Nuclear Receptors, Metabolism, and the Circadian Clock

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As ligand-dependent transcription factors, the nuclear receptor superfamily governs a remarkable array of rhythmic physiological processes such as metabolism and reproduction. To provide a “molecular blueprint” for nuclear receptor function in circadian biology, we established a diurnal expression profile of all mouse nuclear receptors in critical metabolic tissues. Our finding of broad expression and tissue-specific oscillation of nuclear receptors along with their key target genes suggests that diurnal nuclear receptor expression may contribute to established rhythms in metabolic physiology and that nuclear receptors may be involved in coupling peripheral circadian clocks to divergent metabolic outputs. Conversely, nuclear receptors may serve peripheral clock input pathways, integrating signals from the light-sensing central clock in the suprachiasmatic nucleus and other environmental cues, such as nutrients and xenobiotics. Interplay between the core circadian clock and nuclear receptors may define a large-scale signaling network that links biological timing to metabolic physiology.

INTRODUCTION

The prototypes of the nuclear receptor (NR) superfamily were identified as the steroid receptors that mediate gene transcription in response to steroid hormone signaling (Evans 1988). To date, a total of 48 human NR genes have been identified, including classic endocrine receptors for steroid hormones, thyroid hormones, and Vitamin A and D derivatives, and a large number of orphan receptors whose ligands and physiological functions were initially unknown (Giguere et al. 1988; Mangelsdorf et al. 1995; Giguere 1999). The past decade has witnessed stunning advances in orphan receptor research, largely owing to the identification of dietary lipids and metabolites as the ligands for a number of orphan receptors and establishing these adopted orphan receptors as lipid sensors that activate transcriptional programs for metabolic homeostasis (Chawla et al. 2001). For approximately half of the NR superfamily members, no ligands have yet been identified. However, their diverse roles in development, reproduction, and general metabolism have begun to be uncovered.

Many aspects of mammalian physiology are robustly rhythmic. Among these rhythmic phenomena, reproductive physiology, glucose and lipid homeostasis, and toxin clearance broadly depend on hormones and metabolites that serve as ligands for NRs. For example, periodic variations in estrogen and progesterone levels drive female menstrual cycles, whereas a morning surge in cortisol boosts energy production. A large array of metabolites, such as glucose, free fatty acids, and cholesterol and bile acids, also exhibit daily fluctuation, which is thought to be controlled by the intrinsic circadian timing system and in turn may regulate metabolic rhythms through adopted orphan receptors (Back et al. 1969; Yang et al. 2006).

The mammalian circadian timing system comprises a central pacemaker in the suprachiasmatic nucleus (SCN) of the hypothalamus and numerous peripheral tissue oscillators. The central clock is directly entrained by light from the retina via the retinohypothalamic tract, whereas the peripheral oscillators can be synchronized either by neuronal and hormonal signals from the central clock or by other environmental cues such as daily feeding/fasting or activity/rest cycles (Kohsaka and Bass 2007; Levi and Schibler 2007). This hierarchy of circadian clocks coordinates daily cycles of physiology and behavior, allowing animals to adapt to predictable changes in the environment, likely promoting fitness, health, and longevity of the organism.

In this chapter, we discuss molecular and functional links between nuclear receptors and circadian clocks, with emphasis on implications for metabolic physiology (Fig. 1).

NRs WITHIN THE CORE CLOCKWORK

Circadian clocks are self-sustained, cell-autonomous molecular oscillators. The current view of the clockwork is two interlocked transcriptional/posttranslational feedback loops comprising a battery of transcriptional activators and repressors (Hardin 2006; Ko and Takahashi 2006). A heterodimeric complex of BMAL1 and either CLOCK or NPAS2 activate the transcription of Period genes (Per1, Per2, and Per3) and Cryptochrome genes (Cry1 and Cry2) by recognizing E-box cis-regulatory elements in their promoters. Upon accumulating to a critical concentration, PER and CRY move to the nucleus and inhibit the transcription of their own genes by blocking BMAL1–CLOCK/NPAS2 activity. Clock expression is generally constant, whereas rhythmic transcription of Bmal1 is driven by a second feedback loop, involving the orphan nuclear receptors RORα and REV-ERBα (Preitner et al. 2002; Sato et al. 2004).

