Chronobiology Revisited in Psychiatric Disorders: From a Translational Perspective

Simge Seren Kirlioglu, MD, Yasin Hasan Balcioglu, MD
S.S.K. ORCID ID: 0000-0001-9778-6617
Y.H.B. ORCID ID: 0000-0002-1336-1724

Department of Psychiatry, Bakirkoy Prof Mazhar Osman Training and Research Hospital for Psychiatry, Neurology and Neurosurgery, 34147, Istanbul, Turkey

Author Note:
Please address correspondence and full-text requests to Yasin Hasan Balcioglu, MD, Department of Psychiatry, Bakirkoy Prof Mazhar Osman Training and Research Hospital for Psychiatry, Neurology, and Neurosurgery, 34147, Istanbul, Turkey.
E-mail: yhasanbalcioglu@gmail.com
Phone: 0090 212 409 1515
Submitted to: Psychiatry Investigation
Abstract 241 words, text 7351 words, 1 figure, 3 tables, 282 references

The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

The authors received no financial support for the research, authorship, and/or publication of this article.
Abstract

Objective: Several lines of evidence support a relationship between circadian rhythms disruption in the onset, course, and maintenance of mental disorders. Despite the study of circadian phenotypes promising a decent understanding of the pathophysiologic or etiologic mechanisms of psychiatric entities, several questions still need to be addressed. In this review, we aimed to synthesize the literature investigating chronobiologic theories and their associations with psychiatric entities.

Methods: The Medline, Embase, PsycInfo, and Scopus databases were comprehensively and systematically searched and articles published between January 1990 and October 2019 were reviewed. Different combinations of the relevant keywords were polled. We first introduced molecular elements and mechanisms of the circadian system to promote a better understanding of the chronobiologic implications of mental disorders. Then, we comprehensively and systematically reviewed circadian system studies in mood disorders, schizophrenia, and anxiety disorders.

Results: Although subject characteristics and study designs vary across studies, current research has demonstrated that circadian pathologies, including genetic and neurohumoral alterations, represent the neural substrates of the pathophysiology of many psychiatric disorders. Impaired HPA-axis function-related glucocorticoid rhythm and disrupted melatonin homeostasis have been prominently demonstrated in schizophrenia and other psychotic disorders, while alterations of molecular expressions of circadian rhythm genes including CLOCK, PER, and CRY have been reported to be involved in the pathogenesis of mood disorders.

Discussion: Further translational work is needed to identify the causal relationship between circadian physiology abnormalities and mental disorders and related psychopathology, and to develop sound pharmacologic interventions.

Keywords: biological clocks; circadian rhythm disorders; mental disorders; melatonin; HPA-axis

Highlights

- Sleep and circadian biorhythms are major physiologic functions responsible for emotional, cognitive, and somatic responses of the living organism.
- Mental disorders are often associated with disruptions in circadian rhythm functions.
- Molecular elements and expressions of genes including CLOCK, PER, and CRY, which are directly involved in the circadian system, are reported altered in many psychiatric disorders, particularly in mood disorders.
- Glucocorticoid rhythm supported by the hypothalamus–pituitary–adrenal (HPA) axis and melatonergic activity have a crucial role in the regulation of biorhythm, and oscillations of tissue and organ systems including the central nervous system, and both systems have been demonstrated impaired in major mental illnesses including schizophrenia and other psychotic disorders.
“There is a time for many words, and there is also a time for sleep.”

Homer, 850 BC

Introduction

Rhythmicity is a fundamental characteristic of the nature of life. Time as a dynamic and complex phenomenon, plays a pivotal role to sustain rhythmicity for the biologic essentials and needs of living organisms. Chronobiology aims to define basic principles of vital reactions that occur nearly 24 hours per day through circadian rhythms and biologic processes in anything from single cells to human beings. The first scientific awareness of circadian rhythms started with observations of the mimosa plant (*Mimosa pudica*) folding independent of daylight by the French astronomer Jean Jacques d'Ortous de Mairan, in 1729. In the 1930s, the German biologist Erwin Büning subsequently noticed that the movement of the bean plant had an intrinsic period that did not change under constant light conditions and inferred that such periodic alterations were arranged with an endogenous clock.

