



# Regulating behavior with the flip of a translational switch

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From the pioneering discovery of the first single-gene mutants that controlled circadian behavior by Konopka and Benzer (1), the quest to probe the fundamentally important relationship between gene expression and behavior has been ongoing. The ability to control cellular function and behavior with exquisite precision *in vivo* through genetic, optogenetic, and pharmacological manipulation already exists, but the search is still afoot for approaches to rapidly and reversibly control gene expression. Genetic code expansion (GCE) has proved to be a powerful tool to control protein function *in vitro* and inside the cellular milieu (2) and, more recently, in a wide array of vertebrates *in vivo* (3). In PNAS, Maywood et al. (4) use GCE to create a fast, conditional, and reversible translational switch to reconstitute the molecular circadian clock in the suprachiasmatic nucleus (SCN) and circadian behavior in arrhythmic mice. The rapid kinetics with which protein expression is controlled with this translational switch provides unprecedented temporal control over neuronal function *in vivo*, restoring complex behavioral responses reversibly on the timescale of about a day. This pioneering approach should be broadly applicable to many other systems, from neurobiology to the periphery, opening the door to a new era of genetic control and biological insight.

In mammals, the SCN acts as the master circadian pacemaker that provides temporal control over nearly every biological process, from the secretion of melatonin to body temperature (5). From its position in the hypothalamus, the SCN receives retinal input that is used to entrain clocks to the environmental light–dark cycle. It also provides molecular cues to synchronize the rhythms of molecular circadian clocks in peripheral tissues to globally coordinate physiological and behavioral rhythms. Therefore, manipulating circadian clocks in the SCN has powerful control over behavioral responses, making it a particularly tractable and interesting initial target for the reversible control of behavior upon reconstitution of a single gene. At the molecular level, circadian rhythms are driven by a set

of interlocked transcriptional–translational negative feedback loops. The core loop is established by the transcription factor CLOCK:BMAL1, which activates transcription of *Period* and *Cryptochrome* (*Cry*) genes that feed back to repress their expression and complete the feedback loop (6). At least one cryptochrome must be present to interact directly with CLOCK:BMAL1 (7, 8) and coordinate the assembly of large, multimeric repressive complexes (9) that close the feedback loop.

Traditional approaches via single-gene knockouts have demonstrated that *Cry1*<sup>−/−</sup> mice have a short period, while *Cry2*<sup>−/−</sup> mice have a long period (10, 11), consistent with the observation that CRY1 is a “stronger” repressor that binds to CLOCK:BMAL1 more tightly (12). Prior work from Hastings and colleagues (13) demonstrated that reconstitution of an arrhythmic SCN clock in a *Cry*-deficient mouse via adeno-associated virus delivery of a *Cry1*::EGFP fusion expressed under its own minimal promoter (*pCry1*) could reconstitute the clock with the expected long period (14). In PNAS, Maywood et al. expand upon this approach by using GCE to reversibly and rapidly control *Cry1*::EGFP in neurons to advance our understanding of the minimal determinants needed to acutely establish robust circadian rhythms.

GCE utilizes an engineered aminoacyl-tRNA synthetase and tRNA<sub>CUA</sub> pair that allows the incorporation of a noncanonical amino acid (ncAA) into a protein of interest in response to an amber stop codon (i.e., UAG). Maywood et al. used a pyrrolysyl-tRNA synthetase/pyrrolysyl-tRNA<sub>CUA</sub> pair that templated the insertion of the ncAA alkyne lysine N6-[(2-propynyloxy)carbonyl]-L-lysine (AlkK). Providing AlkK in the culture media for SCN slices *ex vivo*, or in the drinking water for mice, allowed for the rapid and reversible translation of *Cry1*::EGFP. Remarkably, this led to reconstitution of the molecular circadian clock and its resultant behavioral control on a timescale of about a day (Fig. 1). Carefully removing AlkK through a series of washouts once again eliminated *Cry1*::EGFP expression and the loss of molecular and behavioral circadian rhythms.