The ROR (α, β, γ) and REV-ERB (α and β) proteins represent closely related families of NRs that recognize similar cis response elements (ROREs) on target genes (Forman et al. 1994). RORs act as constitutive transcriptional activators, whereas REV-ERBs are constitutive...
Figure 1. Potential roles of nuclear receptors in the circadian timing system. The central clock in the suprachiasmatic nucleus (SCN) is directly entrained by light to the solar time. The clocks in peripheral tissues can be synchronized either by neuronal and hormonal signals from the central clock or by other environmental cues such as daily feeding and activity cycles. NRs may function in multiple layers of this circadian clock hierarchy. Some NRs such as RORα and REV-ERBβ are components of the core clockwork. By sensing fat-soluble hormones, vitamins, and dietary lipids in the circulation, NRs may transmit circadian signals from the central clock and/or the feeding cycle into the peripheral clocks. NRs may also act as circadian output factors to drive rhythmic physiological processes including metabolism, immune response, and reproduction.

Repressors. As a result, RORα and REV-ERBβ regulate Bmal1 transcription in an opposing manner by competitively binding to ROEs in its promoter (Preitner et al. 2002; Sato et al. 2004; Akashi and Takumi 2005). Alternate action of these two NRs over each light/dark cycle leads to rhythmic expression of Bmal1. Closing the feedback loop, BMAL1-CLOCK directly regulates the transcription of Rev-erbβ via E-box elements in its promoter (Preitner et al. 2002). The promoter of the Rev-erbβ gene also contains a functional RORE through which it is repressed by itself and induced by RORα (Delerive et al. 2002; Raspe et al. 2002). This additional layer of regulation may ensure antiphase expression of RORα and Rev-erbβ, thus enhancing the precision, robustness, or sustainability of the clock.

The functions of closely related REV-ERBβ, RORβ, and RORγ isoforms are not well understood. We found that expression of Rev-erbβ mRNA is similar to that of Rev-erbα in both tissue distribution and temporal profile (Yang et al. 2006). Although REV-ERBβ has been more intensively studied with regard to circadian rhythms and the mechanisms by which it can repress Bmal1 transcription have been described in some molecular detail (Yin and Lazar 2005), it has been reported that REV-ERBβ is capable of repressing Bmal1 transcription with a strength similar to that of REV-ERBα (Guillaumond et al. 2005). Thus, REV-ERBβ may contribute to Bmal1 expression in a manner similar and partially redundant to REVERBα. Mice with a null mutation of the Rev-erbα locus were found to have a slightly shorter free-running period in constant darkness (DD) and increased phase shifts in response to light pulses compared to wild-type controls (Preitner et al. 2002). Perhaps mice lacking both Rev-erbα and Rev-erbβ function would display a stronger disruption of circadian function. Further studies on Rev-erbα- and Rev-erbβ-deficient mouse models would illuminate the separate and distinct roles of the REV-ERB family members in the functioning of the circadian oscillator.

In contrast to the uniform diurnal expression patterns of Rev-erbα and Rev-erbβ across tissues, RORα, β, and γ isoforms exhibit diverse temporal and spatial expression patterns (Akashi and Takumi 2005; Guillaumond et al. 2005; Bookout et al. 2006; Yang et al. 2006). Although widely expressed, RORα transcripts robustly cycle in the SCN but are seemingly arrhythmic or display low-amplitude diurnal rhythms of expression in the four peripheral tissues that we examined (Ueda et al. 2002; Yang et al. 2006). Staggerer mice, which lack functional RORβ, have a decreased period of free-running locomotor activity in DD and slightly reduced Bmal1 expression in the SCN (Sato et al. 2004; Akashi and Takumi 2005). The expression of other core clock genes is unaffected. This modest phenotype may be due to partial compensation by RORγ, which is also highly expressed in the SCN with a circadian rhythm of expression similar to that of the Period gene transcripts (Sumi et al. 2002). Genetic disruption of RORβ increases the free-running period in mice (Andre et al. 1998; Masana et al. 2007). RORγ mRNA is highly expressed in the periphery and cycles in selective tissues (Yang et al. 2006). Mice lacking functional RORγ have not been described.