The term ‘circadian’ was first used by Franz Halberg in 1959. It means ‘about a day’ and an endogenous day slightly shorter or longer than 24 hours (from the Latin term circa: about and diem: day) depending on constant conditions, preserved from environmental factors. Uncovering interactions between molecules and cells within an endogenous day was a major advancement in the discovery of the essential mechanism of circadian rhythm, which was a remarkable scientific milestone in chronobiology. It had been eagerly attempted to explain the further molecular mechanisms of circadian rhythm; however, the oscillation process could not be unraveled until 1971. Konopka and Benzer first determined a gene by observing the differences of circadian period lengths among three mutant flies. They demonstrated three mutants, one was arrhythmic, another had a shorter period of 19 h, and the third had a longer period of 28 h; flies with neither the short-period gene nor the long-period gene or the arrhythmic gene would not produce a normal rhythm. They concluded that the same functional gene with a point mutation appeared to be affected in all cases. This work inspired Jeffery C. Hall, Michael Rosbash, and Michael Young, independently. They cloned and rescued the *Drosophila Per* gene, which was recognized as the first clock gene, found in 1984. They defined the transcriptional translational feedback loop (TTFL) model with the analysis of *Per* gene expression and they demonstrated additional genes and proteins in further work. The simple genetic model they postulated revealed the generation of an autonomous oscillator, including transcription-translation cycles from interacting positive and negative feedback loops that depend on ribonucleic acid (RNA) and protein levels, which is still used to understand circadian rhythms. Consequently, they were awarded the Nobel Prize in Physiology and Medicine in 2017 for their explanatory findings of molecular mechanisms controlling the circadian rhythm.

Despite the fact that the understanding of the neural basis of rhythmicity and central nervous system (CNS) involvement in circadian mechanisms is not long-standing knowledge, the discovery of the suprachiasmatic nucleus (SCN) of the anterior hypothalamus, which was later described as the master circadian pacemaker in mammals, is actually not very recent. The SCN was first defined as a cluster of different neurons in the 1880s and was subsequently recognized in a number of mammalian species’
brains through comparative studies of the hypothalamus by Crosby and Woodburne 7,8. However, the
discovery of its regulatory function on circadian rhythm occurred nearly 100 years later. The SCN
contains a complex neurochemical organization and its functional organization had been revealed with
comprehensive experimental studies regarding the function of localization, the neuronal mini-network it
contains, and its role in the circadian system. Consequently, the SCN is recognized as a coordinator of
biologic processes regulating numerous cellular clocks of the brain and other organ systems.

The findings of considerable studies revealing that a broad range of cell types in the body and brain have
biologic clocks raised questions regarding the specific function of circadian rhythm and its contribution
to illnesses. Circadian rhythms in peripheral organ systems and their impeccable relationship with the
SCN and other physiologic and metabolic mechanisms are essential for physical and mental health. The
internal desynchronization between the central and peripheral clocks which may be a result of shiftwork
or diversity of clock genotypes or circadian rhythm sleep disorders including delayed sleep phase
disorder, advanced sleep phase disorder, non-entrained type, irregular sleep-wake rhythm, shift work
sleep disorder and jet lag disorder have been associated with many illnesses including metabolic
dysfunctions, obesity, cancer, and mental disorders 9,10.

Circadian rhythms disruption refers to a range nosological penumbra that includes changes in phase and
amplitude of circadian rhythms, circadian misalignment, altered phase relationship between the sleep-
wake cycles and endogenous circadian rhythms. Therefore, we would like to provide brief information
about descriptions of circadian rhythms disruption types. The amplitude of a function is the distance
between the mean value and the peak. Therefore, the amplitude resembles half the range of oscillation.
Amplitude of the circadian rhythm is involved in the sleep-wake cycle and the changes in amplitude lead
to the changes of timing and consolidation of sleep and wakefulness 11. Phase is also one of the
parameters that characterize a circadian rhythm. The circadian pacemaker is known to drive a number of
physiologic variables including body temperature and the rhythms of melatonin and cortisol through
suprachiasmatic nuclei 12. The phase of the circadian rhythms is mostly determined by both genetic and
environmental factors such as routine daily activities and sunlight. The internal phase advance of
biological rhythms which reflects any disturbance of either sleep-wake cycle or endogenous circadian
rhythms is related to illnesses and aging 11. The term “circadian misalignment” describes a range of
circumstances, such as inappropriately timed sleep and wake, misalignment of sleep/wake with feeding
rhythms, or misaligned central and peripheral clocks 13. Another subtype of circadian misalignment
includes the misalignment of body rhythms with environmental cycles that is usually found in night-shift
workers characterizes a condition of chronic desynchronization similar to that produced by persistent jet
lag. Different types of circadian misalignment have been associated with increased risk for both physical
and psychiatric disorders 13.

Circadian rhythms abnormality, a common manifestation of nearly all psychiatric disorders, is not a
surprising predisposing factor for mental disorders, because sleep is considered as a cardinal
psychological and vital function and requires routine evaluation in every mental state examination. The
common disrupted mechanisms related to the circadian rhythm in psychiatric disorders could be determined as the melatonergic system, its effects of sleep pattern, and the hypothalamus–pituitary–adrenal (HPA) axis. Besides, studies of human circadian rhythm genes revealed that genetic polymorphisms of these genes predisposed to psychiatric disorders 14–16. Therefore, circadian disturbances seem to be the common thread to all these possible underlying mechanisms that contribute to illness onset, maintenance, and even the response to treatment. Special attention ought to be paid toward the physiology and pathology of circadian rhythm to understand the etiology of psychiatric disorders, and to develop appropriate treatment strategies because chronobiology is an essential field of work in mental disorders. Related literature provides information on circadian rhythm disturbances for certain psychiatric diagnoses such as schizophrenia, mood and anxiety disorders. However, we are aware of a lack of a comprehensive perspective of molecular and neural substrates of common disrupted circadian mechanisms to clinical manifestations in psychiatric disorders. There have been recent reviews relevant to the subject. For instance, Jones and Benca reviewed circadian disruptions in psychiatric disorders, however, they only focused on schizophrenia and mood disorders 17. Wulff and colleagues comprehensively reviewed sleep and circadian rhythms disruption in psychiatric disorders 18. Nevertheless, their review was lack of findings regarding current genetic, molecular and neurohumoral models of circadian pathologies. Therefore, we aimed to present a comprehensive review from a translational perspective regarding the reciprocal relationship between neurobiologic underpinnings of circadian rhythm pathologies and psychiatric disorders in this article.

