Combining Theoretical and Experimental Approaches to Understand the Circadian Clock

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ABSTRACT

This review is intended as a summary of our work carried out as part of the German Research Association (DFG) Center Program on Circadian Rhythms. Over the last six years, our approach to understanding circadian systems combined theoretical and experimental tools, and Gonyaulax and Neurospora have proven ideal for these efforts. Both of these model organisms demonstrate that even simple circadian systems can have multiple light input pathways and more than one rhythm generator. They have both been used to elaborate basic circadian features in conjunction with formal models. The models introduce the “zeitnehmer,” i.e., a clock-regulated input pathway, to the conceptual framework of circadian systems, and proposes networks of individual feedbacks as the basis for circadian rhythmicity.

Key Words: Neurospora; Gonyaulax; Zeitnehmer; Feedback entrain; Light.

INTRODUCTION: A BASIS FOR COMPLEXITY

Circa 24h oscillations that continue in the absence of external stimuli have been described in organisms from prokaryotes to man. These circadian rhythms are manifest at many levels, from behavior and physiology to gene expression. Although a key
experimental observation in the laboratory is self-sustained rhythmicity under constant conditions, circadian systems are almost exclusively entrained or synchronized with the environment in nature. An important relationship between free-running rhythms and the entrained circadian system is the “phase of entrainment” (Roenneberg et al., 2003). The phase refers to the time of day at which a discernable event occurs within the cycle (e.g., for birds, a marked increase in locomotor activity commences in anticipation of dawn). Entrainment refers to a stable phase relationship between biological rhythms and physical oscillations (e.g., a light cycle). Although we know that circadian systems will be more complex than mechanical oscillators, some features of the former are highly similar to the behavior of the latter (e.g., phase shifting or resetting behavior). As for a mechanical oscillator, the longer the free-running period (τ) of a biological oscillator, the later within the day will the rhythm entrain, and vice versa: earlier entrainment occurs with shorter free-running periods (hence the connection between τ and phase of entrainment). The reciprocal effect is seen when organisms with the same τ are put into longer or shorter entraining cycles (called T cycles). In shorter Ts, the entrained phase is later than in long ones (Aschoff, 1979). So, investigations into what determines the length of τ and how it is influenced by environmental factors also pertain to entrainment. Early insights into molecular aspects of circadian rhythmicity were based on extensive physiological experimentation. More recently, genetic approaches have been relied on for determining clock components. In our work on simple circadian model systems, the unicellular alga *Gonyaulax* and the filamentous fungus *Neurospora*, we have found the combination of physiology and genetics to be a powerful tool.

An initial step to unravel a biological system is to conceptualize and model it. Any circadian system will have an input pathway to transduce information from the exogenous environment to the endogenous clock. The external signals that are capable of entraining circadian systems are called zeitgebers (German for “time giver”). Zeitgebers include light, temperature, food, and social cues. Input pathways feed into a rhythm generator, responsible for the generation of the circa 24h oscillation. The last building blocks of circadian systems are output pathways, driving the rhythmic phenotypes.

The physiological and ecological description of the unicellular alga, *Gonyaulax polyedra*, has contributed greatly to our understanding of circadian programs, in general. The algae have numerous clock-controlled outputs, some of which can be monitored simultaneously. Two forms of bioluminescent expression can be recorded: a sustained “glow” at the end of the night, and much brighter, short flashes occurring throughout the night. The adaptive function of the bioluminescence has never been experimentally demonstrated, but is thought to deter predator attacks by copepods (Buskey and Mills, 1983). Other rhythmic outputs are found in, or are facilitated by, the algae’s migration behavior. During the day, the algae rise to the surface, where they harvest light for photosynthesis. At night they scavenge nitrogen from the depths; thus, energy, oxygen, carbon, and nitrogen sources are facilitated by the circadian program. At the ocean’s surface, *Gonyaulax* populations form aggregations. With the combination of a camera and a photomultiplier tube, both bioluminescence and aggregation can be recorded and quantified. When the algae are held in constant white light with sufficient nitrogen, all circadian outputs maintain a stable phase relationship with one another—they, therefore, have the same τ. When cultures are incubated in constant red light or deprived of nitrogen, the rhythms of bioluminescence and aggregation can be dissociated, and are sometimes observed as free-running with different (circadian) τ’s (Roenneberg and Morse, 1993;
Ronneberg and Rehman, 1996). Thus, at least two circadian subsystems contribute to the temporal program of this unicellular organism.

