Novel approaches for assessing circadian rhythmicity in humans: A review

Abstract:
Temporal organisation of molecular and physiological processes is driven by environmental and behavioural cycles, as well as by self-sustained molecular circadian oscillators. Quantification of phase, amplitude, period, and disruption of circadian rhythms is essential for understanding their contribution to sleep-wake disorders, social jet-lag, inter-individual differences in entrainment and the development of chrono-therapeutics.

Traditionally, assessment of the human circadian system, and the output of the SCN in particular, required collection of long time series of univariate markers such as melatonin or core body temperature. Data were collected in specialised laboratory protocols designed to control for environmental and behavioural influences on rhythmicity. These protocols are time-consuming, expensive, and are not practical for assessing circadian status in patients or in participants in epidemiologic studies.

Novel approaches for assessment of circadian parameters of the SCN or peripheral oscillators have been developed. They are based on machine learning or mathematical model-informed analyses of features extracted from one or a few samples of high dimensional data such as transcriptomes, metabolomes, long term simultaneous recording of activity, light exposure, skin temperature, and heart rate, or in vitro approaches. Here, we review whether these approaches successfully quantify parameters of central and peripheral circadian oscillators as indexed by gold standard markers. While several approaches perform well under entrained conditions when sleep occurs at night, the methods either perform worse in other conditions such as shift work, or they have not been assessed under any conditions other than entrainment and thus we do not yet know how robust they are.

Novel approaches for the assessment of circadian parameters hold promise for circadian medicine, chrono-therapeutics, and chrono-epidemiology. There remains a need to validate these approaches against gold standard markers, in individuals of all sexes and ages, in patient populations, and, in particular, under conditions in which behavioural cycles are displaced.
Novel approaches for assessing circadian rhythmicity in humans: A review

Derk-Jan Dijk PhD\textsuperscript{1,2} and Jeanne F. Duffy PhD\textsuperscript{3}

\textsuperscript{1} Surrey Sleep Research Centre, Faculty of Health and Medical Sciences, University of Surrey, Guildford GU27XP UK

\textsuperscript{2} UK Dementia Research Institute at the University of Surrey

\textsuperscript{3} Division of Sleep and Circadian Disorders, Departments of Medicine and Neurology, Brigham and Women’s Hospital, Boston, MA, USA; Division of Sleep Medicine, Harvard Medical School, Boston, MA, USA

Short title: Novel approaches for assessing circadian rhythmicity

Correspondence: D.J.Dijk@surrey.ac.uk

Keywords: biomarkers, machine learning, transcriptomics, data science, mathematical models, light, heart rate, skin temperature
Abstract

Temporal organisation of molecular and physiological processes is driven by environmental and behaviour cycles, as well as by self-sustained molecular circadian oscillators. Quantification of phase, amplitude, period, and disruption of circadian rhythms is essential for understanding their contribution to sleep-wake disorders, social jet-lag, inter-individual differences in entrainment and the development of chrono-therapeutics.

Traditionally, assessment of the human circadian system, and the output of the SCN in particular, required collection of long time series of univariate markers such as melatonin or core body temperature. Data were collected in specialised laboratory protocols designed to control for environmental and behavioural influences on rhythmicity. These protocols are time-consuming, expensive, and are not practical for assessing circadian status in patients or in participants in epidemiologic studies.

Novel approaches for assessment of circadian parameters of the SCN or peripheral oscillators have been developed. They are based on machine learning or mathematical model-informed analyses of features extracted from one or a few samples of high dimensional data such as transcriptomes, metabolomes, long term simultaneous recording of activity, light exposure, skin temperature, and heart rate, or in vitro approaches. Here, we review whether these approaches successfully quantify parameters of central and peripheral circadian oscillators as indexed by gold standard markers. While several approaches perform well under entrained conditions when sleep occurs at night, the methods either perform worse in other conditions such as shift work, or they have not been assessed under any conditions other than entrainment and thus we do not yet know how robust they are.

Novel approaches for the assessment of circadian parameters hold promise for circadian medicine, chrono-therapeutics, and chrono-epidemiology. There remains a need to validate these approaches against gold standard markers, in individuals of all sexes and ages, in patient populations, and, in particular, under conditions in which behavioural cycles are displaced.
Introduction

Assessing the phase, period, and amplitude of circadian oscillators is central to the study of circadian rhythms, be it in cyanobacteria, mice, or humans (Kuhlman et al., 2018). Accurate phase measurements enable description of the phase relationship (relative timing) of circadian oscillations with the environmental and behavioural cycles in the study of entrainment in humans (Duffy et al., 1999) (Wright et al., 2005). Accurate phase assessments are also a prerequisite to describe the interrelations between circadian oscillators in any multi-oscillator system, be it hierarchically organised or not (Honma, 2018) (Dijk and Lockley, 2002). Accurate assessments of period and amplitude enable identification of mechanisms underlying abnormal entrainment or lack of robustness in circadian regulation of physiological processes. Developing accurate and unobtrusive methods to assess period, phase, amplitude, robustness, and disruption is critical to understand the role of circadian rhythms in physical and mental health and their disorders. Traditional human circadian rhythm research areas include shift-work, jet lag, and circadian rhythm sleep-wake disorders (Sack et al., 2007a). More recently, phenomena like social-jet lag and applications like chrono-therapeutics and chrono-medicine have gained attention, and the epidemiology of circadian disruption is an emerging area of interest (Roenneberg and Merrow, 2016)]. Methods that can accurately assess circadian parameters and be implemented at scale and at low cost are critical for the translation of basic circadian rhythm research to all these areas (Cederroth et al., 2019) (Mullington et al., 2016) (Munch and Kramer, 2019). Recent years have seen the introduction of novel approaches to the assessment of circadian parameters and, in particular, circadian phase in humans. Some of these methodologies use machine learning approaches to extract features that predict circadian parameters from high dimensional ‘omics’ data, whereas others are based on multiple behavioural, environmental and physiological variables collected from research or consumer grade wearables, combinations of mathematical modelling and wearable-based data acquisition, or analyses of in vitro circadian behaviour in human cell cultures. Here we will revisit (Duffy and Dijk, 2002) some of the issues, pitfalls, and requirements for the assessment of circadian parameters in humans and discuss some of the novel approaches within that context.

Diurnal rhythmicity, circadian rhythmicity, endogenous circadian components, behavioural masking

Rhythmicity may be observed in any physiological or behavioural variable and quantification of this rhythmicity may be of intrinsic interest. However, often rhythmicity is assessed not for its own sake, but to quantify characteristics of an underlying circadian oscillator, which is assumed to drive the rhythmicity being assessed. A central concept in circadian rhythm research is that an overt 24-h diurnal rhythm can only be considered a circadian rhythm when it persists in the absence of masking by external environmental and behavioural 24-h cycles (Kuhlman et al., 2018). Many laboratory experiments in which organisms, including humans, were studied while shielded from 24-h environmental cycles have demonstrated that at least part of overt rhythmicity is driven by endogenous circadian oscillators. However, those studies have also revealed that 24-h environmental and behavioural cycles also contribute to 24-h rhythmicity in many aspects of physiology. This distinction between diurnal rhythmicity (due to 24-h rhythms in the environment and in behaviour) and circadian rhythmicity (due to endogenous processes) is critical to take into account when developing and applying methods for the quantitative assessment of circadian rhythmicity. Unfortunately, this important distinction is often overlooked, and we believe it has hampered progress in developing novel methods for assessing circadian timing and in understanding the role of circadian oscillators in disease (Lyall et al., 2018).
In the context of the distinction between \textit{diurnal} and \textit{circadian} rhythmicity, human circadian rhythm researchers also make a distinction between \textit{endogenous} circadian components of a rhythm and \textit{evoked} components. This distinction relates to the concept of behavioural masking and implies that assessment of the endogenous circadian component of a rhythm not only requires that components driven (masked) by environmental cycles are removed, but also that components driven (masked) by behavioural cycles must be controlled (Rietveld et al., 1993). It has now become clear that behavioural and environmental masking extends to rhythmicity at the molecular level. Thus, rhythmicity in the brain and blood transcriptome is to a large extent driven by the timing of the sleep-wake cycle (Maret et al., 2007) (Hor et al., 2019) (Archer et al., 2014) (Archer and Oster, 2015) and is affected by insufficient sleep (Moller-Levet et al., 2013) (Laing et al., 2019b). Likewise, rhythmicity in the liver is to a large extent driven by feeding rhythms and influenced by light (Atger et al., 2015) (Greenwell et al., 2019; Koronowski et al., 2019). Protocols such as the constant routine or forced desynchrony, which eliminate or distribute masking uniformly across the circadian cycle, were developed to control this behavioural masking so that aspects of rhythmicity driven ‘directly’ by circadian clocks could be quantified. These protocols have been used primarily in human studies and the confounding effects of behavioural cycles, and the rest-activity cycle in particular, have often been ignored in animal studies.

Novel circadian biomarkers must be robust to altered environmental and behavioural influences for them to be useful in patients with circadian rhythm disorders, in people who do shift work or have recently travelled to another time zone, and in individuals with social jet lag or who keep irregular sleep schedules. Therefore, in our evaluation of novel methods to quantify circadian rhythmicity, we consider the extent to which they can distinguish between environmental, behavioural, or endogenous circadian components of overt rhythmicity.

A biomarker for circadian phase, amplitude, and period of which oscillator?

