling between the FRQ/WCC and FLO oscillators (Granshaw et al. 2003; Lombardi et al. 2007). An underlying assumption in this work is that the rhythm in conidiation is a true representation of the circadian oscillator, although as discussed above, not all developmental rhythms dis-
played by Neurospora are circadian nor are they necessarily even regulated by the circadian system.

To acquire a clearer picture of the FRQ/WCC oscillator with regard to these other oscillators, it is necessary to be able to follow both in the same culture over the same time frame. In this case, the activity of the FRQ/WCC feedback needed to be monitored at the same time that the long-period, choline-limited rhythm of conidiation was being expressed. This became possible with the development of a codon-optimized luciferase that has activity in Neurospora (Gooch et al. 2007). Using the frq promoter to drive luciferase expression allows the FRQ/WCC to drive luciferase expression allows the FRQ/WCC feed-back needed to be monitored at conditions when it no longer controls conidiation. Using luciferase to follow frq expression in race tubes of the chol-1 strain under limiting choline showed the chol-1, frqP-luc strains replicating a long-period rhythm in conidiation of about 78 hours; this rhythm was frq-independent, as previously shown. A strain, chol-1, frq+, frqP-luc, carrying the long-period frq+ allele, showed a similar long-period rhythm of development on the race tube. However, when luciferase was monitored, the activity showed a clear 22-hour rhythm in the frq+ strain and a long-period, 29-hour rhythm in the chol-1, frq+, frqP-luc strain (Fig. 8). The choline-starvation-induced rhythm is known not to be temperature-compensated (Lakin-Thomas 1998), but the luciferase activity rhythm in the chol-1, frqP-luc strain demonstrated temperature compensation, equivalent to the temperature compensation of the FRQ/WCC oscillator (Shi et al. 2007). These experiments unequivocally demonstrated that the FRQ/WCC oscillator is functionally wild type under choline starvation in the chol-1 strain, but it is no longer controlling the timing or expression of asexual developmen-
t. These observations indicate that the choline-starvation-induced rhythm may reflect morphological cycles that mask or uncouple the circadian oscillator from development. It is clear, in the case of chol-1, that two distinctly different rhythms can coexist in the same culture, be of very different periodicities, and therefore not be controlled by the same oscillatory system (Fig. 7).

CONCLUSIONS

Circadian output comprises the functional end of circadian systems in terms of adaptive significance. Among fungi and animals, the regulatory logic of the core transcription-translation feedback loop is similar, and even in plants, where the core mechanism contains additional interlocked loops, a principal and primary form of output is the daily clock regulation of gene expression. For this reason, work on Neurospora, which pioneered the analysis of clock-controlled genes, continues to illuminate the pathways through which clocks can control metabolism and overt rhythms.

In addition to providing an overview of output in Neurospora, we have developed three recent stories. The first described the cloning of the bd gene and its identification as an allele of ras-1. In Neurospora, virtually every strain examined in the context of rhythms bears the bd allele that helps to clarify the overt rhythm in asexual development. ras-1bd influences a wide variety of signaling pathways and results in both enhanced light responses and increased asexual development, altering the profile of cccs in the cell and organism. Interestingly, both of these can be largely phenocopied by treatments that increase levels of intracellular reactive oxygen species.

A second story provides an example of an output from the clock that feeds back to affect circadian input. Cloning of prd-4 showed it to be an allele of checkpoint kinase-2. Normally, Chk2 is quiescent until it is activated by...
such as overt development and mask a wild-type circadian clock to bypass the requirement for DNA damage to activate this kinase; as a result, FRQ is always precociously phosphorylated, resulting in more rapid turnover and a faster cycling clock with a short-period length. In the wild-type clock, Chk2 phosphorylates FRQ in response to DNA damage in a time-of-day-dependent manner, resulting in clock-regulated phase shifting.

Finally, with improved methods for the study of single cells and the synchronization of cells in culture, more and more noncircadian or circadian-coupled feedback loops are being described. Arguably one of the most closely examined circadian cells is *Neurospora*, in which at least nine distinct oscillators have been described in addition to the circadian FRQ/WCC loop. In clock wild-type cells, many of these are normally coupled to the clock and run with circadian period lengths, but when the clock is lesioned, for instance, through mutation of *frq* or *wc* genes, these FLOs can continue to cycle on their own. In doing so, some retain circadian characteristics, suggesting the possibility that they, like the FRQ/WCC loop, contribute to the circadian oscillator. As long as the core feedback loop could be followed for long periods only through its control of the banding rhythm, it was difficult to study the FRQ/WCC loop simultaneously with other loops.

However, the recent adaptation of luciferase to function at the FRQ/WCC loop simultaneously with other loops. In clock wild-type cells, the circadian FRQ/WCC oscillator continues to compensate against changes in temperature or nutrition, 60–120-hour period length in conidia formation that is not controlled by the circadian clock. In clock wild-type cells, Chk2 phosphorylates FRQ in response to DNA damage in a time-of-day-dependent manner, resulting in clock-regulated phase shifting.

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