The nuclear receptor coactivator PGC1α, which was originally cloned as a cofactor of the PPAR family of NRs and has been shown to bind and regulate the activity of multiple NRs (Puigserver et al. 1998; Knutti et al. 2000), was recently shown to coactivate RORα and RORγ on the Bmal1 promoter RORE (Liu et al. 2007). PGC1α−/− mice seem to have a slightly increased free-running period in DD, although it is unclear whether this result is statistically significant (Liu et al. 2007). In contrast, RORα mutant Staggerer mice have a slightly decreased free-running period of locomotor activity, suggesting that a mild effect of PGC1α loss on free-running period would not be due primarily to loss of coactivation of RORα but may result from the loss of coactivation of RORβ, RORγ, or other NRs. PGC1β−/− mice exhibit decreased nocturnal locomotor activity under light/dark conditions, which may be due to a disturbance of the circadian system (Sonoda et al. 2006).

An earlier report suggested a role for the nuclear receptor Ear2 (also known as COUP-TFIII) in the maintenance of circadian rhythmicity because Ear2−/− mice were found to have increased error in the timing of locomotor activity onsets and their behavioral rhythms degraded in response to lower intensities of constant light (LL) compared to wild-type controls (Warnecke et al. 2005). However, the same animals exhibited distorted architecture of several neuronal structures, raising the concern that their disrupted circadian behaviors may be due to developmental defects rather than to disruption of the circadian clock in adult animals. Conditional deletion of Ear2/COUP-TFIII
in adult neurons would resolve this question. In our study, transcripts encoding all three members of the COUP-TF family of NRs were expressed arrhythmically in all tissues examined (Yang et al. 2006).

Transcripts encoding the highly related nuclear receptors NGFI-B, NOR1 and NURR1 (also known as NR4A1, NR4A2, and NR4A3) displayed striking high-amplitude diurnal patterns of expression in most tissues examined in our study. The Ngfi-b transcript was also reported to be strongly induced by light in the hamster SCN (Morris et al. 1998), which led to the hypothesis that NGFI-B may have a role in either light entrainment or maintenance of circadian rhythmicity. However, Ngfi-b–/– mice did not exhibit any disruption of circadian locomotor behavior under constant conditions and displayed normal behavioral phase shifting in response to a variety of light stimuli (Kilduff et al. 1998). Because there are three highly homologous receptors in the NR4A family and they have similar diurnal rhythms of expression, they are likely redundant for some functions, possibly including circadian clock entrainment. The function of the ubiquitous, strongly diurnal expression of the NR4A receptors remains unexplained.

NURS MAY MEDIATE ENTRAINMENT OF PERIPHERAL CLOCKS BY METABOLIC SIGNALS

Elegant experiments within the last decade established that fasting and feeding patterns are the primary determinant of the timing of peripheral circadian clocks (Damiola et al. 2000; Stokkan et al. 2001). However, the mechanisms by which feeding time sets peripheral clocks remain obscure and may include both neural and humoral components. Circulating factors that are involved in the entrainment of peripheral clocks by feeding time would be expected to display robust diurnal rhythms in the circulation that are altered by changing the time of restricted feeding. Many NR ligands meet these criteria, including glucocorticoids, triiodothyronine (T3), thyroxine (T4), retinoic acid, and dietary lipids. One or many of such glucocorticoids, triiodothyronine (T3), thyroxine (T4), retinoic acid, and dietary lipids. One or many of such molecules may determine the timing of all peripheral clocks; alternatively, unique factors or combinations of factors, possibly including one or more NR ligands, may entrain various peripheral organ clocks.