Searching strategy and selection criteria of reviewed studies
An electronic database search was performed by the authors in the MEDLINE, Embase, PsycInfo, and Scopus databases for relevant articles published between January 1990 and October 2019. We searched reference lists of relevant reviews. Different combinations of the keywords psychiatric disorder, mental disorder, mood disorder, bipolar disorder, depression, unipolar depression, major depressive disorder, schizophrenia, psychotic disorders, anxiety disorders, circadian rhythms, circadian markers, chronotype, chronobiology, circadian gene, clock gene, melatonin, and HPA axis were polled. Articles published only in English were reviewed. Unpublished studies, case reports, theses, and conference papers were excluded. Several highly cited and regarded comprehensive review articles and meta-analyses are cited due to space considerations. Eligible open-access and institutional-access articles were recruited. The articles were filtered through an inspection of the abstracts in order to select the most suitable articles related to the topic. In addition to database searches, the reference lists of the relevant articles were also evaluated manually for additional publications matching the scope of our review. The authors avoided incorporating duplicated samples of the key papers; however, studies with similar methodology were included if they provide essential findings to the literature (Figure 1).

Molecular regulation of the circadian rhythm
We believe that it is noteworthy to briefly summarize the molecular underpinnings of circadian science that gave input to the research into neural substrates of rhythmicity. Although the aforementioned discovery of the period gene was a remarkable finding that identified a genetic determination of the
biological clock, it did not mean comprehension of all circadian molecular mechanisms. The circadian rhythm started to be more understandable with the determination of alterations in PER protein and period mRNA levels during a day. Hall and Rosbash ascertained that levels of period mRNA peaked in the early night, several hours earlier than the peak PER protein abundance. The TTFL model emerged with the discovery of further circadian rhythm genes found in subsequent studies. According to this model, PER and TIM (a protein encoded by the timeless gene) proteins transformed into a heterodimer form in the cytoplasm in order to translocate into the nucleus. TIM protein allows nuclear entry of PER. Besides CLOCK and CYCLE [orthologues of mammalian CLOCK and BMAL-1 (a protein encoded by the brain muscle ARNT-like protein-1 (Bmal-1) gene), respectively] constitute a protein couple that supports the transcription of period and timeless genes [the equivalent of period 1-3 and cryptochrome 1-2(Cry)] in mammalian cells] in the nucleus. When the PER-TIM heterodimer binds to the CLOCK-CYCLE couple, CLOCK-CYCLE segregates from DNA and the transcription of downstream genes related to PER and TIM conclude. In other words, the PER and TIM heterodimer terminate their transcription. However, in the event of a decrement in PER and TIM protein levels, the CLOCK and CYCLE couple activates their transcription once again, and TTFL starts over. All of these biochemical reactions include transcription and translation processes that occur rapidly. However, a near 24-h period needs a delay period and timeless gene transcriptions. The explanation about the regulation of the needed delay comes from the discovery of the doubletime gene, another member of the clock genes. The doubletime gene’s product casein kinase-1 (CSNK-1ε; casein kinase 1 epsilon in mammals) phosphorylates PER for degradation. Thus, activity of the doubletime gene reduces the stability and accumulation of PER, thereby promoting a delay between PER-TIM transcription and PER-TIM nuclear function. This molecular mechanism occurs both in the SCN and nearly all peripheral cells.

The maestro of chronophysiologic rhythms including body temperature, sleep-wake cycle motor activity, and neuroendocrine functions, is located in the SCN of the hypothalamus. The clock genes in the peripheral cells such as hepatocytes, adipocytes or epidermal and dermal cells have their own rhythmicity; however, cyclic processes in which the SCN is involved provide an integrative organization of the physiologic functions and behavioral outputs of the body. The circadian system sustains an endogenous rhythmic activity in spite of environmental cues. Regardless of the presence of light, the neuronal activity in the SCN occurs at a higher frequency during the day compared with the night. The neurons of the SCN tend to be excitable in the day to maintain spontaneous activity through persistent Na++ currents, oscillations in chloride pumps, K+ channels, and Ca++ pools in the morning. Conversely, hyperpolarized neurons are inhibited and keep the silence in the SCN at night. CRY and PER proteins gather in the cytoplasm before translocating into the nucleus where they inhibit CLOCK-BMAL-1 activity during the night. In other words, CRY and PER proteins terminate their own transcription when they inhibit CLOCK-BMAL-1 complex activity. After that, degradation of PER and CRY manages the inhibition of CLOCK-BMAL1 toward the morning, followed by resumed transcription of period/cryptochrome and other clock genes.