Now that it has been established that circadian oscillators are present in every tissue, organ, and cell, it is more important than ever to be explicit about the circadian oscillator of interest when assessing a rhythmic output (Mohawk et al., 2012) (Mure et al., 2018) [Fig 1]. Traditionally, and particularly so in research on circadian rhythm sleep-wake disorders, there has been a focus on assessing parameters of the ‘master circadian pacemaker’ located in the SCN (Sack et al., 2007a) (Sack et al., 2007b). Obviously, there is more to the human circadian system than the SCN. Cardiologists may be interested in the phase, period, or amplitude of circadian oscillators in the heart (Thosar et al., 2018), or how the timing of anti-hypertensives impact blood pressure control (Smolensky et al., 2017); clinicians or researchers of metabolism may want to assess circadian parameters in the pancreas or adipocytes (Qian and Scheer, 2016). In fact, insulin sensitivity rhythms, glucose rhythms, and many other rhythms related to metabolism have already been characterised in humans (Poggiogalle et al., 2018). Immunologists may be interested in the extent to which the various white blood cell types vary across the diurnal cycle (Pick et al., 2019), how molecular processes related to immune function oscillate within each of these cell types, and how far these intracellular or intercellular rhythms are directly modulated by local circadian clocks (Baxter and Ray, 2019) (Downton et al., 2019). Oncologists are interested in rhythmicity in tumours and how circadian phase assessment may allow the most effective timing of chemo- or radio-therapies (Shafi and Knudsen, 2019) (Shuboni-Mulligan et al., 2019). Neurologists and psychiatrists are interested in circadian rhythms in mood, seizures, and neurodegeneration (Logan and McClung, 2019) (Khan et al., 2018; Leng et al., 2019) (Pavlova et al., 2009) (Lucey et al., 2017). Whether these rhythms in heart rate, glucose, insulin sensitivity or leukocytes are driven by local tissue clocks and/or through central control from the SCN, and to what extent these rhythms reflect...
‘endogenous’ circadian rhythms, or diurnal rhythmicity, or instead are driven by rhythmic
behaviours, is not easily established. Nevertheless, novel methods and biomarkers to quantify
rhythmicity and biomarkers may facilitate the characterisation of tissue-specific local oscillators,
allowing for increased understanding of many normal and pathological physiological processes and
the application of chronotherapies targeting specific organs and tissues.

Standard parameters of circadian oscillators and the rhythms they generate are phase and intrinsic
period, but aspects such as amplitude and waveform of rhythms can also be used to characterise
rhythms, although the latter two are rarely assessed.

**Traditional peripheral markers for SCN phase and period**

The SCN drives many daily rhythms, including those observed in the autonomic nervous system
(Buijs et al., 2013), endocrine rhythms(Czeisler and Klerman, 1999) (Morris et al., 2012), and in
particular the circadian component of the sleep propensity rhythm (Dijk and Czeisler, 1995). The
desire to assess characteristics of the circadian pacemaker located in the SCN arose from hypotheses
predicting that changes in endogenous circadian parameters were the cause of changes in sleep
timing, such as observed in ageing, circadian rhythm sleep-wake disorders, or between chronotypes
(Sack et al., 2007b). Diagnosis and understanding of the aetiology of circadian rhythm sleep-wake
disorders may be informed by accurate assessment of the SCN’s intrinsic period, amplitude, and/or
phase. This desire to know about the status of the SCN pacemaker is also driven by the recognition
that effects of sleep-wake therapeutics, be it light treatment or melatonin administration, are
dependent on the circadian phase at which they are administered (Duffy and Wright, 2005) (Keijzer
et al., 2014). Effective timing of chrono-therapies aiming to correct SCN driven rhythms requires
accurate assessment of the SCN phase to know when they should be applied to obtain the desired
results.

Because the SCN is not directly accessible in humans, the timing of peripheral markers is used as a
proxy for SCN phase. The choice of which marker(s) to use is influenced by the ease of assessment,
cost, and reliability of the marker itself. While sleep-wake propensity is influenced by the circadian
system, the timing of the sleep-wake cycle is not considered a reliable marker of SCN phase or
period in humans (Czeisler et al., 1999). Even though the circadian (SCN) phase at which sleep occurs
impacts the duration and structure of sleep, studies of shift-workers and jet-lagged travellers as well
as laboratory studies of spontaneous and forced desynchrony have demonstrated that sleep can
occur at many phases and that the relationship between the timing of the sleep-wake cycle and SCN
phase varies across conditions and both between and within individuals (Dijk and Lockley, 2002).

Peripheral rhythms such as the core body temperature, cortisol, and melatonin rhythms have been
used extensively as markers for circadian phase of the SCN. Several lines of evidence support the
validity of these peripheral markers for SCN phase. Neuroanatomical tracer and lesion studies show
that through its projections to the PVN and other hypothalamic nuclei, the SCN drives rhythmicity in
melatonin and cortisol as well as core body temperature (Moore, 2013). Further evidence comes
from the observation that exposure to light induces equivalent phase shifts in these phase markers
(Czeisler et al., 1990), and the sleep-propensity rhythm is closely coupled to the core body
temperature and melatonin rhythms (Dijk et al., 1997). Reductions in amplitude induced by light
pulses or phase shifts of the light-dark and sleep-wake cycle are correlated across melatonin,
cortisol, and core body temperature (Dijk et al., 2012) (Czeisler et al., 1990).

Given the validity of these phase markers, assessing the phase of the SCN relative to clock time or to
the external light-dark cycle, or even relative to another oscillator, would seem to be
For Peer Review

straightforward. However, the key pitfall in using these phase markers remains that their overt rhythmicity is composed of both ‘endogenous’ circadian components and ‘evoked’ components. In humans, the plasma melatonin rhythm serves as an example of this. In constant darkness, a prominent rhythm of melatonin concentration in blood or saliva can be observed, with high values during the biological (subjective) night and low values during the biological (subjective) day. However, light has long been known to suppress melatonin. While initially it was thought that only bright light could do so, it is now recognized that light intensities as low as 6 lux can acutely suppress melatonin by 50% in some participants, thereby ‘masking’ the endogenous SCN phase and amplitude (Phillips et al., 2019) [Zeitzer et al., 2000][Fig 2D]. Therefore, accurate estimation of the phase of the melatonin rhythm requires a time series of blood or saliva samples to be collected in very dim light (Benloucif et al., 2008), a requirement which is not easily met in the real world. The cortisol rhythm, besides being affected by light (Rahman et al., 2019), is also masked by stress and fasting, but just like melatonin, is little affected by sleep (Oster et al., 2017) [Fig 2B].

Whereas in the case of melatonin the primary confounding variable is environmental (i.e. light), for other markers of SCN phase the primary confounding variables are behavioural. Core body temperature (CBT) is lower at night than during the day. The amplitude and phase of the observed CBT rhythm is very much influenced by the timing of rest/sleep. This is demonstrated by data showing that when sleep occurs at night the amplitude of the temperature rhythm is high, whereas when sleep occurs during the day the amplitude of the rhythm is greatly reduced [Fig 2A]. In fact, this masking can be so extreme as to cause an observed temperature nadir to occur during the day, even though the true endogenous nadir (revealed under appropriate conditions) is located at night.

The masking impact of sleep on core temperature is due to a combination of supine posture, inactivity, and sleep itself (Krauchi and Deboer, 2010). The difficulty of assessing circadian phase on the basis of temperature may be further illustrated by contrasting core temperature and skin temperature: whereas sleep lowers core temperature, skin temperature rises when we lie down and fall asleep, and the pattern of temperature varies across distal and proximal parts of the body (Krauchi et al., 2000) (Krauchi and Wirz-Justice, 1994). The conditions in which assessment of SCN phase is of most interest are also conditions in which this assessment is most challenging. In shift-work and jet-lag, sleep will often be displaced from the normal circadian phase and/or normal clock time. Altered phase relationships between sleep and SCN phase also occur in Circadian Rhythm Sleep-Wake Disorders (CRSWD), social jet lag, and neurodegeneration (Rahman et al., 2009).

Furthermore, the sleep-wake cycle is almost always associated with cycles of dark-light and fasting-feeding, and these latter cycles also affect physiological, endocrine, and molecular rhythms. Thus, these peripheral markers (melatonin, cortisol, core temperature) can only be used to assess SCN status when the masking effects of environmental and behavioural cycles are adequately controlled. For the same reasons, adequate environmental and behavioural control should be a prerequisite in the search for any novel methods to assess SCN status.

Univariate markers such as melatonin, cortisol, or core temperature can only be used as markers of the SCN status by collection of a timeseries. The timeseries should be sufficiently long to identify the phase, i.e. minimum, maximum, onset or offset, of the variable under study. If amplitude of the central clock is of interest, the time series must be at least a full circadian cycle (24 hours). In the search for novel methods to assess SCN status, a minimum duration timeseries of at least 24 hours should be used.

In summary, peripheral univariate markers can be used to assess circadian parameters such as phase and amplitude, but require carefully controlled conditions, costly and labour-intensive protocols,
with repeated sampling of blood, urine, or saliva over extended periods of time, and are often burdensome for the research participant.

The dim light melatonin onset (DLMO) (Lewy et al., 1999) assessed in blood or saliva has emerged as the gold standard marker of choice, but urinary 6-sulphatoxy melatonin has also been used successfully (e.g. (Lockley et al., 2015), although the duration of collection is longer and the temporal resolution of the phase assessment may be lower than for the DLMO.

**Dim-light melatonin onset as a biomarker for SCN phase**

The term biomarker is frequently used in medicine and clinical trials and is defined as “a characteristic that is objectively measured and evaluated as an indicator of normal biological processes, pathogenic processes, or pharmacologic responses to a therapeutic intervention” (Menetski et al., 2019). Some of the required or desirable characteristics of a biomarker are that it is present in easily accessible tissues or fluids; that it can be reliably quantified through established essays; that it is sensitive, specific, and valid in a wide variety of situations (robustness) and populations. The dim light melatonin onset (DLMO) is essentially a biomarker for SCN phase, and the DLMO meets many biomarker requirements. Importantly, even though melatonin is masked by light, the masking effects of the sleep-wake cycle itself on melatonin are small. This implies that melatonin is a robust marker for central circadian phase in those conditions in which the phase relationship between sleep and SCN may be changed.