Glucocorticoids

Circulating glucocorticoids display strong diurnal rhythms, and glucocorticoids have been shown to induce the expression of circadian transcripts in cultured cells (Balsalobre et al. 2000a,b), which made these steroid hormones attractive candidates for peripheral entrainment cues. However, recent evidence argues against their role as a major time cue for peripheral clocks (Le Minh et al. 2001). Similar detailed analysis of the potential for other NR ligands to entrain peripheral circadian clocks has not yet been done and would be required to make conclusions about their physiological roles in peripheral clock entrainment.

Thyroid Hormones

Thyroid stimulating hormone (TSH), T3, and T4 are robustly rhythmic in the circulation in rodents, and the circadian rhythm of TSH has been shown to be driven by circadian clock function in human subjects (Allan and Czeisler 1994). Furthermore, the phases of T3 and T4 rhythms are completely reversed by reversed-phase restricted feeding (Ahima et al. 1998). Finally, thyroidectomy alters the diurnal expression of clock genes outside of the SCN (Amir and Robinson 2006).

Retinoic Acid

Retinoic acid is the ligand for heterodimeric NR complexes containing RAR and RXR. RAR and RXR associate with CLOCK and NPAS2 (also known as MOP4) in a hormone-dependent fashion (McNamara et al. 2001). The same study also found that retinoic acid could inhibit CLOCK-mediated transcription both in cultured muscle cells and in cardiovascular organs from intact animals. Moderate phase shifts of peripheral clocks in the heart and aorta were observed after injection of retinoic acid. An independent screen tested 299 peptides and bioactive lipids for the ability to mediate entrainment of circadian rhythms in cultured fibroblasts expressing Per2-luciferase. Of the 12 targets active in their screen, 3 were all trans-retinoic acid, 9-cis retinoic acid, and 13-cis retinoic acid (Nakahata et al. 2006). Moreover, there is evidence that retinoic acid signaling might be involved in light sensing and the central clock resetting (Thompson et al. 2004; Fu et al. 2005).

PPAR Ligands

The intestinal synthesis and degradation of oleylethanolamide (OEA) occur diurnally and are regulated by food intake (Fu et al. 2003, 2007). OEA is an endogenous ligand for PPARα and inhibits appetite via PPARα activation (Fu et al. 2003), presumably at hypothalamic sites. The timing and regulation of OEA synthesis and degradation are consistent with a potential role in the entrainment of peripheral circadian clocks. PPARα regulates the transcription of both Bmal1 and Rev-erba via PPRE cis-regulatory elements in their promoters (Canaple et al. 2006), suggesting a potential mechanism of contribution to peripheral clock entrainment. In addition, the endogenous PPARγ ligand 15-deoxy-Delta12,14-prostaglandin J2 was another of the 12 candidates found to entrain circadian rhythms in cultured fibroblasts by real-time monitoring of Per2-luciferase rhythms, further emphasizing the potential of dietary responsive NRs to influence the clock (Nakahata et al. 2006).

Non-NR Ligands

It is also possible that NRs participate in the entrainment of peripheral clocks in response to nutrient-derived signals that are not direct NR ligands such as glucose and insulin. Degradation of REV-ERBα protein is blocked by GSK3-mediated phosphorylation (Yin et al. 2006). Because...
GSK3 activity is inhibited in response to acute bouts of feeding by insulin-stimulated AKT-mediated phosphorylation, degradation of REV-ERβα in response to feeding may contribute to the entrainment of peripheral clocks.