**Involvement of master clock in the regulation of circadian rhythm**
The master clock synchronizes the endogenous rhythm to the external world, mainly in the presence of major environmental input – light. A specialized tract, called the retino-hypothalamic tract, which starts from the retinal ganglion cells that include the essential photoreceptor pigment melanopsin, and terminating at the SCN. This tract aids upregulation of clock gene expression and increases neuronal activity in the SCN. Nevertheless, functions of the SCN, such as synchronization by the light/dark cycle, do not only depend on this molecular mechanism. Many inputs of the SCN have been determined including melatonin, food intake, blood pressure, and physical activity. In addition, the SCN receives non-photic timing inputs from the raphe nucleus, which means the serotoninergic system plays a substantial role in the regulation of circadian rhythm. Furthermore, the SCN serves in the excretion of numerous neurotransmitters that interact with other hypothalamic structures, hence neuropeptidergic signaling maintains circadian rhythm of the SCN. Consequently, the biologic interactions between the brain and body are modulated by the SCN, which is critically involved in the organism’s adjustment to the environment through the impact of internal signals, which are mediated by hormonal rhythms, the autonomic nervous system, and external time indicators such as light and food intake. The master clock regulates the endogenous rhythm in response to environmental inputs and dysfunction of the master clock could contribute to a wide range of illnesses including obesity, diabetes mellitus, autoimmune disorders, and particularly mental disorders. Disruption that arises due to a misalignment between inner physiology and the external world or a clock gene polymorphism may facilitate the emergence of diseases, increased disease severity and worsened prognosis, and heightened risk for poor treatment outcomes.

(Table 1 to be inserted here)

**Neurohumoral and hormonal regulation of circadian rhythm**

The SCN collects information about the endogenous clocks through nervous projections and peripheral hormones. The SCN’s monosynaptic outputs mainly target the pre-autonomic neurons of the paraventricular nucleus (PVN) in the hypothalamus. The SCN is directly involved in the hypothalamic output to the preganglionic parasympathetic regions of the brainstem and to sympathetic preganglionic motor neurons of the spinal cord. These projections allow the SCN to command the rhythmic control of hormone release and metabolism of all visceral structures through parasympathetic and sympathetic outputs. It has been determined that the SCN could increase glucose production from the liver through the sympathetic output to the liver with its projections that reach to the PVN. Similarly, the SCN could increase corticosterone secretion in the adrenal or support glucose uptake into the muscle cells via sympathetic activation. Besides, hormonal signals predominantly controlled by the SCN have a critical role in the regulation of internal synchronization. Internal synchronization is supplied by adrenal glucocorticoids, pineal melatonin, adipocyte-derived leptin, pancreatic insulin or stomach ghrelin induced by the SCN. Internal synchronization included many multi-synaptic neuronal pathways that modulate behavior. For example, leptin increases during food intake in rats, ghrelin increases following a fasting period, and adrenaline increases with locomotor activity.
Glucocorticoids are produced in the adrenal glands from cholesterol and rhythmically released at ultradian (pulsatile) and circadian (daily) scales. Glucocorticoid release peaks typically prior to the onset of physical activity and depends on the fluctuations of corticotropin (adrenocorticotropic hormone, ACTH), a polypeptide secreted from the anterior pituitary under the control of corticotropin-releasing hormone (CRH), during the day. Glucocorticoid levels are regulated by a complex interaction between the adrenal clock and sympathetic outputs from the PVN and SCN. Furthermore, the daily variation of glucocorticoids is influenced by stressful life events that activate the HPA axis and the autonomous nervous system. Glucocorticoid rhythm has a crucial role in the regulation of other hormonal rhythms and peripheral oscillations of metabolic gene expressions in the cells of tissues such as liver and white adipose tissue.

On the other hand, adrenal glucocorticoids can modulate the synchronization of the master clock to light via serotonergic projections from the raphe nucleus. Serotonergic neurons release serotonin in the presence of glucocorticoid and locomotor activity. Such neuronal activity ensures transmitting feedback to the SCN in order to sustain the functioning of the clock itself. In other words, serotonergic projections stimulated by locomotor activity provide a re-synchronization of the SCN. Furthermore, brain serotonin synthesis and catabolism have their own circadian rhythm, closely related to the SCN. Neuronal serotonin release in the SCN is provided in the absence of photic stimulation, and serotonin levels increase in the raphe nucleus after the beginning of the dark phase. Tryptophan hydroxylase (TpH), the rate-limiting enzyme in the synthesis of serotonin, is one of the regulators of circadian rhythm in the raphe nucleus. It is known that TpH peaks during the dark phase, helping the interaction between the serotonergic system and the SCN through the increment of serotonin levels. Also, serotonergic neurotransmission alterations could cause phase shifts and changes in SCN activity affecting the phosphorylation of CLOCK proteins.