Even though the DLMO is considered a gold standard, assessment of it is not without error. This maybe simple measurement error such as assay errors or errors related to imperfect implementation of the dim light protocol. In addition, it may be that the phase relation between the melatonin rhythm and the relevant SCN rhythm, e.g. the phase response curve to light, or the sleep-propensity rhythm varies between individuals or conditions.

Kronauer and colleagues estimated the error of melatonin based phase assessments of the human circadian pacemaker in a detailed comparison of the variability of phase assessments using melatonin, cortisol and core body temperature (Klerman et al., 2002). It was concluded that melatonin based methods were superior and that the standard deviation for melatonin based methods ranged from 14 to 21 min. This implies that within an individual, phase differences of more than approximately 30 minutes phase can be reliably assessed. For a description of how accuracy and uncertainty of melatonin based phase assessments are affected by sampling frequency, analysis methods, populations studied and thresholds applied we refer to (Klerman et al., 2002) (Danilenko et al., 2014) (Lewy et al., 1999).

In its standard implementation, the main drawbacks of the DLMO are that: a) it requires special environmental conditions (dim light); and 2) it requires a time series of at least several hours of samples. The required sampling frequency and duration of the time series depends on any *a priori* knowledge about the approximate phase and on the required precision of the phase assessment. In many situations it can be assumed that melatonin will rise sometime in the evening hours or early night (a few hours prior to usual bedtime), whereas in other situations no such assumptions can be made (e.g. non-24 hour sleep-wake disorder, shift work, or jet lag), and a time series of 24 hours may be required to capture the DLMO. Whereas the DLMO is most commonly assessed under controlled laboratory conditions, protocols for the use of the DLMO in the home environment have been developed and validated in patient populations (Keijzer et al., 2011) (Pullman et al., 2012) (Burgess et al., 2016).

**Requirements/desirables for novel approaches to assess parameters of the SCN in humans**
The main aim of novel approaches is to overcome the limitations of current approaches. At the same time, these novel approaches should meet the requirements outlined above, such as being robust against masking effects and be able to accurately reflect SCN phase. Thus, novel phase markers should be evaluated against a gold standard phase marker for SCN, e.g. DLMO. The precision of the biomarker should meet the requirements of its application or use-case. Importantly, for many applications/use-cases the biomarker should assess the circadian parameter of interest with sufficient precision to quantify the parameter at the level of the individual, rather than the ‘group’ level.

How to quantify performance of a new marker? Performance of a new marker against a gold standard can be quantified in a variety of ways and different publications use different methods. Some of the metrics used are the error and its standard deviation, the absolute error and its standard deviation, the median error and its range, the fraction of samples with an error less than a particular threshold, the correlation between gold standard and novel marker, regression analysis of gold standard vs. novel method, and there are probably others. All these methods have their advantages and disadvantages and their validity depends on whether underlying assumptions about the distribution of the data, e.g. a normal distribution, are valid. An issue relevant for assessing errors of phase is that the circular nature of the data needs to be taken into account, e.g. a phase at 23:59 and 00:02 h are very close. Another point to keep in mind is that a mean error may simply represent a systematic bias and can be corrected for and an absolute error is only useful if a systematic error (bias) cannot be corrected for.

Novel approaches to assess circadian phase in humans

The novel approaches can be broadly classified into: a) circadian phase assessments based on one or a few samples of high dimensional (i.e. multivariate) material, like the transcriptome or metabolome; b) circadian phase assessments based on long-term passive sampling of behavioural, physiological, and/or environmental variables [Fig 3].

- Omics based assessment of circadian phase of central pacemaker using few samples

High dimensional or multivariate refers to samples that contain a large number of variables or features, the constellation of which varies with circadian phase such that one timepoint suffices to assess any circadian phase. It has long been recognised that a single time point sample of high dimensional data may contain as much information about circadian phase as a timeseries of univariate data. The classical example is Linnaeus’ flower clock. Various species of flowers open at different times of day, and any given time of day is characterised by a constellation of open/closed flowers. It turns out that Linnaeus may never have actually planted a flower clock, and the accuracy of flower clocks that were planted by botanical gardens were affected by weather, latitude, and seasonal changes (https://en.wikipedia.org/wiki/Linnaeus%27s_flower_clock). The transcriptome, proteome, and metabolome are high dimensional data, and a considerable fraction of the thousands of variables in each of these -omics data sets has been shown to be rhythmic in human blood, adipose tissue, skin, and brain (Moller-Levet et al., 2013) (Christou et al., 2019) (Depner et al., 2018) (Ang et al., 2012) (Dallmann et al., 2012) (Wu et al., 2018).

A main problem in the development of biomarkers from high dimensional data is the selection of relevant features. Several approaches exist and may be subdivided in those that use a priori-knowledge, e.g. focus on RNAs from core circadian genes, or unbiased approaches (for a discussion see (Laing et al., 2017) (Laing et al., 2019b). When unbiased approaches are applied to, for example, the transcriptome, we are exposed to the curse of high dimensionality (e.g. ~20,000 transcripts) and
the resultant risk of ‘overfitting’. When there are many more features than data points to be predicted or described it is always possible to find a set of features that will fit or describe a particular data set (like the training data set) but this set of features is much less likely to fit an independent data set (a test set). This problem of overfitting is addressed in machine learning approaches to feature selection and will not be discussed in detail here (Smith et al., 2014).

A first single-time point sampling method for the assessment of circadian phase was developed in mice. This was an approach based on a prior knowledge since it was based on more than 100 time-indicating genes in the liver (Ueda et al., 2004). Subsequently, a single-time point method for assessment of circadian phase in mice was developed based on plasma metabolites (Minami et al., 2009). The strengths of these single-time point methods in animals are that the methods were developed while controlling for effects of feeding, sex, age and the light-dark cycle, as well as that they were validated against gold standard SCN markers such as corticosterone.

In humans, a first blood metabolite-based timetable method was developed based on data from three participants and validated in six participants using plasma cortisol and melatonin as gold standards (Kasukawa et al., 2012). Data were collected under constant routine conditions at the beginning and end of a forced desynchrony protocol, with blood samples drawn every two hours during a 39-hour episode of wakefulness following sleep episodes which were located during the biological night. Using two blood samples taken 12 hours apart, the reported accuracy was approximately three hours. The set of metabolites used for the timetable construction consisted of 58 rhythmic metabolites, a large fraction of which belonged to metabolism pathways of steroid hormones such as cortisol.

A first method to use the human blood transcriptome for the prediction of SCN phase was developed based on 329 mRNA samples from 26 participants to build the model, and a validation set of 349 mRNA samples from 27 participants (Laing et al., 2017). The blood samples were collected during a residential stay in a clinical research centre while participants were scheduled to a normal sleep-wake cycle, a misplaced sleep-wake cycle, or underwent a period of approximately 40 hours of sleep deprivation following a week of sufficient or insufficient sleep (Moller-Levet et al., 2013; Archer et al., 2014). These conditions to some extent mimic conditions like shift-work. Plasma melatonin data were used as a gold standard proxy for SCN phase.

The development of a circadian biomarker based on these data was not based on an initial identification of rhythmic transcripts but used an unbiased ‘machine learning approach’ (partial least squares regression) to build a model. Transcripts were quantified by microarray technology using an Agilent platform. For the development of the model, data from all conditions were used, i.e. not just ‘baseline’ but also conditions in which the circadian system was perturbed. The rationale for using samples from various conditions for model development was that in real-world situations a sample may come from either normal or perturbed conditions and this will often be unknown. That the effects of sleep timing on putative transcriptome based biomarkers can be very substantial has been demonstrated in forced desynchrony experiments [Fig 2B]. A sample collected in a shift work setting may come from a shift worker in whom SCN phase may or may not have adapted to shift work and it will be unknown whether sleep is or is not displaced relative to SCN phase.

This approach identified a set of 100 mRNA abundance features which was able to predict melatonin phase in the validation set with an accuracy (SD or error) of three hours based on one sample, and 1h:40 min based on two samples taken 12 hours apart (82% of samples had an error less than two hours). The latter approach essentially is a within-subject normalisation procedure that improves performance of the method because it removes the large between-subject variation in the blood
transcriptome, i.e. the blood transcriptome is to some extent trait-like. The median error (which means that 50% of the samples have an error less than this number) was less than one hour for the differential model. Importantly, even though the model was developed from samples across a variety of sleep-wake conditions, the accuracy was still less when sleep occurred out of phase (See Fig 4).

The unbiased approach used in Laing et al. (Laing et al., 2017) outperformed timetable methods and ZeitZeiger when applied to the same data set.

Comparing the features identified by each of these methods revealed that many of the features are not directly related to clock genes but many are related to glucocorticoid signalling pathways.

Hughey and colleagues applied an algorithm (ZeitZeiger), in which a rhythmic spline is fitted to the data (Hughey, 2017) to publicly available data sets collected at the University of Surrey and the University of Pennsylvania (Moller-Levet et al., 2013) (Archer et al., 2014) (Arnardottir et al., 2014). The algorithm builds a predictor for what is called ‘circadian time’. Importantly, in this analysis ‘circadian time’ does not refer to the circadian phase of a sample based on melatonin phase in that individual, but to the time of the sample relative to sunrise or the melatonin phase averaged across individuals. The model was developed on data collected under baseline sleep-wake conditions and when applied to baseline data the absolute mean error was 2.1 hours. When the model was tested on conditions during which the sleep-wake cycle was displaced, the performance was worse, such that the variability in prediction error increased by 42% with a median absolute error greater than three hours. Hughey and colleagues then developed strategies to improve prediction at the individual level by using more than one sample per participant, such that using two samples taken 8-9 hours apart resulted in an average improvement in prediction of 0.43 hours to approximately 1.67 hours. A strength of the ‘ZeitZeiger predictor’ is that it is based on a small set of 15 genes. In accordance with the results from Laing et al. (Laing et al., 2017), only a very few of these 15 genes were core clock genes.