Another contribution from the Gonyaulax work is the demonstration of clock-regulated light input pathways. Using spectrally defined light and pharmacological agents, blue and blue–red sensitive light transduction pathways to the circadian system can be distinguished (Ronneberg and Hastings, 1988; Roenneberg and Deng, 1997). The blue input pathway is active only during the subjective night, facilitating large (type 0) phase shifts, while the blue–red input is active throughout the circadian cycle and leads to small (type 1) phase shifts. Thus the responsiveness and the spectral sensitivity of the light input pathways are under clock control. Using other model organisms, clock regulation of light input pathways has also been clearly shown (Bognar et al., 1999; Fleissner and Fleissner, 1996; Somers et al., 1998). In retrospect, given the extensive regulation of cellular physiology by the circadian clock, this observation is not surprising.

How do these physiological experiments contribute to an understanding of the molecular mechanism of circadian rhythms? While several of the properties and responses of circadian systems resemble those of simple mechanical oscillators, probing the Gonyaulax circadian system reveals a multioscillator and multi-input construction.

**MODELING: COMBINING TWO FEEDBACK LOOPS**

When considered together, two oscillators in a single cell and clock-regulated light input pathways are conceptually problematic. In any oscillatory system, the inputs have profound effects on phase, period, and amplitude (Glass and Mackey, 1988). Thus, alterations (e.g., a mutation) of an endogenously oscillating input pathway could have effects indistinguishable from alterations of a central component of the rhythm generator.

The problem becomes more acute in view of the molecular genetic description of the circadian clock. Mutant hunts have delivered clock genes in flies, fungi, animals, plants, and cyanobacteria. Due to the loss of clock function in clock mutant strains, these genes are generally considered to produce central components of the rhythm generation mechanism. Expression patterns of the first clock gene, *dPeriod*, crystallized the transcription/translation negative feedback loop hypothesis, whereby the protein product feeds back negatively—directly or indirectly—on its own transcription (Hardin et al., 1990). A circadian negative feedback was also experimentally probed in *Neurospora* using the clock gene *frequency* (*frq*). Induced overexpression of the gene results in loss of overt circadian rhythms as well as a decrease in RNA expression (Aronson et al., 1994b). If these clock genes function either on input pathways or within the rhythm generator, they could similarly impact circadian rhythms in mutant strains or overexpression experiments, and it would be difficult to distinguish between these possibilities.

To assess the potential of a clock gene functioning on an input pathway to a rhythm generator, we modeled an “input feedback” (Ronneberg and Merrow, 1998) based on the following assumptions: (a) feedbacks are commonly found in biological pathways (controlling responsiveness, homeostasis, or adaptation) and (b) light input pathways are commonly clock-regulated (Bognar et al., 1999; Fleissner and Fleissner, 1987). The model essentially consists of two mutually coupled feedback loops (Fig. 1). Both the feedback within the input pathway and the downstream oscillator are simple, two-component
negative feedback loops (e.g., RNA and protein of a clock gene). The parameters of the model were set so that the isolated oscillator loop is self-sustained, while the isolated input loop damps. When the components are put together, the system produces a self-sustained rhythm.