**NRs AS CIRCADIAN EFFECTORS OF METABOLISM**

Our survey of the diurnal expression profile of all 49 mouse nuclear receptors reveals wide expression and tissue-specific oscillation of NRs in a variety of metabolic tissues (Yang et al. 2006). From this analysis, we are not able to determine which cycling NR transcripts are driven directly by peripheral circadian clocks and which are responsive to physiological rhythms or to secondary clock-driven transcription factors. Nor did we measure NR protein levels and are thus unable to say which NRs are rhythmic at the protein level. However, there is evidence that several NR transcripts are directly regulated by CLOCK/BMAL1, and a few NR protein rhythms have been described that closely follow the rhythmic expression of the corresponding transcripts. The dynamic and coordinated changes in NR expression along with expression of their key target genes suggest that NRs may contribute to the regulation of divergent metabolic readouts by peripheral circadian clocks.

The NR transcripts most likely to be directly regulated by peripheral circadian clocks are those with peak expression at ZT 4 or ZT 8 in peripheral metabolic tissues, when CLOCK/BMAL1 heterodimers are most active in those tissues. The rhythmic expression of NR transcripts that oscillate in a tissue-specific manner may be driven by CLOCK-BMAL1 but may require additional cofactors that are expressed in a tissue-specific manner. We discuss a few examples of NRs that appear most likely to be directly controlled by peripheral circadian clocks and that would be expected to have a large impact on metabolic physiology, thus potentially linking peripheral circadian clocks to physiological rhythms.

**PPARs**

The roles of the PPAR family of transcription factors in various aspects of mammalian metabolic physiology are well established (Lee et al. 2003; Evans et al. 2004). Although the amplitudes of their diurnal expression patterns in some of the tissues that we examined are low, the rhythmicity of at least PPARα is likely to be physiologically relevant. Both the PPARα transcript and protein oscillate in mouse liver with a time of peak expression consistent with regulation by local circadian clocks (Lemberger et al. 1996). Pparα transcription can be directly regulated by CLOCK and BMAL1 in vitro and in the liver and intestine in vivo (Inoue et al. 2005; Oishi et al. 2005; Canaple et al. 2006). As discussed above, the endogenous PPARα ligand OEA is synthesized diurnally in the gut epithelium, probably in response to nutrient availability. Perhaps the rhythmic expression of PPARα in the liver synergizes with OEA production to increase the amplitude of PPARα activity on its target promoters. This mechanism may also have a role in the diurnal regulation of appetite, as OEA activation of PPARα in the central nervous system (CNS) is a satiety signal. We did not measure the transcription of NRs in the CNS, so it is not clear whether a similar amplification of the signal is likely there.

**CAR**

The so-called constitutive androstane receptor (CAR) would be an interesting candidate for circadian regulation as it is a potent modulator of xenobiotic metabolism (Qatanani and Moore 2005). A recent elegant study showed that transcription of CAR is regulated by the PARbZIP family of transcription factors, including DBP, HLF, and TEF, which in turn are regulated by CLOCK:BMAL1-dependent transcription and indeed are among the transcripts with the highest amplitude of circadian transcription in many organs (Gachon et al. 2006). In control animals, CAR expression peaks in the early night (ZT 12), whereas compound null mutations of Dhp, Hlf, and Tef result in loss of diurnal expression of CAR and its target genes, including many cytochrome family enzymes involved in the clearance of exogenous compounds by the liver. Physiologically, Dhp−/−; Tef−/−; Hlf−/− animals have increased liver weight and reduced tolerance for anesthetic and chemotherapeutic agents. The combined diurnal expression of CAR with peak expression during the night phase when mice are actively ingesting food and diurnal sensitivity to exogenous agents that are both lost upon loss of the PARbZIP transcription factors suggests that these proteins have a critical role in optimizing circadian timing of toxin clearance to the phase of food ingestion.

**SHP**

We found that the Shp (small heterodimeric partner) transcript is robustly rhythmic in the liver but is not detectably expressed in adipose tissues or skeletal muscle (Yang et al. 2006). The SHP promoter contains E boxes and can be directly activated by CLOCK and BMAL1 (Oiwa et al. 2007). Furthermore, the ability of CLOCK and BMAL1 to drive transcription from the SHP promoter was found to be increased more than fivefold by coexpression of the nuclear receptor LRH-1 (Oiwa et al. 2007), which we found to be highly expressed in the liver compared to the other tissues examined, probably accounting for the tissue-specific oscillation of SHP transcription that we observed.