Braun et al. (Braun et al., 2018) developed a new algorithm using the University of Surrey and the University of Pennsylvania data sets, as well as a new data set consisting of RNAseq data obtained from healthy participants collected during unperturbed baseline conditions. The authors used a 2-sample approach, and their method requires at least two samples which for optimal performance should be drawn 10-12 hours apart. Unfortunately, the authors did not use a circadian phase marker to validate their method but instead used external clock time (this limitation has been discussed previously (Laing et al., 2019a) (Braun et al., 2019)). They identified a set consisting of approximately 40 genes, few of which were core clock genes. The reported median error based on two samples was two hours.

Whereas the Laing et al., Hughey et al., and Braun [CHANGE-18] et al. approaches were based on whole blood, Wittenbrink and colleagues (Wittenbrink et al., 2018) used monocytes as the source of their transcripts, assuming that rhythmicity would be more robust in this cell type. In addition, the authors argued that because microarray and RNAseq platforms are relatively inaccurate in detecting transcript levels, it may be beneficial to use other platforms for the implementation of circadian phase biomarker sets. Wittenbrink and colleagues took this into consideration when they developed a biomarker for circadian phase using a two-step process. In the first step, they used ZeitZeiger to identify a set of transcripts quantified by RNAseq. This predictor set was then implemented on a NanoString platform. Their initial search used samples collected in a group of healthy young men in a constant routine protocol, and their validation dataset was from a group of individuals with extreme chronotypes living in real-world conditions from whom samples were collected in the morning and
afternoon. In both cases, the gold standard proxy for circadian phase was salivary DLMO, which in the validation set varied between approximately 17:00 and 01:00 hours. The authors created 4 predictors: 1-sample-12 genes; 1-sample-2 genes; 2-sample, 13-genes and 2-sample,2-genes. In the validation set the median absolute error using the Nanostring platform was less than an hour across all predictors and timing of validation samples (morning, afternoon). The 1-sample 12-genes predictor had an absolute median error of 0.7 hours for a morning sample and 0.8 hours for afternoon sample. The performance of the two-sample predictors was similar to the one sample predictors (See Fig 4b). A weakness of this otherwise elegant study is that performance was not assessed under conditions in which sleep was displaced, such as in forced desynchrony, jet-lag, or shift work. Thus, we do not yet know to what extent the performance of this method will be affected by the masking effects of sleep, activity, posture, or feeding-fasting, which have been shown to be considerable.

**Long term passive sampling-based assessment of circadian phase using activity, skin temperature, heart rate, light exposure, and other variables.**

The desire to assess SCN phase based on passive monitoring of physiological and behavioural variables has a long history. Traditionally, these approaches used linear methods in which estimated masking effects were added or subtracted from the observed variables. However, the effects of masking differ across individuals and interact with circadian phase (that is, the masking effects are larger at some circadian phases than others), rendering simple addition/subtraction methods inadequate (Klerman et al., 1999). Recently, more sophisticated approaches in which multiple variables are recorded simultaneously and algorithms are used to predict melatonin phase have been developed using multiple regression or artificial neural network approaches. The various approaches differ primarily with respect to the included variables and required duration of data collection.

**Skin temperature, light, and activity**

Whereas core body temperature has long been considered a valuable marker of circadian phase it is cumbersome to measure by either rectal sensors or thermistor pills that are swallowed. Skin temperature can be measured in a less intrusive manner. The circadian and sleep-wake and activity dependent regulation of skin temperature has been investigated extensively (Krauchi and Wirz-Justice, 1994). Based on these findings skin temperature has been evaluated as a source of information about central circadian phase. Kolodyazhniy and colleagues published two approaches based on ambulatory skin temperature recordings (from six locations), (blue) light recordings from a sensor mounted on glasses, and motion (Kolodyazhniy et al., 2011; Kolodyazhniy et al., 2012). Models were constructed using either multiple regression or artificial neural networks, and gold standard estimation of SCN phase was based on salivary melatonin collected under constant routine conditions. Participants were healthy but of various chronotypes, with a range of melatonin phases of slightly more than five hours. Performance of the algorithms was derived from leave-one-participant-out cross-validation. The best model correlated well with melatonin phase (r=0.97) with a SD of the error of only 23 minutes. In one of the very few instances in which a model developed by one group was tested by another group, Stone et al. put this model to the test (Stone et al., 2019b). Importantly, in this study the model was tested not only in participants on a normal sleep-wake schedule but also in shift workers, and the gold standard proxy for SCN phase was either DLMO or 6-sulphatoxy melatonin. Whereas in this independent validation the model performed well under baseline conditions, the performance deteriorated dramatically for assessments in night shift workers such that the error was more than two hours in approximately half of the assessments (See Fig 4c). This poor performance persisted even when the model was trained on night shift data.
Performance was even worse when the model was trained on non-shift-work data. The findings of this comprehensive study underscore the necessity to validate any method for circadian phase assessment in situations in which sleep and other masking effects are displaced, not only because these are more challenging conditions for most methods but because they represent situations in which circadian biomarkers will be used.

**Combining a mathematical model for light with light exposure and activity data**

Although in the studies of Kolodyazhniy et al. and Stone et al. light information was used to build a predictor algorithm, no specific model for how light affects the human circadian pacemaker was employed. Kronauer’s mathematical model for the effects of light (and activity) on the human circadian pacemaker (Jewett et al., 1999; St Hilaire et al., 2007) (Kronauer et al., 1982), which is based on the extensive laboratory studies by Czeisler and colleagues (Duffy et al., 1996) (Boivin et al., 1996) (Czeisler et al., 1989) (Jewett et al., 1991) (Gronfier et al., 2004) (Chang et al., 2012), remains the only quantitative model to date (Duffy and Wright, 2005). It uses light, quantified as lux, as input and no provision for the spectral composition of light is available. The primary assumption underlying the use of Kronauer’s model to predict circadian phase is that light is the most powerful zeitgeber for the human SCN and that variations in timing and intensity of light exposure, which are in part driven by the timing of sleep and social constraints (Skeldon et al., 2017), are the main determinants of variations in circadian phase. Determinants of phase of entrainment are well understood at a theoretical level (Granada et al., 2013). Phase of entrainment is determined by individual differences in intrinsic period [Wright et al., 2005]] and theoretically also by individual differences in light sensitivity. Woelders and colleagues used Kronauer’s model and reported that DLMO was associated with individual differences in light exposure, such that Light+Model explained 52% of the variance (Woelders et al., 2017). When light data and Kronauer’s model were supplemented with activity data, DLMO could be predicted with a SD or error of 1.1 hours, i.e. 95% of the predictions had an error of 2.2 hours or less. It is noted that although the participants varied with respect to chronotype with a considerable range of measured DLMOs of 9.3 hours, they were all sleeping at their habitual bedtimes for the duration of the study. In a study of shift workers on either a diurnal or night schedule, Stone and colleagues used a similar approach, i.e. they either used a photic-only model or a combination of photic input to Kronauer’s model and activity (Stone et al., 2019a). The proxy for SCN phase was 6-sulphatoxy melatonin. Performance on the diurnal schedule was comparable and even slightly better than in the Woelders et al. study. Although performance of the predictors deteriorated somewhat on the night schedule, performance was still rather good with a SD of 1.39 hours and 80% of predictions within two hours of the observed values (See Fig 4C). St. Hilaire and colleagues have also used individual light exposure data input to the Kronauer model to predict phase shifts in a simulated shift work study where salivary DLMO was assessed before and after a series of night shifts. They found that 85% of the model predictions were within 2 hours of the observed DLMO shifts (St. Hilaire & Duffy, personal communication).

**Heart rate**

Heart rate (HR) and heart rate variability (HRV) are under control of the autonomic nervous system, which in turn is under SCN control. The masking effect of activity and sleep on heart rate is considerable, but constant routine protocols have demonstrated an endogenous circadian component to heart rate (Viola et al., 2002) (Vandewalle et al., 2007). In principle, this implies that ambulatory monitoring of heart rate may provide information on SCN phase provided that masking can be accounted for. Gil and colleagues investigated the potential of HR measures for circadian phase prediction using only 24 hours of data (Gil et al., 2013) (Gil et al., 2014). The proxy for SCN phase was DLMO measured in 11 participants for model development and 19 participants for
performance testing. Participants slept at their habitual times and the range of chronotypes and
DLMOs was relatively small, i.e. approximately three hours. The model was based on an
autoregressive moving average with a linear combination of external inputs, e.g. heart rate or light.
Using only heart rate as a predictor resulted in a SD of 56 minutes, similar to performance of a light
input only model (with light transformed with a power function, as in Kronauer’s model). Combining
heart rate and light resulted in performance with a SD of 39 minutes and adding activity did not
improve performance significantly. Although performance of these models appears impressive, it
should be noted that the range of DLMO was limited and this model has not been tested under
conditions of displaced sleep.