Melatonin, a member of the class of acetamides, is another hormone related to biologic rhythm. It is primarily released by the pineal gland, particularly at night. Melatonin release is adjusted by the length of night time and melatonin per se regulates the seasonality of energy metabolism and reproduction in photoperiodic species. Melatonin secretion peaks a few hours before sleep or at the time of minimal vigilance propensity, and decreases as wakefulness approaches under normal conditions. In contrast, core body temperature reaches the highest degree during the day and has a nocturnal decline related to the melatonin peak. This inverse relationship between melatonin and core body temperature is organized by the SCN. The nocturnal release of melatonin is induced by the SCN input to the PVN noradrenergic (sympathetic) afferents to the pineal gland. Melatonin accumulates sleep both by setting the SCN and inhibiting neural centers such as the locus coeruleus (LC) and raphe nuclei, which mediate arousal through the ventrolateral preoptic nucleus of the hypothalamus (VLPO). It has been determined that melatonin receptor agonists increase monoaminergic neuronal activity and contribute to the regulation of dopamine and 5-HT neurotransmission. In other words, melatonin has a modulatory role on the monoaminergic activity by linking the circadian and monoamine systems. The SCN modulates the release of melatonin mainly through γ-aminobutyric acid (GABA) neurons that project from the SCN to the PVN. The daylight in the morning and the bright light in the evening activate the SCN neurons.
that inhibit the same PVN neurons through GABAergic projections and cease the secretion melatonin. The daily rhythm of melatonin has remarkable effects on the molecular clockworks of both the brain and body alongside regulating the sleep/wake cycle. Melatonin receptors (MT1 and MT2) are mainly localized in the CNS but also have been detected beyond the CNS in a wide range of somatic cells. This diversity could be interpreted as melatonin having an integrative role in the light-induced circadian rhythms controlled by the SCN in the whole organism.

**Circadian rhythm and its implications on psychiatric disorders**

At the core of any psychiatric disorder is an abnormality in neurotransmitter signaling. It is well known that the disruption of circadian physiology has widespread effects on all aspects of neural and neuroendocrine function, which leads to psychiatric disorders. The aforementioned information regarding neural substrates of biologic rhythm is frequently reported impaired in many mental disorders. Following the comprehensive conceptual framework of neural substrates of chronobiologic processes mentioned above, we will next discuss the reciprocal associations between circadian rhythm disturbances and psychiatric disorders, and draw a clinical picture for common diagnoses.

(Table 2 to be inserted here)

**Mood disorders**

In 1681, Robert Burton defined the autumn as the most melancholic season in his best-known classic, The Anatomy of Melancholia. Circadian rhythm abnormalities in mood disorders have been pointed towards by the observers of melancholia for sixty years. A wide range of body functions such as core body temperature, blood pressure, pulse rate, and hormones such as plasma cortisol levels, thyroid-stimulating hormone, and melatonin have been found disturbed in patients with manic depression compared with people without a mental disorders. Moreover, mood and other symptoms of the disorder have been previously reported to show diurnal variation in depression. Disordered sleep/wake cycle is considered as another clue for physicians in patients with bipolar disorder (BD) and major depressive disorder (MDD). In addition, it was recognized that disrupted rhythms were resynchronized after antidepressant or mood-stabilizing treatment. Another significant feature is that mood episodes recur seasonally and previous studies showed that there could be an association between light and the emergence of mood states. Thus, all of these findings suggested the possibility of circadian rhythm disturbance in mood disorders. Consequently, the earliest mention of seasonality took place in the Diagnostic and Statistical Manual of Mental Disorders Third Edition, Revised Version (DSM-III-R), and seasonal pattern was defined as a specifier in the affective disorders section.

Chronotype is another concept associated with mental disorders, particularly with affective disorders, and resembles individual physiologic functions and activities such as sleeping, eating, or hormone release. Chronotype has usually been used to denote sleep habits: morning and evening types. The relationship between chronotypes and several psychiatric disorders has been studied to date and the evening
chronotype has been related to a vulnerability to depression and increased alcohol and stimulant drug use.

Although sleep/wake cycle alteration, which is considered as a consequence of circadian system disruption, had been the best-known contributor to the pathophysiology of mood disorders for years, today, it is well-recognized that circadian rhythm is entangled with a wide range of molecular and cellular processes that are hypothesized to lead to mood disorders. Accordingly, below we discuss in detail internal and external factors that may play a role in the emergence of mood disorders through various psychophysiological mechanisms within the circadian rhythm processes.