SHP is an orphan nuclear receptor that dimerizes with other NRs, including LXRα, FXR, and PXR, and represses their activities. Multiple NR partners of SHP have been shown to regulate the transcription of Cyp7a1 and Cyp8b2, among other transcripts involved in metabolizing dietary lipids and exogenous toxins (Schoonjans and Auwerx 2002). The loss of SHP function alone is sufficient to significantly increase the expression of Cyp7a1 and Cyp8b2 in the liver. Furthermore, when Shp−/− mice are fed a diet high in cholesterol and/or cholic acid, their expression of detoxifying enzymes is strikingly higher than in control animals under similar conditions and they avoid the hepatic steatosis suffered by control animals (Wang et al. 2003). Taken together, these results imply that the liver-specific rhythmic expression of SHP is prob-
ably driven by local circadian clocks and is expected to drive rhythmic repression of multiple NRs, thus contributing to daily rhythms in clearance of excess dietary lipids and other toxins.

**COMPLEXITY AND SPECIFICITY OF CIRCADIAN REGULATION BY NRs**

In metabolic pathways, certain circadian responsive target genes can be subject to direct regulation by multiple NRs and other factors. For example, evidence suggests that the Cyp7a1 gene is regulated by at least six NRs: LXRα, LRH-1, FXRα, PXR, RXRα, and SHP (Chawla et al. 2000; Schoomjans and Auwerx 2002). Conversely, individual NRs participate in gene regulation in diverse cellular pathways. For example, PPARγ in adipocytes regulates genes involved in lipogenesis and lipid storage, glucose uptake, energy expenditure, and adipokine production (Lee et al. 2003; Evans et al. 2004). Furthermore, crosstalk within the NR superfamily has been widely documented. In the liver, the Shp gene can be regulated by FXRα, LHRH-1, and ERRγ (Lu et al. 2000; Sanyal et al. 2002), whereas the Rev-erba gene can be regulated by RORα, PPARα, PPARγ, and LXR (Gervois et al. 1999; Raspe et al. 2002; Fontaine et al. 2003). These NRs also interact functionally with other classes of transcriptional regulators such as SREBP-1c (Repa et al. 2000; Chen et al. 2004). Taken together, we propose that a large pool of NRs in any given peripheral tissue comprises an interlaced transcriptional network that coordinates multiple metabolic pathways in response to circadian and other cues.

In this complex regulatory network, how does the oscillation of an array of NRs give rise to specific rhythmic outputs? Several mechanisms could be involved. First, as suggested by our results and others (Panda et al. 2002), NRs appear to preferentially target rate-limiting genes in a metabolic pathway for circadian regulation. Second, for a battery of NRs that target a single gene, cycling of a minimal number of nuclear receptors seems to be sufficient to boost the oscillation of the target gene. This notion is supported by our observation that, among six nuclear receptors that are known to regulate Cyp7a1, only SHP is rhythmically expressed and appears to be the primary contributor to Cyp7a1 cycling. Third, for NRs that act as heterodimers, cyclic expression of one subunit may be sufficient for periodic changes in the heterodimer activity. Because RXR serves as the partners for many other NRs (Mangelsdorf and Evans 1995), any dramatic changes in their levels would be detrimental. Indeed, we found that all RXR subtypes are continually expressed at fairly constant levels. In the absence of the cyclical induction of an RXR ligand, this places the critical regulation of the partner and/or its partner ligand; here, the PPARs may be one example.