New methods to assess intrinsic circadian period

Gold standard assessment of the intrinsic period of the SCN in sighted people is achieved through
forced desynchrony of the sleep-wake cycle from the endogenous circadian rhythms by scheduling
sleep-wake and the associated dim light-dark cycle to a non-circadian cycle length that is outside the
range of entrainment (Klerman et al., 1996) (Czeisler et al., 1999; Duffy et al., 2011) (Lazar et al.,
2013). In totally blind individuals, intrinsic period can be assessed by repeated phase assessments of
48-hour urinary 6-sulphatoxy melatonin while the participants are living in their habitual
environment (Hack et al., 2003; Lockley et al., 2015). These experiments have provided important
insights into the role of interindividual variation in intrinsic period in entrainment. Several variants of
the forced desynchrony protocol have been developed, and a consistent finding across studies is
that that in participants without circadian rhythm sleep-wake disorders, the average period is about
24.2 hours with a small standard deviation (Czeisler et al., 1999; Duffy et al., 2011) (Micic et al.,
2016) (Eastman et al., 2017) (Hasan et al., 2012). Two alternative period assessment methods have
emerged: a) in vitro recording of circadian rhythms in human cell cultures, in which cells are
modified to express a luciferase gene under the control of the promoter of a ‘clock’ gene; and b) reducing
the residual variance of light exposure and mathematical model-derived prediction of
circadian phase (DLMO) by optimising the period parameter of Kronauer’s model at the level of the
individual. The latter method has yielded a realistic population average and SD for circadian period,
but individual estimates have not been validated against gold standard assessments. Furthermore,
the period estimates were based on only one circadian phase assessment per participant, and the
optimisation procedure may therefore not reflect intrinsic period but rather the period which best
predicts this single circadian phase (Woelders et al., 2017).

For the in vitro circadian period assessment methods, a number of validated studies are available in
both blind and sighted individuals. However, in these studies the average in vitro period is longer
than that derived from forced desynchrony studies, and systematically longer than that derived in
the same individuals by melatonin (Pagani et al., 2010; Hasan et al., 2012). In fact, the correlation
between the in vitro periods and the periods as assessed by urinary 6-sulphatoxy melatonin (in blind
individuals) or by plasma melatonin in forced desynchrony studies in sighted people are weak and
sometimes not even significant. It thus appears that the current in vitro assessments will not be able
to accurately assess intrinsic period at the level of the individual.

New methods to assess intrinsic circadian amplitude/circadian disruption

Changes in the amplitude of overt rhythms such as activity or body temperature are often observed,
in for example ageing, and it is also often tacitly assumed that these changes reflect changes in the
amplitude of the endogenous circadian components of these variables. As we have previously
argued, protocols such as the constant routine are needed to ascertain this (Duffy and Dijk, 2002)
[Duffy and Dijk]. However, in general, the concept of circadian amplitude is not well-defined and a
simple gold standard measure for circadian amplitude has not been agreed upon. For example, there is substantial variation in melatonin amplitude between individuals, and little evidence that those individual differences reflect differences in SCN amplitude. Studies in which a constant routine has been carried out before and after the circadian system has been perturbed have demonstrated that changes in the amplitudes of cortisol, melatonin, and core body temperature in response to an intervention are generally correlated [Jewett et al. 1991 [(Dijk et al., 2012). This suggests that the amplitudes of these measures may reflect the amplitude of oscillations within the SCN, and that it may be possible to develop a metric for circadian amplitude.

New approaches to assess endogenous circadian amplitude are based on multivariate approaches. These methods may not necessarily aim to assess the amplitude of the SCN but may be targeted at assessment of peripheral oscillators. In fact, these methods may not aim to assess the simple construct of amplitude but instead metrics that reflect the robustness or ‘normality’ of circadian processes. The assumption is that under normal conditions diurnal oscillations in a particular organ or tissue are characterised by a typical progression of expression levels of genes (a tissue specific temporal program), and in particular genes that are at the core of the generation of circadian rhythmicity. Shilts and colleagues designed a method to quantify ‘circadian disruption’ by computing a metric called ‘clock correlation distance’ (CCD) which is based on the co-expression patterns of 12 ‘clock genes’ (Shilts et al., 2018). A larger CCD indicates less normal, or less robust rhythmicity. The method was developed on samples from various mouse tissues but then applied to samples from human blood, skin, brain and \textit{in vitro} cell cultures. The robustness of circadian rhythmicity as detected by this method varied between blood, skin, brain and \textit{in vitro} cell cultures. The method does not require samples to be labelled with time of day, works best if the entire circadian cycle is covered but can perform reasonably well if only part (e.g. one third) of the circadian cycle is covered. For the method to produce reliable results in humans it requires approximately 30 samples, and the method cannot be applied to a single sample. One application for this method is in cancer research and it was indeed shown that this method detects disruption of circadian rhythmicity in tumours. Vlachou and co-workers also aimed to develop a method to quantify circadian disruption as well as circadian time with intended application in oncology [BioRxiv doi: http://dx.doi.org/10.1101/622050]. Their method, called ‘time teller’ is based on 10-15 genes, requires only one sample, and is designed to be used to both assess circadian phase as well as circadian ‘dysfunction’. As mentioned, these methods to assess circadian disruption are built on the notion that during a normal circadian cycle, at any point in the cycle there is a specific constellation of rhythmic genes being expressed. This assumption can be used to tell time from a single sample but also can be used to detect disruption of circadian organisation. Obviously, these methods are very much focussed on the local circadian organisation of the tissue from which the sample was obtained. Circadian disruption in the periphery does not necessarily imply circadian disruption in the central circadian clock. For example, severe disruption of circadian organisation of the transcriptome, including expression of core clock genes, has been observed in the whole blood transcriptome when sleep was displaced to the daytime while at the same time the phase and amplitude of the plasma melatonin rhythm were similar to when sleep occurred at night (Archer et al., 2014).

These observations highlight both the potential of using multivariate data in the periphery to detect circadian disruption or quantify ‘amplitude’ while at the same time emphasizing the need for careful validation and interpretation of these data. For the assessment of the physiological patency of a local tissue, the cause of the disruption of temporal programs may not be that important. However, in any multi-oscillator system, local rhythmicity and its disruption may sometimes reflect disruption
of local clocks, disruption of circadian rhythmicity downstream from those local clocks, or disruption of central clocks imposing rhythmicity on those local tissues.

**Concluding remarks**

Wearables and -omics data combined with machine learning and mathematical modelling hold great promise for the development of novel methods to quantify circadian processes in humans. Optimal choice of variables to be collected by wearables, the -omics to be used in ‘one sample’ methods, or the source of samples, i.e. blood, skin, saliva, remain to be established. This choice of variable(s) will be influenced by the purpose of the biomarker (See Fig 1,2). Until now, much emphasis has been on the development of ‘sophisticated’ algorithms without clearly stating the purpose of the biomarker. Little effort has been devoted to comparison of these biomarkers to gold standards, validating them in realistic protocols, or defining the required accuracy across use cases. A variety of performance measures have been used in different studies, and reaching a consensus on the performance metrics of biomarkers will facilitate comparison across methods.

It will be useful to evaluate these new methods within a framework that is based on concepts developed in circadian rhythm research together with concepts from the field of biomarker development. Organisations such as the FDA and NIH together with Industry partners have provided guidelines for the evidence needed for biomarker qualification and a description of the workflow and decision processes in biomarker development (Leptak et al., 2017). Some of the key concepts that can be applied to circadian biomarkers are ‘context of use’ which relates to ‘what question does the biomarker address’, the ‘biological rationale for use of biomarker’, ‘independent data sets for qualification’ and ‘comparison to current standard’.

Circadian concepts such as diurnal vs. circadian rhythmicity, masked vs. endogenous circadian rhythms will remain useful when evaluating new markers for circadian processes. Likewise, biomarker concepts such as robustness, reliability, sensitivity, and specificity should be formalized for our field and applied to any novel method designed for quantifying circadian rhythms. It may be unrealistic to expect that a particular biomarker for, for example circadian phase, is universally robust, i.e. can be applied in a wide range of situation and populations and it may be that ‘situation’ ‘population’ or ‘diagnosis’ specific approaches can be developed. However, in practise the precise situation, population or diagnosis will be often unknown. Therefore, for these novel, be it universal or specific, biomarkers to be useful they should be validated in controlled laboratory settings where the rest-activity schedules of participants are manipulated, tested in men and women of all ages, tested in large groups of normal individuals in “real life” situations, and tested in patient populations, particularly those patient populations in which these methods will be applied.

**Acknowledgements**

The authors’ research was supported by grants from the BBSRC(BB/F022883), AFOSR (FA9550-08-1-0080), the UK Dementia Research Institute, and the NIH (R01 AG044416, P01 AG09975).

The authors do not have patents related to biomarkers and declare no competing interests.

The authors thank Drs. Simon N. Archer and Anne C. Skeldon for comments on the manuscript and Dr. Carla S. Möller-Levet for additional data analyses.

**References**
Ang JE, Revell V, Mann A, Mantele S, Otway DT, Johnston JD, Thumser AE, Skene DJ, and Raynaud F (2012) Identification of human plasma metabolites exhibiting time-of-day variation using an untargeted liquid chromatography-mass spectrometry metabolomic approach. Chronobiol Int 29:868-881.

Archer SN, Laing EE, Moller-Levet CS, van der Veen DR, Bucca G, Lazar AS, Santhi N, Slak A, Kabiljo R, von Schantz M, Smith CP, and Dijk DJ (2014) Mistimed sleep disrupts circadian regulation of the human transcriptome. Proc Natl Acad Sci U S A 111:E682-691.

Archer SN, and Oster H (2015) How sleep and wakefulness influence circadian rhythmicity: effects of insufficient and mistimed sleep on the animal and human transcriptome. J Sleep Res 24:476-493.