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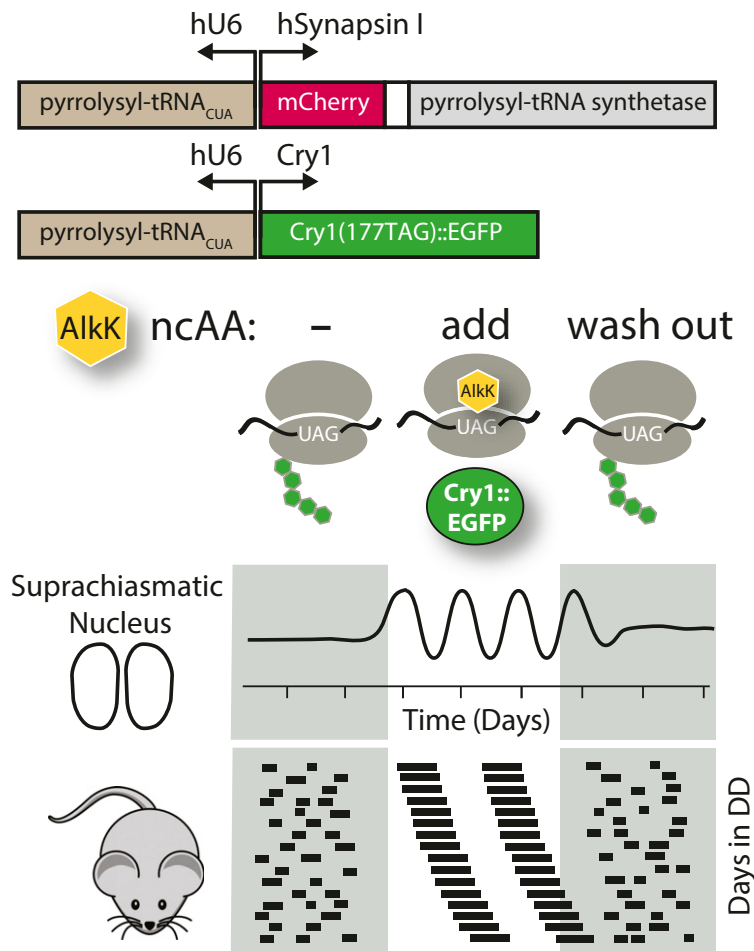
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**Fig. 1.** Reversible control of the circadian clock using translational switching of the core clock gene, *Cry1*. Using the hSynapsin promoter (Synapsin I), expression of *Cry1::EGFP* was restricted to neurons that expressed the required orthogonal tRNA synthetase (marked by coexpressed mCherry protein) and the tRNA<sub>CUA</sub> under the hU6 promoter. *Cry1::Egfp* mRNA was controlled by its own minimal promoter (*pCry1*), which directs its rhythmic transcription. Translational of *Cry1::EGFP* was strictly conditional on the presence of AlkK, an ncAA that is inserted into the protein upon readout of an amber stop codon (i.e., UAG). Circadian rhythms were monitored before and after AlkK administration with *Per2::Luc* bioluminescence in SCN explants or behavioral analysis in mice.

One powerful feature of this approach is that increasing the concentration of AlkK allowed for dose-dependent control over the expression of *Cry1::EGFP* and circadian period, which could be valuable for future studies of analysis of network perturbation by titrating clock protein stoichiometries. Additionally, use of ncAAs like AlkK opens the chemical biology toolkit by allowing for bioorthogonal labeling approaches via click chemistry (15). Here, Maywood et al. use AlkK to label nascent *Cry1::EGFP* protein with an azido fluorophore for imaging purposes. The ability to rapidly label a specific cellular population with superbright fluorophores, cross-linkers, and other chemical moieties could open the door to new studies of rare and/or transient endogenous protein complexes in situ.

GCE offers several improvements over conventional methods to manipulate protein expression. Current strategies generally target gene transcription, making use of well-studied constitutive or inducible promoters to control expression of the protein of interest. However, in many cases, changes in gene expression are permanent, such as with the use of Cre-loxP and Flp-FRT systems. Other systems, like those based on tetracycline (Tet)-controlled transcriptional control (Tet-on/off), can reversibly regulate protein expression upon addition of doxycycline. However, leaky control of gene expression and the use of orthogonal constitutive

promoters can disrupt circadian rhythms, because many clock genes need to be expressed with temporal precision that is encoded by their feedback loops (14, 16). This study therefore establishes an elegant tool that should be applicable to other systems in which transcription can remain under tight regulation, further expanding the current genetic toolkit for studying and controlling dynamic processes.

Through the use of the neuron-specific hSynapsin promoter, this work demonstrates that, despite arrhythmic behavior in the absence of *Cry* expression, the circadian network is apparently primed and ready to run upon reconstitution with the CRY protein. Given the complex network compensation that occurs upon permanent gene knockouts in the circadian network (17), the reversibility of this approach could be useful in parsing out compensation by paralogs and bringing a deeper understanding of this complicated and dynamic process. Furthermore, recent work has highlighted an exciting role for astrocyte–neuron communication in establishing circadian networks in the SCN (18, 19). Using cell type-specific promoters, GCE now opens the door to further studying the roles of intercellular communication in vivo.

In the future, it will be interesting to see whether GCE strategies could be employed to reversibly manipulate posttranslational

modifications of clock proteins. Most core components of the clock are regulated by posttranslational modifications at different points throughout the circadian cycle (20). Recent advances in the use of GCE approaches to encode site-specific phosphorylation in mammalian cells (21) tease the possibility of probing clock function at high temporal precision, thereby eliminating potential artifacts from expressing genes with constitutive mimetic substitutions.

The future looks bright: As our toolkit for the genetic code expands, so will our insight into biology.

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