**MULTIPLE LOOPS BETWEEN NRs AND THE CORE CLOCK**

In the above sections, we have depicted a scheme in which various NRs may be involved in circadian input and output pathways, as well as in the core clock mechanism. Feedback regulation is implicated at many points in circadian signal transduction. Not only does the core clock involve double interlocked feedback loops (Glossop et al. 1999; Shearman et al. 2000), but this type of feedback may also regulate communication between the core clockwork and input/output pathways. For example, PPARγ has been proposed to serve the input pathway by promoting Bmal1 and Rev-erba expression (Gervois et al. 1999; Canaple et al. 2006). Conversely, BMAL1 and CLOCK directly regulate Pparα transcription (Inoue et al. 2005; Oishi et al. 2005; Canaple et al. 2006). Xenobiotic metabolism is an important output of the circadian clock and xenobiotic compounds may, in turn, act as NR ligands to alter core clock gene expression (Claudel et al. 2007). We postulate that NRs and the core clock components are integrated into multiple feedback loops, which constitute a large-scale signaling network.

The signaling network connecting NRs and circadian clocks may serve a number of functions. First, as the double loops in the core clock are likely vulnerable to stochastic perturbation, additional loops involving NRs may improve the precision and robustness of peripheral clocks’ 24-hour oscillations. In addition, this NR-clock network may sense a broad range of external cues, such as light, diet, and stress, by hormonal mechanisms, and integrate multiple input signals. In addition, NRs in this network regulate a variety of outputs that may link the core clock to diverse molecular genetic programs, thus orchestrating metabolism and physiology over the light/dark cycle. Finally, nearly half of the NR superfamily, including members of the ROR, REV-ERB, COUP-TF, NGFI-B, RAR, RXR, TR, GR, PPAR, and ERR subfamilies, are ubiquitously expressed in all tested tissues (Bookout et al. 2006); other NRs are expressed either constitutively or diurnally in a tissue-specific manner. By regulating tissue-specific transcriptional programs, those NRs may create local versions of the circadian network.

**CONCLUDING REMARKS**

During the last decade, remarkable progress has been made in uncovering the molecular basis of the core circadian clockwork. However, we are just beginning to understand its biological inputs and outputs at a level ranging from gene expression to physiology and metabolism. In this chapter, we discussed evidence suggesting that the NR superfamily might constitute critical signaling cascades involved in coupling the circadian clock to divergent physiological outputs as well as in the clock entrainment by various zeitgebers. We have proposed that a conserved large-scale circadian network may emerge from feedback regulation between the NR and core clock genes. However, the existence and architecture of such NR-clock signaling network have yet to be determined. Although rhythmic expression is prevalent in the NR superfamily, it remains unknown which NRs (if any) are involved in peripheral clock resetting, which are solely responsive to the ticking output of the clock and which can manifest a diurnal rhythm as a result of other oscillatory cues.

Efficacy of NR signaling depends on the availability of receptors as well as their ligands. Hormonal ligands are
usually produced in specific tissues such as the thyroid and adrenal glands and are delivered to target tissues via the circulation, although some active ligands can be generated in local tissues. Important questions yet to be answered include how the circadian clock affects endocrine and locally produced autocrine ligands and how ligand cycling is correlated with receptor cycling in regulation of rhythmic physiological processes. Furthermore, it remains elusive whether feeding/fasting cycles give rise to cyclic accumulation of dietary lipids and metabolites in the body, which would serve as ligands to stimulate the activity of adopted orphan receptors in a diurnal manner, thereby inducing cyclic expression of their target genes.

Understanding the mechanisms by which cell-autonomous clocks are in sync at the tissue, organ, and system levels is a major challenge in the field of circadian biology. A plausible model is that, by sensing light, the central clock dictates rhythmic activity of the endocrine system which in turn entrains whole-body physiology to the light/dark cycle. Conditions of restricted feeding may overcome such SCN-driven entrainment by directly altering some endocrine functions. Nuclear hormone receptors represent a family of candidates that may be involved in the synchronization mechanisms of individual organs. Overall, the NR signaling system and the circadian timing system appear to be integrated at many levels (see Fig. 1), and it will be interesting to further elucidate their connections as we move toward understanding circadian inputs and outputs in greater detail.

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