The model was used to investigate what happens to circadian qualities when the input feedback is compromised or removed. The results show that most of the properties found in clock mutants can be simulated by changing the properties of the input feedback (Roenneberg and Merrow, 1998). Namely:

1. The input feedback has a profound effect on $\tau$ and can even turn the self-sustained oscillation into a damped rhythm, eventually leading to arhythmicity.
2. Constitutive overexpression of the input component stops the rhythm.
3. Temperature compensation (different temperatures were simulated by assigning temperature-dependent rates to the different equations) is lost when the input loop is uncoupled from the oscillator (see Rensing et al., 1997; Ruoff and Rensing 1996; Ruoff et al., 1996, 1997 for further mathematical exploration of temperature-compensated circadian systems).
4. Finally, the phase response curve (PRC) based on light pulses (simulated by increasing the level of the input at discrete times) depends on the properties of the input feedback.

**EXPERIMENTS (1): THE POWER OF ENTRAINMENT**

How can theoretical models be used practically to gain insights into real circadian systems? There are indications that the two oscillators in *Gonyaulax* are targets of one or the other light input pathway (Morse et al., 1994; Roenneberg and Deng, 1997). To investigate these possibilities further, the molecular mechanisms of the two oscillators as well as the identity of the photopigments and their transduction pathways have to be elucidated. Unfortunately, *Gonyaulax* does not lend itself to molecular genetic experimentation [its scrambled genome is about 50 times larger than the human genome (Lee et al., 1993), and at this time it is not easily transformable], so it is a poor choice for further work on the molecular mechanism of the circadian system. In contrast, *Neurospora* is
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Amenable to forward and reverse genetics, and several components of the clock mechanism as well as of the light input pathway have been cloned. It has a small, (recently) sequenced genome and is haploid, so that even recessive mutations are detectable without backcrossing. In addition, \textit{Neurospora} is easy to grow, nonpathogenic, and inexpensive to maintain. Over the last three decades, \textit{N. crassa} has been among the premier molecular genetic systems for both clock and photobiology research.

Using the readout of a 22h free-running rhythm in asexual spore formation, \textit{Neurospora} was used to generate a panel of clock mutants (Feldman and Hoyle, 1973). The \textit{frq} gene was the first to be cloned (McClung et al., 1989) and, over the last 10 yrs, a molecular coregulatory, clock network has been described. On the oscillator side, \textit{frq} transcription is positively regulated by \textit{WHITE COLLAR-1} (WC-1) and \textit{WHITE COLLAR-2} (WC-2) (Crosthwaite et al., 1997), and the FRQ protein feeds back negatively on its own transcription (Aronson et al., 1994). Furthermore, FRQ has positive effects on \textit{wc-1} transcription (Merrow et al., 2001b), and on WC-1, and WC-2 protein levels (Cheng et al., 2001; Lee et al., 2000); finally, WC-1 has negative effects on wc-2 RNA (Cheng et al., 2002). On the light input side, \textit{frq} and \textit{wc-1} are rapidly induced by light (Ballario et al., 1996; Crosthwaite et al., 1995), and WC-1 and WC-2 are phosphorylated (Schwerdtfeger and Linden, 2000; Talora et al., 1999) in a light-dependent manner. Both WC proteins contain PAS domains, and a third \textit{Neurospora} PAS protein, VIVID (VVD) also has a role in circadian light signaling, e.g., on release from light to darkness, conidiation is delayed by about 4h in some—though not all—\textit{vvd} mutants (Heintzen et al., 2001; Shrode et al., 2001). A full-length, antisense \textit{frq} transcript also regulates the timing of conidiation (Kramer et al., 2003). Finally, the WC-1 protein was recently demonstrated to function as a blue-light photoreceptor (Froehlich et al., 2002; He et al., 2002), and in addition, there are a number of kinases that are critical for stability of the FRQ protein and thus the circadian, free-running period (Görl et al., 2001; Liu et al., 2000; Yang et al., 2002).