Arnardottir ES, Nikonova EV, Shockley KR, Podtelezhnikov AA, Anafi RC, Tanis KQ, Maislin G, Stone DJ, Renger JJ, Winrow CJ, and Pack AI (2014) Blood-gene expression reveals reduced circadian rhythmicity in individuals resistant to sleep deprivation. Sleep 37:1589-1600.

Atger F, Gobet C, Marquis J, Martin E, Wang J, Weger B, Lefebvre G, Descombes P, Naef F, and Gachon F (2015) Circadian and feeding rhythms differentially affect rhythmic mRNA transcription and translation in mouse liver. Proc Natl Acad Sci U S A 112:E6579-6588.

Baxter M, and Ray DW (2019) Circadian rhythms in innate immunity and stress responses.

Benloucif S, Burgess HJ, Klerman EB, Lewy AJ, Middleton B, Murphy PJ, Parry BL, and Revell VL (2008) Measuring melatonin in humans. J Clin Sleep Med 4:66-69.

Boivin DB, Duffy JF, Kronauer RE, and Czeisler CA (1996) Dose-response relationships for resetting of human circadian clock by light. Nature 379:540-542.

Bollinger T, and Schibler U (2014) Circadian rhythms - from genes to physiology and disease. Swiss Med Wkly 144:w13984.

Braun R, Kath WL, Iwanaszko M, Kula-Eversole E, Abbott SM, Reid KJ, Zee PC, and Allada R (2018) Universal method for robust detection of circadian state from gene expression. Proc Natl Acad Sci U S A 115:E9247-E9256.

Christou S, Wehrens SMT, Isherwood C, Moller-Levet CS, Wu H, Revell VL, Bucca G, Skene DJ, Laing EE, Archer SN, and Johnston JD (2019) Circadian regulation in human white adipose tissue revealed by transcriptome and metabolic network analysis. Sci Rep 9:2641.

Czeisler CA, Duffy JF, Shanahan TL, Brown EN, Mitchell JF, Rimmer DW, Ronda JM, Silva EJ, Allan JS, Emens JS, Dijk DJ, and Kronauer RE (1999) Stability, precision, and near-24-hour period of the human circadian pacemaker. Science 284:2177-2181.

Czeisler CA, Johnson MP, Duffy JF, Brown EN, Ronda JM, and Kronauer RE (1990) Exposure to bright light and darkness to treat physiologic maladaptation to night work. N Engl J Med 322:1253-1259.

https://mc.manuscriptcentral.com/jbrhythms
Czeisler CA, and Klerman EB (1999) Circadian and sleep-dependent regulation of hormone release in humans. Recent Prog Horm Res 54:97-130; discussion 130-132.

Czeisler CA, Kronauer RE, Allan JS, Duffy JF, Jewett ME, Brown EN, and Ronda JM (1989) Bright light induction of strong (type 0) resetting of the human circadian pacemaker. Science 244:1328-1333.

Dallmann R, Viola AU, Tarokh L, Cajochen C, and Brown SA (2012) The human circadian metabolome. Proc Natl Acad Sci U S A 109:2625-2629.

Danilenko KV, Verevkin EG, Antyufeev VS, Wirz-Justice A, and Cajochen C (2014) The hockey-stick method to estimate evening dim light melatonin onset (DLMO) in humans. Chronobiol Int 31:349-355.

Depner CM, Melanson EL, McHill AW, and Wright KP, Jr. (2018) Mistimed food intake and sleep alters 24-hour time-of-day patterns of the human plasma proteome. Proc Natl Acad Sci U S A 115:E5390-E5399.

Dijk DJ, and Czeisler CA (1995) Contribution of the circadian pacemaker and the sleep homeostat to sleep propensity, sleep structure, electroencephalographic slow waves, and sleep spindle activity in humans. J Neurosci 15:3526-3538.

Dijk DJ, Duffy JF, Silva EJ, Shanahan TL, Boivin DB, and Czeisler CA (2012) Amplitude reduction and phase shifts of melatonin, cortisol and other circadian rhythms after a gradual advance of sleep and light exposure in humans. PLoS One 7:e30037.

Dijk DJ, and Lockley SW (2002) Integration of human sleep-wake regulation and circadian rhythmicity. J Appl Physiol (1985) 92:852-862.

Dijk DJ, Shanahan TL, Duffy JF, Ronda JM, and Czeisler CA (1997) Variation of electroencephalographic activity during non-rapid eye movement and rapid eye movement sleep with phase of circadian melatonin rhythm in humans. J Physiol 505 (Pt 3):851-858.

Dowton P, Early JO, and Gibbs JE (2019) Circadian rhythms in adaptive immunity. Immunology.

Duffy JF, Cain SW, Chang AM, Phillips AJ, Munch MY, Gronfier C, Wyatt JK, Dijk DJ, Wright KP, Jr., and Czeisler CA (2011) Sex difference in the near-24-hour intrinsic period of the human circadian timing system. Proc Natl Acad Sci U S A 108 Suppl 3:15602-15608.

Duffy JF, and Dijk DJ (2002) Getting through to circadian oscillators: why use constant routines? J Biol Rhythms 17:4-13.

Duffy JF, Dijk DJ, Hall EF, and Czeisler CA (1999) Relationship of endogenous circadian melatonin and temperature rhythms to self-reported preference for morning or evening activity in young and older people. J Investig Med 47:141-150.

Duffy JF, Kronauer RE, and Czeisler CA (1996) Phase-shifting human circadian rhythms: influence of sleep timing, social contact and light exposure. J Physiol 495 (Pt 1):289-297.

Duffy JF, and Wright KP, Jr. (2005) Entrainment of the human circadian system by light. J Biol Rhythms 20:326-338.

Eastman CI, Tomaka VA, and Crowley SJ (2017) Sex and ancestry determine the free-running circadian period. J Sleep Res 26:547-550.

Gil EA, Aubert XL, and Beersma DG (2014) Ambulatory estimation of human circadian phase using models of varying complexity based on non-invasive signal modalities. Conf Proc IEEE Eng Med Biol Soc 2014:2278-2281.

Gil EA, Aubert XL, Most El, and Beersma DG (2013) Human circadian phase estimation from signals collected in ambulatory conditions using an autoregressive model. J Biol Rhythms 28:152-163.

Granada AE, Bordyugov G, Kramer A, and Herzel H (2013) Human chronotypes from a theoretical perspective. PLoS One 8:e59464.

Greenwell BJ, Trott AJ, Beytebiere JR, Pao S, Bosley A, Beach E, Finegan P, Hernandez C, and Menet JS (2019) Rhythmic Food Intake Drives Rhythmic Gene Expression More Potently than the Hepatic Circadian Clock in Mice. Cell Rep 27:649-657 e645.
Gronfier C, Wright KP, Jr., Kronauer RE, Jewett ME, and Czeisler CA (2004) Efficacy of a single sequence of intermittent bright light pulses for delaying circadian phase in humans. Am J Physiol Endocrinol Metab 287:E174-181.

Hack LM, Lockley SW, Arendt J, and Skene DJ (2003) The effects of low-dose 0.5-mg melatonin on the free-running circadian rhythms of blind subjects. J Biol Rhythms 18:420-429.

Hasan S, Santhi N, Lazar AS, Slak A, Lo J, von Schantz M, Archer SN, Johnston JD, and Dijk DJ (2012) Assessment of circadian rhythms in humans: comparison of real-time fibroblast reporter imaging with plasma melatonin. FASEB J 26:2414-2423.

Honma S (2018) The mammalian circadian system: a hierarchical multi-oscillator structure for generating circadian rhythm. J Physiol Sci 68:207-219.

Hor CN, Yeung J, Jan M, Emmenegger Y, Hubbard J, Xenarios I, Naef F, and Franken P (2019) Sleep-wake-driven and circadian contributions to daily rhythms in gene expression and chromatin accessibility in the murine cortex. Proc Natl Acad Sci U S A 116:25773-25783.

Hughey JJ (2017) Machine learning identifies a compact gene set for monitoring the circadian clock in human blood. Genome Med 9:19.

Jewett ME, Forger DB, and Kronauer RE (1999) Revised limit cycle oscillator model of human circadian pacemaker. J Biol Rhythms 14:493-499.

Jewett ME, Kronauer RE, and Czeisler CA (1991) Light-induced suppression of endogenous circadian amplitude in humans. Nature 350:59-62.

Kasukawa T, Sugimoto M, Hida A, Minami Y, Mori M, Honma S, Honma K, Mishima K, Soga T, and Ueda HR (2012) Human blood metabolite timetable indicates internal body time. Proc Natl Acad Sci U S A 109:15036-15041.

Keijzer H, Smits MG, Duffy JF, and Curfs LM (2014) Why the dim light melatonin onset (DLO) should be measured before treatment of patients with circadian rhythm sleep disorders. Sleep Med Rev 18:333-339.

Keijzer H, Smits MG, Peeters T, Looman CW, Endenburg SC, and Gunnewiek JM (2011) Evaluation of salivary melatonin measurements for Dim Light Melatonin Onset calculations in patients with possible sleep-wake rhythm disorders. Clin Chim Acta 412:1616-1620.

Khan S, Nobili L, Khatami R, Loddenkemper T, Cajochen C, Dijk DJ, and Eriksson SH (2018) Circadian rhythm and epilepsy. Lancet Neurol 17:1098-1108.

Klerman EB, Dijk DJ, Kronauer RE, and Czeisler CA (1996) Simulations of light effects on the human circadian pacemaker: implications for assessment of intrinsic period. Am J Physiol 270:R271-282.

Klerman EB, Gershengorn HB, Duffy JF, and Kronauer RE (2002) Comparisons of the variability of three markers of the human circadian pacemaker. J Biol Rhythms 17:181-193.