**Major depressive disorder**

As a cardinal element of chronobiologic processes, sleep behavior and its disturbances have received the strongest spotlight regarding research into their undisputed etiologic and prognostic association with mood disorders. The concomitance of sleep disruption and depression had been the main focus of research into the contribution of circadian rhythms disruption to depression development since the 1970s. The relationship between sleep and mood could easily be observed even in healthy individuals exposed to jet lag or shiftwork. The presence of sleep disruption may cause negative effects, irritability, and fatigue. Sleep behavior changes, such as difficulties in initiating/maintaining sleep or early morning awakening have been determined in 90% of patients with MDD. Sleep-wake disruptions are among the criteria for the diagnosis of depression, and comorbid parasomnias are associated with poor treatment outcomes, increased suicidality, and greater relapse risk in depression. Sleep architecture alterations including shortened latency of the initial rapid eye movement (REM) sleep, prolonged first REM period, increased total REM time, increased REM density and proportion of REM sleep, and decreased non-REM sleep have been demonstrated in depression. It has been suggested that there is a reciprocal relationship between sleep-wake cycle variables such as wakefulness and sleep latency and sleep architecture features like REM sleep latency. For instance, preclinical studies have consistently demonstrated that prolonged REM sleep was associated with decreased wakefulness in depressive subjects. In addition, an endogenous circadian rhythm abnormality, the phase advance, is related to decreased REM latency after falling asleep among individuals with depression. These findings suggest that sleep itself has multiple and complex regulators related with homeostatic mechanisms along with the endogenous circadian rhythm.

Melatonin output and the timing of its release have been found closely associated with other rhythms as mentioned above. Numerous studies have been conducted to show alterations of melatonin release and its phase to determine circadian misalignment between internal and external clocks in patients with mood disorders. To date, the most consistent results suggested lower nocturnal melatonin levels, delayed melatonin secretion onset, and offset in patients with depression. Besides, the length of the interval between melatonin secretion and sleep onset has been found related to depression severity. However, a few studies demonstrated increased nocturnal melatonin levels in depressive patients. Such conflicting findings might be related to the heterogeneity of the study group and inclusion of depressive...
patients with psychotic symptoms. Abnormal body temperature variations including the absence of the nocturnal decline of core body temperature and daily mean temperature degrees are also observed in patients with depression and these higher values normalized with antidepressant treatment. This is probably due to the impaired control of the melatonergic system over thermoregulatory processes in depressive patients.

There is an irrefutable association between circadian genes and mood regulation. Even though mood disorders are not directly related to clock gene mutations, findings suggest that individual genetic polymorphisms of clock genes may influence the clinical features of the disorder, such as age at disease onset and treatment response. Genetic studies have implicated clock, timeless, cryptochrome-1 (Cry-1), period-2,3 (Per-2,3), Bmal-1,2, neuronal pas domain protein 2 (Npas-2), nuclear receptor subfamily-1, group d, member 1 (Nrl1d-1), retinoid-related orphan receptor a (Rora), CSNK-Ic, D site of albumin promoter binding protein (Dbp), acetylserotonin methyltransferase (Asmt), melatonin receptor 1b (Mnr1-B), arylalkylamine N-acetyltransferase (Aanat) genes in unipolar depression. However, most of these studies have small sample sizes and need to be replicated in larger groups.

Glucocorticoids are adrenal steroid hormones and have multifunctional roles in the body and brain such as metabolism, immunity, arousal, neuronal survival, and neurogenesis. Glucocorticoids have their own circadian rhythm and an important role in synchronizing peripheral clocks and the SCN. In addition, they have anti-inflammatory properties and regulate the immune system response. Since Carroll defined the resistance of the dexamethasone suppression test in patients with depression in 1968, hypothalamic-pituitary-adrenal (HPA) axis dysregulation has been one of the most consistent findings in mental disorders, particularly in depression. Hypercortisolemia-flattened HPA axis circadian rhythm and disrupted response of the HPA axis to glucocorticoid feedback are commonly observed in patients with depression. Dehydroepiandrosterone (DHEA), is another adrenal steroid that has a neuroprotective role and modulates corticosterone-induced cell death. An increased cortisol/DHEA ratio, which assesses the degree of ‘functional’ hypercortisolemia, is seen in adults and adolescents with depression. Glucocorticoid receptor hypofunction has also been found in peripheral tissue cells including mononuclear cells and skin cells. Furthermore, findings support that antidepressant treatment repairs the impaired HPA axis dysfunction in depression.