The convergence of light input and clock components makes \textit{Neurospora} an ideal model organism for testing the hypothesis of clock genes as part of an input feedback. A simple prediction of the model described above is that circadian qualities remain when the input feedback is removed, for example, by mutation of one of its genes (Roenneberg and Merrow, 1998). In such a restricted system, the circadian free-running rhythms may not be self-sustained, and entrainment by light may be lost. Remaining qualities of the circadian system could, however, be uncovered by using an alternative input pathway for entrainment, e.g., temperature. As described in the introduction, the characteristics of circadian entrainment can be experimentally demonstrated in zeitgeber cycles of different lengths (T cycles). With shorter T cycles, the phase of entrainment settles to a later time relative to the zeitgeber transitions (Aschoff and Pohl, 1978). At a certain point (at the limit of entrainment), the cycle becomes too short to entrain the system, which then free-runs through the zeitgeber cycle with relative coordination (Holst, 1939). When T becomes even shorter (\(T \approx \tau/2\)), the entrained output is seen once per two cycles, called a frequency demultiplication (Bruce, 1960). These characteristics of entrainment offer an excellent benchmark to probe for residual circadian properties in arrhythmic clock mutants. We, therefore, studied the entrainment of different \textit{frq} mutants in \textit{Neurospora} by light and temperature cycles of different lengths.

In temperature cycles, the lab \textit{Neurospora} strain (band) follows the rules of circadian entrainment (Merrow et al., 1999). The entrained phase changes systematically with T, moving progressively later as the T shortens [Fig. 2(A)]. Frequency demultiplication...
occurs in 12h cycles (e.g., one conidial band per 24h), and finally, the phase of entrainment is a function of zeitgeber strength.

Surprisingly, in light/dark (LD) cycles, none of these prototypical circadian entrainment characteristics are apparent in *Neurospora* (Merrow et al., 1999): the onset of conidiation is pegged to the L → D transition, occurring 7–8h later, regardless of the length of T [Fig. 2(B)]. “Fixed” phases of entrainment were also found for the long and short period mutants (*frq*^+) and *frq*^−, respectively), and the delay after the L → D transition was a function of the respective mutant’s τ: relative to *frq*^+ (τ ≈ 22h), onset of conidiation was earlier for *frq*^− (τ ≈ 16h) and later for *frq*^+ (τ ≈ 29h). The fixed phases were, in contrast to the results from temperature cycles, insensitive to different zeitgeber strengths, i.e., they did not change until a threshold was reached at low fluences below which the rhythms ran free. These results indicate a driven, noncircadian light-resetting behavior. There are, however, two observations indicating that synchronization by light in *Neurospora* may still be achieved via circadian entrainment: (a) After transition from constant darkness to LD conditions, transients of several days can occur until a stable, entrained state is reached (a driven response would stabilize in the first or second cycle) and (b) In T cycles close to 24h, the phase for the wild type strain was advanced by one to two hours relative to shorter cycles [see ski-shaped line in Fig. 2(B)].