Klerman EB, Lee Y, Czeisler CA, and Kronauer RE (1999) Linear demasking techniques are unreliable for estimating the circadian phase of ambulatory temperature data. J Biol Rhythms 14:260-274.

Kolodyazhniy V, Spati J, Frey S, Gotz T, Wirz-Justice A, Krauchi K, Cajochen C, and Wilhelm FH (2011) Estimation of human circadian phase via a multi-channel ambulatory monitoring system and a multiple regression model. J Biol Rhythms 26:55-67.

Kolodyazhniy V, Spati J, Frey S, Gotz T, Wirz-Justice A, Krauchi K, Cajochen C, and Wilhelm FH (2012) An improved method for estimating human circadian phase derived from multichannel ambulatory monitoring and artificial neural networks. Chronobiol Int 29:1078-1097.

Koronowski KB, Kinouchi K, Welz PS, Smith JG, Zinna VM, Shi J, Samad M, Chen S, Magnan CN, Kinchen JM, Li W, Baldi P, Benitah SA, and Sassone-Corsi P (2019) Defining the Independence of the Liver Circadian Clock. Cell 177:1448-1462 e1414.

Krauchi K, Cajochen C, Werth E, and Wirz-Justice A (2000) Functional link between distal vasodilation and sleep-onset latency? Am J Physiol Regul Integr Comp Physiol 278:R741-748.

Krauchi K, and Deboer T (2010) The interrelationship between sleep regulation and thermoregulation. Front Biosci (Landmark Ed) 15:604-625.

https://mc.manuscriptcentral.com/jbrhythms
Krauchi K, and Wirz-Justice A (1994) Circadian rhythm of heat production, heart rate, and skin and core temperature under unmasking conditions in men. Am J Physiol 267:R819-829.

Kronauer RE, Czeisler CA, Pilato SF, Moore-Ede MC, and Weitzman ED (1982) Mathematical model of the human circadian system with two interacting oscillators. Am J Physiol 242:R3-17.

Kuhlman SJ, Craig LM, and Duffy JF (2018) Introduction to Chronobiology. Cold Spring Harb Perspect Biol 10.

Laing EE, Moller-Levet CS, Archer SN, and Dijk DJ (2019a) Universal and robust assessment of circadian time? Proc Natl Acad Sci U S A 116:5205.

Laing EE, Moller-Levet CS, Dijk DJ, and Archer SN (2019b) Identifying and validating blood mRNA biomarkers for acute and chronic insufficient sleep in humans: a machine learning approach. Sleep 42.

Laing EE, Moller-Levet CS, Poh N, Santhi N, Archer SN, and Dijk DJ (2017) Blood transcriptome based biomarkers for human circadian phase. Elife 6.

Lazar AS, Santhi N, Hasan S, Lo JC, Johnston JD, Von Schantz M, Archer SN, and Dijk DJ (2013) Circadian period and the timing of melatonin onset in men and women: predictors of sleep during the weekend and in the laboratory. J Sleep Res 22:155-159.

Leng Y, Musiek ES, Hu K, Cappuccio FP, and Yaffe K (2019) Association between circadian rhythms and neurodegenerative diseases. Lancet Neurol 18:307-318.

Leptak C, Menetski JP, Wagner JA, Aubrecht J, Brady L, Brumfield M, Chin WW, Hoffmann S, Kelloff G, Lavezzari G, Ranganathan R, Sauer JM, Sistare FD, Zabka T, and Wholley D (2017) What evidence do we need for biomarker qualification? Sci Transl Med 9.

Lewy AJ, Cutler NL, and Sack RL (1999) The endogenous melatonin profile as a marker for circadian phase position. J Biol Rhythms 14:227-236.

Lockley SW, Dressman MA, Licamele L, Xiao C, Fisher DM, Flynn-Evans EE, Hull JT, Torres R, Lavedan C, and Polymeropoulos MH (2015) Tasimelteon for non-24-hour sleep-wake disorder in totally blind people (SET and RESET): two multicentre, randomised, double-masked, placebo-controlled phase 3 trials. Lancet 386:1754-1764.

Logan RW, and McClung CA (2019) Rhythms of life: circadian disruption and brain disorders across the lifespan. Nat Rev Neurosci 20:49-65.

Lucey BP, Fagan AM, Holtzman DM, Morris JC, and Bateman RJ (2017) Diurnal oscillation of CSF Abeta and other AD biomarkers. Mol Neurodegener 12:36.

Lyall LM, Wyse CA, Graham N, Ferguson A, Lyall DM, Cullen B, Celis Morales CA, Biello SM, Mackay D, Ward J, Strawbridge RJ, Gill JMR, Bailey MES, Pell JP, and Smith DJ (2018) Association of disrupted circadian rhythmicity with mood disorders, subjective wellbeing, and cognitive function: a cross-sectional study of 91 105 participants from the UK Biobank. Lancet Psychiatry 5:507-514.

Maret S, Dorsaz S, Gurcel L, Pradervand S, Petit B, Pfister C, Hagenbuche O, O'Hara BF, Franken P, and Tafti M (2007) Homer1a is a core brain molecular correlate of sleep loss. Proc Natl Acad Sci U S A 104:20090-20095.

Menetski JP, Hoffmann SC, Cush SS, Kamphaus TN, Austin CP, Herrling PL, and Wagner JA (2019) The Foundation for the National Institutes of Health Biomarkers Consortium: Past Accomplishments and New Strategic Direction. Clin Pharmacol Ther 105:829-843.

Micic G, Lovato N, Gradisar M, Burgess HJ, Ferguson SA, and Lack L (2016) Circadian Melatonin and Temperature Taus in Delayed Sleep-wake Phase Disorder and Non-24-hour Sleep-wake Rhythm Disorder Patients: An Ultradian Constant Routine Study. J Biol Rhythms 31:387-405.

Minami Y, Kasukawa T, Kazaku Y, Iigo M, Sugimoto M, Ikeda S, Yasui A, van der Horst GT, Soga T, and Ueda HR (2009) Measurement of internal body time by blood metabolomics. Proc Natl Acad Sci U S A 106:9890-9895.

Mohawk JA, Green CB, and Takahashi JS (2012) Central and peripheral circadian clocks in mammals. Annu Rev Neurosci 35:445-462.
Moller-Levet CS, Archer SN, Bucca G, Laing EE, Slak A, Kabiljo R, Lo JC, Santhi N, von Schantz M, Smith CP, and Dijk DJ (2013) Effects of insufficient sleep on circadian rhythmicity and expression amplitude of the human blood transcriptome. Proc Natl Acad Sci U S A 110:E1132-1141.

Moore RY (2013) The suprachiasmatic nucleus and the circadian timing system. Prog Mol Biol Transl Sci 119:1-28.

Morris CJ, Aeschbach D, and Scheer FA (2012) Circadian system, sleep and endocrinology. Mol Cell Endocrinol 349:91-104.

Mullington JM, Abbott SM, Carroll JE, Davis CJ, Dijk DJ, Dinges DF, Gehrmann PR, Ginsburg GS, Gozal D, Haack M, Lim DC, Macrea M, Pack AI, Plante DT, Teske JA, and Zee PC (2016) Developing Biomarker Arrays Predicting Sleep and Circadian-Coupled Risks to Health. Sleep 39:727-736.

Munch M, and Kramer A (2019) Timing matters: New tools for personalized chronomedicine and circadian health. Acta Physiol (Oxf) 227:e13300.

Mure LS, Le HD, Benegiamo G, Chang MW, Rios L, Jillani N, Ngotho M, Kariuki T, Dkhissi-Benyahya O, Cooper HM, and Panda S (2018) Diurnal transcriptome atlas of a primate across major neural and peripheral tissues. Science 359.

Oster H, Challet E, Ott V, Arvat E, de Kloet ER, Dijk DJ, Lightman S, Vgontzas A, and Van Cauter E (2017) The Functional and Clinical Significance of the 24-Hour Rhythm of Circulating Glucocorticoids. Endocr Rev 38:3-45.

Pavlova MK, Shea SA, Scheer FA, and Bromfield EB (2009) Is there a circadian variation of epileptiform abnormalities in idiopathic generalized epilepsy? Epilepsy Behav 16:461-467.

Phillips AJK, Vidafar P, Burns AC, McGlashan EM, Anderson C, Rajaratnam SMW, Lockley SW, and Cain SW (2019) High sensitivity and interindividual variability in the response of the human circadian system to evening light. Proc Natl Acad Sci U S A 116:12019-12024.

Pick R, He W, Chen CS, and Scheiermann C (2019) Time-of-Day-Dependent Trafficking and Function of Leukocyte Subsets. Trends Immunol 40:524-537.

Poggiogalle E, Jamshed H, and Peterson CM (2018) Circadian regulation of glucose, lipid, and energy metabolism in humans. Metabolism 84:11-27.

Pullman RE, Roepke SE, and Duffy JF (2012) Laboratory validation of an in-home method for assessing circadian phase using dim light melatonin onset (DLMO). Sleep Med 13:703-706.

Qian J, and Scheer F (2016) Circadian System and Glucose Metabolism: Implications for Physiology and Disease. Trends Endocrinol Metab 27:282-293.

Rahman SA, Kayumov L, Tchmoutina EA, and Shapiro CM (2009) Clinical efficacy of dim light melatonin onset testing in diagnosing delayed sleep phase syndrome. Sleep Med 10:549-555.

Rahman SA, Wright KP, Jr., Lockley SW, Czeisler CA, and Gronfier C (2019) Characterizing the temporal Dynamics of Melatonin and Cortisol Changes in Response to Nocturnal Light Exposure. Sci Rep 9:19720.

Rietveld WJ, Minors DS, and Waterhouse JM (1993) Circadian rhythms and masking: an overview. Chronobiol Int 10:306-312.