Depression and inflammatory disorders such as rheumatoid arthritis, inflammatory bowel disease, and asthma have been found coexisting, and such common comorbidities point to the neuroinflammatory background and immune-associated contributions in the etiopathogenesis of depression. Studies have also shown that pro-inflammatory cytokines could induce a depression-like symptom cluster including anhedonia, fatigue, increased sleep, and decreased locomotor activity. Inflammatory markers such as interleukin (IL)-1β, IL-2, IL-6, tumor necrosis factor (TNF)-α, C-reactive protein (CRP), and prostaglandin E2 (PGE2) have been reported increased in patients with depression. Sleep-wake cycle changes and circadian misalignment between the internal and external clocks may be other contributors to increased pro-inflammatory cytokine levels in depression. The arrhythmic clock system interacts with the nuclear factor-kappa B (NF-kB) signaling pathway, which is one of the major
regulators of inflammation in the body and activates the inflammatory response \(^{127,128}\). Besides, sleep disturbances and long sleep duration were found related with the increased cytokines levels and the risk for depression \(^{129}\). We may interpret the aforementioned findings as the circadian system’s involvement in the pathophysiology of MDD being not limited to sleep/wake cycle disruption, it is also related to complex associations between biologic rhythm, environment-gene interactions, HPA axis dysfunction, and immune system alterations.

**Bipolar disorder**

Sleep disturbances have been the core common characteristic feature in bipolar mood episodes, both mania and depression, since the first definition of Kraepelin \(^{130}\). In turn, insomnia or hypersomnia and decreased need for sleep are typical for manic and depressive episodes. Studies showed that sleep architecture was characterized by increased REM density and reduced REM latency in bipolar manic episodes \(^{131}\). Sleep disturbances are also frequently observed in euthymic patients with BD. Increased REM density and the proportion of REM sleep have been shown in remitted patients with BD \(^{132}\). Moreover, findings revealed that remitted patients with BD have longer sleep latency and sleep duration and lower sleep efficiency \(^{133,134}\). Bipolar depression has similar polysomnographic findings including a tendency for more early awakenings and more fragmented REM sleep periods. However, total REM density was found greater in bipolar depression than in unipolar depression \(^{132}\) (See table 2 for detailed information). Although abnormalities of sleep architecture are seen in episodes and inter-episodes, sleep disturbances worsen before relapses. Sleep loss and reduced sleep duration were defined as reliable predictors of hypomania and mania \(^{132}\). In addition, hypersomnia in euthymia is found associated with the development of upcoming depressive symptoms \(^{135}\). On the other hand, a large amount of euthymic patients describe symptoms that meet the diagnostic criteria for insomnia \(^{134,136}\). Sleep-wake disturbances have been found as one of the reasons for a worse course of illness, relapses, increased symptom severity, and poor treatment outcomes \(^{137-140}\). These findings may explain the reason for the treatment need in remitted patients with BD \(^{89}\).

Involvement of the melatonergic system in the pathogenesis of BD through circadian dysregulations such as changes in the release timing, melatonin rhythm, and the sleep-wake cycle \(^{132}\). In terms of the phase relationship between the sleep-wake cycle and melatonin rhythm, it has been demonstrated that the melatonin secretion in the hours surrounding habitual sleep onset which is considered as a crucial circadian moment for sleep regulation was increased in patients with BD compared to those with unipolar depression \(^{141}\). Although findings of melatonin function in patients with BD are inconsistent, circadian system characteristics generally vary depending on the current episode; mania or depression \(^{82}\). Melatonin levels were found higher in the daytime in manic patients than in healthy controls and patients with depressive episode \(^{142}\). Findings about nocturnal melatonin levels among BD phases are not consistent \(^{72,143-145}\). It remains unclear as to whether these alterations derive from a primary dysfunction of the circadian rhythm or if they are secondary to sleep disturbances related to the BD episode. However,
some studies supported the beneficial effect of exogenous melatonin administration, which provides sleep and mood improvement.  

Some of the clock genes have been found intimately associated with both the onset of BD and illness course. Studies revealed that circadian gene polymorphisms may increase the predisposition to BD and indirectly affect recurrences and symptoms across all BD phases. Genetic linkage and gene expression studies implicated the variant genes related to BD as clock, timeless, Cry-1, Npas-2, Bmal-1, Dhp, Nrl-1, Per-2,3, Rora, Rorb, Asmt, Csnk-1ε, Csnk-1δ, and glycogen synthase kinase-3β (GSK-3β). It has been demonstrated that ClockD19, the mutant gene that occurs with the deletion of exon 19 in the Clock gene, produces a dominant negative CLOCK protein capable of DNA binding but deficient in transcriptional activity. This gene induces dopamine synthesis and increased dopaminergic activity, which result in an increase in tyrosine hydroxylase (TH) expression in the ventral tegmental area (VTA) and manic-like behavior in animal models. Moreover, ClockD19-related higher dopaminergic activity in the VTA normalized after lithium treatment, which suggests increased dopaminergic activity may be the main reason for the manic-like behavior of mice. Recently, several lines of evidence have emphasized the importance of the molecular and synaptic mechanisms of monoaminergic systems and circadian gene interactions, which are closely related to molecular alterations associated with the ClockD19 model in the VTA and nucleus accumbens. On the other hand, lithium, a potent inhibitor of the GSK-3 enzyme, regulates the clock gene Nrl-1 and BMAL-1 through GSK-3. Some polymorphisms including Clockrs3805148, Clockrs534654, Timelessrs11171856, and Timelessrs2291739 are associated with suicidal behavior in BD.