**Figure 2.** Entrainment characteristics of *Neurospora crassa* in temperature and light T cycles. (A) Phases of conidiation in temperature T cycles. The gray area represents the colder (22°C), the white area the warmer (27°C) part of each cycle. Daily onsets of conidiation in both *frq*^+ (—) and the clock mutant *frq*^− (---) form parallel lines that are independent of the temperature transitions. (B) Phases of conidiation in light T cycles (400 nE m^−2 s^−1 of cool white fluorescent light and darkness, shown in gray). Unlike the systematic phase changes in temperature cycles, in light cycles the phases are uniformly 7 to 8h following lights-off, until the cycle reaches 24h (solid line; for comparison, the line for *frq*^+ in temperature cycles is replotted from panel A as a dashed line, for which the gray represents the cold incubation).
With the fundamental characteristics of entrainment described in FRQ-sufficient strains, we investigated entrainment in frq9 (a presumptive functional null mutant) and in frq10 (a knockout) (Aronson et al., 1994a; Loros et al., 1986), both of which have indistinguishable circadian phenotypes. Typically, these mutants do not show a banding rhythm in constant darkness, but there are some conditions in which they do make conidial bands about once per 24h, namely, in extra-long, 60 cm race tubes and fairly specific nutritional conditions (Aronson et al., 1994a; Loros and Feldman, 1986). Under these rhythm-permitting conditions, the frq mutation mainly leads to a large variance in τ but not to a significant change in its mean (Aronson et al., 1994a; Merrow et al., 2001a). The FRQ-less mutants were assayed for entrainment by light and temperature, respectively (Merrow et al., 1999; Roenneberg and Merrow, 2001). In race-tube conditions where a free-running rhythm is not seen in these strains, entrainment with temperature yielded phases that were indicative of an underlying rhythm generator, i.e., the timing of conidial band formation changed relative to the warm-to-cold transition according to the T [dashed line in Fig. 2(A)].

In contrast to temperature cycles, the null frq mutants failed to synchronize to LD cycles (Merrow et al., 1999), even at fluence rates up to 4 μE m⁻² s⁻¹ (1000-fold higher than that required for synchronization of conidiation in the frq-sufficient strain; Merrow and Roenneberg, data not shown). To further understand the blindness of FRQ-less strains, we used a strain where the frq ORF was transformed into the null mutants such that FRQ could be constitutively expressed (Aronson et al., 1994b). Light-regulated conidiation was only restored in these strains when frq was induced (Merrow et al., 2001b). Thus, the lack of light-regulated conidiation in FRQ-less mutants is indeed due to lack of FRQ and not to background mutations. For synchronization by light, there is apparently no need for direct or indirect light-induction of frq, as has been concluded from experiments with the frq+ strain (Crosthwaite et al., 1995; Froehlich et al., 2002). The transcriptional/translational feedback of frq RNA and protein is also dispensable for this response. These experimental results fully support the hypothesis generated by the model: a FRQ-less oscillator (FLO) remains in frq null mutants. Although important for free-running rhythmicity, the frq-FRQ feedback is dispensable for regulation of spore formation by light, as long as FRQ protein is present.

Experiments combining both light and temperature cycles indicate how multiple zeitgebers work together in the “real world” to “fine tune” entrainment. In high amplitude temperature cycles with light cycles applied in antiphase (i.e., light during the cold phase), light had relatively little effect on the phase of entrainment (Liu et al., 1998). When the amplitude of the temperature cycle was reduced to 5°C, however, the entrained phase was dominated by the light cycle (Roenneberg and Merrow, 2001).

**EXPERIMENTS (2): HOW BLIND IS BLIND?**

Subsequent experiments investigated whether the frq null mutants are completely or selectively blind. An additional light response, i.e., light-induced mycelial carotenoid production, was quantified in frq+ and frq10 strains (Merrow et al., 2001b). Both showed robust light-induced carotenoid production—even with similar sensitivities. The only difference was that carotenoid concentrations, at the same, saturating fluences, reached only half the concentration without FRQ compared to the frq+ strain. Thus, carotenogene-
sis and conidiation can either be seen as two distinct light input pathways or two distinct branches off a common light input with FRQ being essential for one but not for the other. Additional support for a separate, or bifurcated light input pathway comes from the very different sensitivities of carotenogenesis and light-regulated conidiation: the latter is 20-fold more sensitive than the former (Merrow et al., 1999; 2001b; Sargent et al., 1956). Because WC-1 is the likely photoreceptor mediating this response (Froehlich et al., 2002; He et al., 2002; Linden et al., 1999) and its levels are regulated by FRQ (Lee et al., 2000; Merrow et al., 2001b), the decreased saturation levels of carotenoids in frq mutant mycelia could be due to the lower level of WC-1 protein (Lee et al., 2000; Merrow et al., 2001b). Investigation of another light-regulated output, gene expression, in the frq null strain confirmed an intact light response, albeit with a decrease in amplitude, again attributable to low WC-1 levels (Arpaia et al., 1993; 1995; Merrow et al., 2001b).