Roenneberg T, and Merrow M (2016) The Circadian Clock and Human Health. Curr Biol 26:R432-443.

Sack RL, Auckley D, Auger RR, Carskadon MA, Wright KP, Jr., Vitiello MV, Zhdanova IV, and American Academy of Sleep M (2007a) Circadian rhythm sleep disorders: part I, basic principles, shift work and jet lag disorders. An American Academy of Sleep Medicine review. Sleep 30:1460-1483.

Sack RL, Auckley D, Auger RR, Carskadon MA, Wright KP, Jr., Vitiello MV, Zhdanova IV, and American Academy of Sleep M (2007b) Circadian rhythm sleep disorders: part II, advanced sleep phase
disorder, delayed sleep phase disorder, free-running disorder, and irregular sleep-wake rhythm. An American Academy of Sleep Medicine review. Sleep 30:1484-1501.

Shafi AA, and Knudsen KE (2019) Cancer and the Circadian Clock. Cancer Res 79:3806-3814.

Shilts J, Chen G, and Hughey JJ (2018) Evidence for widespread dysregulation of circadian clock progression in human cancer. PeerJ 6:e4327.

Shuboni-Mulligan DD, Breton G, Smart D, Gilbert M, and Armstrong TS (2019) Radiation chronotherapy-clinical impact of treatment time-of-day: a systematic review. J Neurooncol 145:415-427.

Skeldon AC, Phillips AJ, and Dijk DJ (2017) The effects of self-selected light-dark cycles and social constraints on human sleep and circadian timing: a modeling approach. Sci Rep 7:45158.

Smith GC, Seaman SR, Wood AM, Royston P, and White IR (2014) Correcting for optimistic prediction in small data sets. Am J Epidemiol 180:318-324.

Smolensky MH, Hermida RC, Ayala DE, Mojón A, and Fernandez JR (2017) Bedtime Chronotherapy with Conventional Hypertension Medications to Target Increased Asleep Blood Pressure Results in Markedly Better Chronoprevention of Cardiovascular and Other Risks than Customary On-awakening Therapy. Heart Fail Clin 13:775-792.

St Hilaire MA, Klerman EB, Khalsa SB, Wright KP, Jr., Czeisler CA, and Kronauer RE (2007) Addition of a non-photic component to a light-based mathematical model of the human circadian pacemaker. J Theor Biol 247:583-599.

Stone JE, Aubert XL, Maass H, Phillips AJK, Magee M, Howard ME, Lockley SW, Rajaratnam SMW, and Sletten TL (2019a) Application of a Limit-Cycle Oscillator Model for Prediction of Circadian Phase in Rotating Night Shift Workers. Sci Rep 9:11032.

Stone JE, Phillips AJK, Ftouni S, Magee M, Howard M, Lockley SW, Sletten TL, Anderson C, Rajaratnam SMW, and Postnova S (2019b) Generalizability of A Neural Network Model for Circadian Phase Prediction in Real-World Conditions. Sci Rep 9:11001.

Thosar SS, Butler MP, and Shea SA (2018) Role of the circadian system in cardiovascular disease. J Clin Invest 128:2157-2167.

Ueda HR, Chen W, Minami Y, Honma S, Honma K, lino M, and Hashimoto S (2004) Molecular-timetable methods for detection of body time and rhythm disorders from single-time-point genome-wide expression profiles. Proc Natl Acad Sci U S A 101:11227-11232.

Vandewalle G, Middleton B, Rajaratnam SM, Stone BM, Thorleifsdottir B, Arendt J, and Dijk DJ (2007) Robust circadian rhythm in heart rate and its variability: influence of exogenous melatonin and photoperiod. J Sleep Res 16:148-155.

Viola AU, Simon C, Ehrhart J, Geny B, Piquard F, Muzet A, and Brandenberger G (2002) Sleep processes exert a predominant influence on the 24-h profile of heart rate variability. J Biol Rhythms 17:539-547.

Wittenbrink N, Ananthasubramaniam B, Munch M, Koller B, Maier B, Weschke C, Bes F, de Zeeuw J, Nowozin C, Wahnschaffe A, Wisniewski S, Zaleska M, Bartok O, Ashwal-Fluss R, Lammert H, Herzel H, Hummel M, Kadener S, Kunz D, and Kramer A (2018) High-accuracy determination of internal circadian time from a single blood sample. J Clin Invest 128:3826-3839.

Woelders T, Beersma DGM, Gordijn MCM, Hut RA, and Wams EJ (2017) Daily Light Exposure Patterns Reveal Phase and Period of the Human Circadian Clock. J Biol Rhythms 32:274-286.

Wright KP, Jr., Gronfier C, Duffy JF, and Czeisler CA (2005) Intrinsic period and light intensity determine the phase relationship between melatonin and sleep in humans. J Biol Rhythms 20:168-177.

Wu G, Ruben MD, Schmidt RE, Francey LJ, Smith DF, Anafi RC, Hughey JJ, Tasseff R, Sherrill JD, Oblong JE, Mills KJ, and Hogenessh JB (2018) Population-level rhythms in human skin with implications for circadian medicine. Proc Natl Acad Sci U S A 115:12313-12318.

Zeitzer JM, Dijk DJ, Kronauer R, Brown E, and Czeisler C (2000) Sensitivity of the human circadian pacemaker to nocturnal light: melatonin phase resetting and suppression. J Physiol 526 Pt 3:695-702.
Figure legends

Fig 1. A biomarker for which rhythm and what generates this rhythm?

Modified from (Bollinger and Schibler, 2014). Structure of the human circadian timing system. Molecular clocks and circadian rhythms are present in the brain including the SCN-based oscillator and periphery. A circadian biomarker may provide information about the rhythms in the SCN or in peripheral tissues and organs. Rhythms in organs and tissues are influenced by external rhythmic signals, SCN driven signals, local circadian oscillators and behavioural rhythms such as sleep or eating. A biological sample will contain many features (transcripts, proteins, metabolites). These various features will be influenced by SCN input, the local circadian oscillator and behaviour. The selection of the final feature set for the biomarker will depend on the purpose of the biomarker, e.g. assessing SCN phase or phase of tissue specific circadian oscillator. In many cases, the tissue or organ of interest will not be accessible and the features will be extracted from for example blood, which makes the identification of robust biomarkers even more challenging.

Fig 2. Effects of sleep-wake cycle and light exposure on rhythmic variables

Panels A Daily rhythm of core body temperature is altered when sleep occurs in phase (at night) vs. out of phase. Recalculated from (Dijk and Czeisler, 1995). Panel B Rhythms of plasma melatonin and cortisol are not much affected by sleeping in phase (during the night) or out of phase (during the day) [data from (Archer et al., 2014). Panel C Frequency distribution of the acrohases of rhythmic transcripts when sleeping in phase (blue) and out of phase. From: (Archer et al., 2014). Panel D Individual level dose-response curves for melatonin suppression and light levels. Blue High sensitivity individual; Red: Low sensitivity individual. From (Phillips et al., 2019).

Fig 3. Univariate multiple sampling vs multivariate single sample

A) A multivariate biomarker will require one or two samples separated by several hours, and by containing information about multiple rhythmic features the relative level of each of those features can classify the overall circadian timing (Laing et al., 2017). B) With a univariate biomarker, a time series of points assessing a single feature is collected. Depending on the variability of the feature, “noise” from periodic behaviors or physiologic changes and/or environmental changes, may influence any one data point or cycle, but multiple cycles of data will provide an accurate assessment of the underlying rhythmic process.

Fig 4. Examples of predictors of circadian melatonin phase and impact of sleeping ‘out of phase’ on accuracy of biomarker prediction

Panel A Prediction of plasma melatonin phase from two samples taken 12 hours apart across the circadian cycle during wakefulness (green symbols), nocturnal sleep (light blue symbols) and misplaced sleep (dark blue circles). From (Laing et al., 2017):

Panel B Prediction of salivary melatonin phase from one sample taken in the afternoon during wakefulness from extreme morning and evening types living on their habitual sleep-wake schedule. Men: triangles; Women circles. Size of circles indicates age of participants. From (Wittenbrink et al., 2018).
Panel C Prediction of urinary 6sulfatoxy melatonin phase from recordings activity, light exposure and a mathematical model for the effects of light in participants living on a ‘nocturnal’ schedule. From (Stone et al., 2019a)

Panel D Absolute error and its standard deviation of various biomarkers when tested on participants sleeping during the night (in phase) or during the day (out of phase) in either the laboratory or in a shift work situation. In all cases the biomarker-predicted phase was compared to a gold standard phase marker (plasma melatonin for the transcriptome predictors (Laing et al., 2017)) and urinary 6-sulfatoxy melatonin for the neural network (Stone et al., 2019b) and light model (Stone et al., 2019a). In all cases, accuracy was worse for the ‘out of phase’ condition.
Fig 1. A biomarker for which rhythm and what generates this rhythm?

- Environmental rhythmic signals e.g., Light-Dark
- SCN driven rhythmic signals e.g., cortisol, melatonin
- Rhythmic behaviours (e.g., sleep, eating)
- Rhythmic changes in physiological and biochemical variables (e.g., body temperature, nutrients)
Fig 2 Effects of sleep-wake cycle and light on rhythmic variables

A

B

C

D

https://mc.manuscriptcentral.com/jbrhythms
Fig 3 Univariate multiple sampling vs Multivariate single sample

[Image of a graph showing univariate and multivariate sampling with time in hours on the x-axis and variable values on the y-axis.]
Fig 4 Examples of predictors and effects of sleep displacement on performance

A

B

C

D

r=0.91, p=1.77x10^-5

Absolute error (minutes)

sleep in phase
sleep out of phase