The possibility of a dual light input pathway system suggests a reinvestigation of the hypothesis that WC-1 governs all light responses (He et al., 2002). Blindness of the wc strains for light-regulated conidiation was, so far, only based on release from constant light to constant darkness (Crosthwaite et al., 1997; Russo, 1988). This assay has, however, limited value because under standard race tube conditions wc mutants are arrhythmic in constant darkness, so that no further conidial bands can be measured. When these mutants are submitted to full photoperiod LD cycles, conidiation was entrained (Dragovic et al., 2002), clearly demonstrating that these mutants have a residual light reception, in spite of being completely blind for light-induced mycelial carotenogenesis. Subsequent experiments on knockout mutants (vs. point or RIP mutants, which were first tested) confirm that the entrainment indeed occurs in the complete absence of either WC-1 or WC-2 (Fig. 3). The evidence suggests not only two light-transduction pathways but also two photoreceptors/pigments. The recent annotation of the Neurospora genome contains numerous additional photoreceptor candidates (link to various Neurospora genome projects through http://www.fgsc.net/).

The appearance of light-regulated conidial banding in the wc mutants depends on media composition, with media containing either no carbon source or quinic acid rather than glucose giving the clearest phenotype. Qualitatively, the light-responsive conidial banding in the wc knockout mutants looks similar to those in experiments where FRQ protein is constitutively expressed from an inducible promoter in the frq mutant strains (Merrow et al., 2001). Given that FRQ is unregulated and at low levels in the wc knockout mutants, apparently the abundance of conidiation and the precision of its onset and offset suffer without regulated FRQ levels. This is not trivial, as precision is an important property of circadian systems (Roenneberg and Merrow, 1998; Pittendrigh, 1960).

REVISITING THE MODEL

Clock-controlled light input pathways appear to be a common feature in many circadian model systems (Bognar et al., 1999; Fleissner and Fleissner, 1987; McWatters et al., 2000). In Neurospora, the convergence of light input components and those involved in the circadian clock suggests a straightforward mechanism for this circadian input regulation. It also shapes how we envision the molecular mechanisms of circadian systems. In plants, light-induction of phytochrome B (phyB) RNA is gated by the clock (Bognar et al., 1999; McWatters et al., 2000), and, in turn, PHYB has effects on $\tau$ (Somers
Figure 3. Light-regulated conidiation in the wc mutant strains. Race tubes were incubated in LD 12h:12h. Single race tubes are double plotted (gray areas indicate darkness; original scans of the race tubes are shown below each plot). The strain and the delay (in h) between lights-off and conidiation-onset (entrained phase) are given on the right. Indicated phases are the average of six race tubes. The light source was cool white fluorescent light enriched for short wavelengths with solarium light (15 μE m$^{-2}$ s$^{-1}$ Osram L36, 5 μE m$^{-2}$ Osram L80).
et al., 1998). In *Neurospora*, it is possible to show time-of-day differences in the robustness of light-induced RNA responses (Heintzen et al., 2001; Merrow et al., 2001b) with stronger induction in the subjective night than in the subjective day in the case of the *wc-1* and *al-3* mRNAs. The response correlates with the amount of WC-1 protein found in tissue grown in constant darkness (i.e., more WC-1 during the night: Lee et al., 2000; Merrow et al., 2001b). The word “correlate” should be stressed here, as it is entirely possible that the amplitude of RNA induction by light could also be modulated by other factors.

The combination of modeling and molecular genetic experiments revealed two coupled feedback loops, one of which fulfills the function of a clock-regulated light input (with predictably strong effects on the free-running rhythm) while the other, on its own, still shows rudimentary circadian qualities under entraining conditions. Indeed, evidence beyond the *Neurospora* clock favors circadian systems as extended networks rather than simple loops. Within the last year, two new, interlocking feedback loops were added to the ones already known for the mammalian system (Honna et al., 2002; Preitner et al., 2002). When these numerous negative feedback loops are treated as individual components within a complex network (Roenneberg and Merrow, 2003), it becomes apparent that the circadian molecular system is difficult to explain based on straightforward genetic analyses.

Figure 4 diagrams the *Neurospora* circadian system similarly to how we have recently drawn the circadian network in mammals (Roenneberg and Merrow, 2003). As for the known clock components, FRQ, WC-1, WC-2, and VVD, only one, FRQ, has an
individual, self-regulated feedback loop. It inhibits its own transcriptional activation (Aronson et al., 1994b; Froehlich et al., 2002), and thus, in theory, given constitutive expression of its regulators, could achieve an oscillation. WC-1 protein, but not RNA, is rhythmically expressed in (DD) (Lee et al., 2000; Merrow et al., 2001b), although RNA levels can be modulated by FRQ (Merrow et al., 2001b), and a self-sustained circadian rhythm is rescued in wc-1 mutants with constitutive overexpression of WC-1 (Cheng et al., 2001). Thus, WC-1 is not placed in a feedback loop per se although its regulation is probably important for the circadian system. Constitutive expression of WC-2 also supports a free-running rhythm, and neither wc-2 RNA nor protein oscillate, so it serves as a parameter of the system. VIVID has negative effects on WC-1, and oscillates for one cycle in DD (Heintzen et al., 2001). It is not light-induced in the wc-1 knockout strain, indicating regulation by WC-1 (Lee et al., 2003). The four feedback loops in the FLO are conjectured, and are representative of some of the many feedbacks embedded in the cell’s biochemistry that must talk to one another.

The diagram is not only an alternative description of the molecular circadian machinery but also suggests distinct domains within the network: several individual feedbacks separately receive direct or indirect light input, and others participate in the generation of circadian rhythmicity that remains entrainable by temperature cycles in the absence of the light input loop(s). Another benefit of conceptualizing the network as individual feedbacks is that it highlights likely locations for new components, e.g., the transcription factors that regulate basal expression levels of wc-1, wc-2, frq, or those involved in rhythmic conidiation in temperature cycles in FRQ-less strains (the FRQ-less oscillator, FLO).

Several experimental results support the fundamental role of a feedback network as a general basis of circadian rhythmicity. Recent experimental results show that the free-running rhythmicity of some clock mutant mice is conditional, e.g., some have poorly or nonsustained circadian rhythms in constant darkness but show self-sustained rhythmic activity in constant light (Steinlechner et al., 2002; Spoelstra et al., 2002). The network concept also predicts that the elimination of one loop may disturb the network, so that its output becomes arrhythmic, while the elimination of an additional loop may reestablish rhythmicity. Recent experiments with mammalian double clock mutants show evidence for this possibility (Oster et al., 2002).

The network view of the molecular circadian system is also supported by modeling the evolution of the circadian clock with a set of simple, coupled feedback loops (Roenneberg and Merrow, 2002). Although circadian research has shown the central role of transcriptional/translational autoregulation, a network of feedbacks must have already existed at all levels of regulation in primitive, clockless cells. Feedbacks are a common control mechanism in metabolic pathways (see Roenneberg and Merrow, 1998), albeit with periods much shorter than 24h. Well-defined, transcriptional/translational feedbacks can have self-sustained periods much shorter than 24h (Hirata et al., 2002; Hoffmann et al., 2002; Elowitz and Leibler, 2000). The basic cellular biochemistry consists of a network of coupled feedback loops, including many potential, short-period oscillators. When such a network is modeled and is “stimulated” by rhythmic inputs (similar to a 24h zeitgeber), the network’s responses are chaotic, although each individual, isolated loop can be readily driven by the “zeitgeber signal.” Thus, an additional mechanism must have allowed complex cellular feedback networks to respond in an organized manner to light and darkness or warm and cold, even before a full fledged circadian system existed.
A solution to this problem is to add a zeitnehmer (German for “time taker”) function, a feedback from the output of the network onto its input pathway (Roenneberg and Merrow, 1998, 2002). Although zeitnehmer functions allow the system to respond adequately to rhythmic stimulation, the system is still not self-sustained with a circadian period under constant conditions. The evolution of a circadian system based on such a feedback network is, however, simulated in the model by changing the coupling strength between the individual feedback loops.

The implications of such a model are several. Many of the genes participating in this network may well be genes that are essential for survival, thus they would be difficult to find in mutant screens. On the other hand, if the model is correct, nonclock characteristics can be used to identify clock gene candidates that would not be found by traditional methods. For example, a mutant screen for loss of an organized response to temperature cycles could be performed either in organisms with no apparent clock (such as yeast) or in mutants with a disrupted clock.

**CONCLUSION**

Studying the circadian system with a forward genetics approach has resulted in the description of a transcription/translation feedback loop as central to the clock mechanism in all model systems. Based on their strong impact on circadian rhythms, these clock components have come to be recognized as clock genes, or “core clock component.”

The exercise of assigning circadian function to a given gene or molecule has an intellectual lineage. The first generation would be the “one-gene-one-enzyme hypothesis,” as was originally described using Neurospora (Beadle, 1945). The second generation (the molecular genetic revolution) produced the transcription/translation feedback loop, faithfully propagating the idea that the phenotype in a mutant is a direct readout of the mutation. Because circadian research has focused its efforts (using similar genetic approaches) on those genes and molecules that have profound effects on circadian rhythms, it may have tilted towards transcriptional regulation as a basis for circadian rhythmicity. The discovery that there are clearly additional domains of circadian regulation, as described here, and by work on Neurospora in other labs (Lakin-Thomas, 2000; Lakin-Thomas and Brody, 2000) may be the beginning of the third generation.

The detailed kinetic description of the known clock components and the continuing discovery of new clock components reveal an extensive coregulation. Quantitative genetic analyses have shown, for example, unexpected interactions among the components of the olfactory pathway in Drosophila (thought to represent epistatic effects (Federowicz et al., 1998)). It could be that sensory transduction pathways commonly feature interdependent regulation, and any parsimonious interpretation of genetic results would become misleading. Has the Neurospora clock evolved from two light input pathways that feed back onto each other with the consequence of circadian, self-sustained rhythmicity or has a circadian molecular network evolved to improve the processing of light information?

Nonstraight-forward genetic interactions among clock mutations have already been demonstrated in Neurospora (Lakin-Thomas and Brody, 1981; Morgan et al., 2001). This cellular system, for which uniform tissue types can easily be manipulated and analyzed, appears especially suited to study coregulatory effects. This task is a particularly difficult challenge in mice, where the circadian system of the whole animal is clearly a composite
of numerous cellular and organ clocks (Herzog and Tossini, 2001), and dissection of discrete genetic effects is confounded by their interactions. With the *Neurospora* genome annotation available, and forward and reverse genetic tools online, the use of novel mutant screens together with physiological experimentation should generate a panel of novel clock genes. The only mutant screens that have been used in *Neurospora* to date are those under constant conditions, but entrainment, as it controls appropriate temporal expression of function, is surely one of the most important circadian properties. We have started screening for mutants under entrained conditions (light or temperature cycles, both 24h and non-24h T). The first results indicate that there are many clock genes that impact the phase of entrainment in this fungus (M. Mason and K. Sveric, unpublished). Given the parallels between the clock mechanism in *Neurospora* and those of all other model systems, these novel clock genes should be relevant for identification of similar components in other circadian model systems.

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