A dysfunctional HPA axis is suggested to play an important role in the pathophysiology of BD, although the mechanism needs to be elucidated. Increased levels of cortisol and ACTH are the most replicated findings in BD. However, CRH levels are not determined to increase in BD. Depressive symptoms and cognitive deficits are thought to be associated with the higher levels of cortisol, and ACTH and cortisol seem to be related to manic episodes. A meta-analysis suggested that abnormalities of stress-related pathways including increased morning cortisol levels were mainly prominent in manic episodes. Such abnormalities are even observed in remitted patients, which means that the long-term pathology of the HPA axis is related to clinical states of BD and contributes to the stress-vulnerability models of illness development and progression.

Immune abnormalities have received increased attention due to their possible role in the pathophysiology of BD, as well as MDD. Systematic reviews on cytokine levels in patients with BD revealed that IL-4, IL-6, IL-10, soluble IL-2 receptor, soluble IL-6 receptor, and TNF-α levels were increased in patients compared with healthy controls, whereas IL-2, IL-8, IFN-gamma, and C-C motif ligand were not different from controls. Moreover, a comparison of cytokine levels in another study determined that proinflammatory cytokines including IL-2, IL-4, IL-6 were higher during manic episodes, and IL-6 levels were higher in depressive state than in healthy controls. It was also demonstrated that mood symptoms had a positive correlation with IL-6 and IL-2 levels. When bipolar depression and unipolar depression were compared, sIL-6R, CRP, sTNF-R1, and monocyte chemoattractant protein-1 (MCP-1) were found
at higher levels than in unipolar depression \textsuperscript{163}. In conclusion, sleep disturbances may be a reliable indicator of an upcoming mood episode in BD.

\textbf{Schizophrenia}

Although the relationship between mood disorders and circadian abnormalities has become clearer in recent times, the links between schizophrenia and disrupted circadian rhythms have yet to be elucidated fully. However, sleep disorders and sleep-wake cycle alterations have been known as common and consistent features of schizophrenia and other psychotic disorders since the first definition of Kraepelin in 1883 \textsuperscript{164}. Schizophrenia has been associated with abnormalities in sleep including delayed and advanced sleep onset, altered resting activity patterns, and irregular sleep-wake cycle \textsuperscript{165}. Research into circadian abnormalities and sleep disruption in schizophrenia has attempted to explain the causal relationship in a reciprocal context. Hyperdopaminergia is a well-known phenomenon in psychosis syndromes and striatal hyperdopaminergic activity may be a result of sleep disruption and circadian abnormalities, and increased dopamine levels may induce sleep disruptions \textsuperscript{166–168}. For instance, the \textit{Clock T3111C} polymorphism, which is associated with increased dopamine levels in the SCN, has been determined in a population of Japanese patients with schizophrenia \textsuperscript{171}. Furthermore, the blind-drunk mutant mouse, which carries a mutation in the gene encoding an exocytic synaptic protein, synaptosomal-associated protein-25 (Snap-25), exhibits schizophrenia-like symptoms \textsuperscript{169,170}. This mouse model of schizophrenia has been shown to display phase advance and fragmentation of the circadian cycle \textsuperscript{171}. Most consistent findings of the circadian genetics studies have been associations between \textit{CLOCK}, \textit{PERIOD1}, \textit{PERIOD3}, and \textit{TIMELESS} genes and schizophrenia \textsuperscript{172}. Circadian rhythms disruption has been reported in approximately 80\% of patients with schizophrenia \textsuperscript{173}. Abnormal sleep patterns in schizophrenia have been described in both unmedicated patients and patients currently receiving antipsychotic treatment \textsuperscript{174}. The major findings in sleep architecture could be aligned, such as long sleep-onset latency, increased intermittent-awakenings, decreased total sleep time, and poor sleep efficiency \textsuperscript{174}. Moreover, reductions in REM latency, REM density, and duration of non-REM Stage 4 are other alterations in micro-sleep architecture \textsuperscript{17,18,175–177}. Sleep disturbances are also important to predict increased suicide attempts in patients with schizophrenia \textsuperscript{178}.

Melatonin is a versatile neurohormone that plays an important role in the pathophysiology of schizophrenia. \textit{5-HT} synthesis regulation, sleep-wake cycle, and anti-oxidant effects against neuroinflammation are impaired due to melatonin dysfunction in schizophrenia \textsuperscript{167,179}. It has been shown that melatonin increases endogenous antioxidants by increasing phosphorylated glycogen synthase kinase-3 (GSK-3) levels and provides an anti-inflammatory effect \textsuperscript{179,180}. Galván-Arrieta et al. reported a reduction in axogenesis associated with lower levels of phosphorylated GSK-3 subtype β and less expression of melatonergic receptors in patients with schizophrenia compared with healthy controls. These findings may indicate a melatonin-derived neurodevelopmental deficit at a cellular level \textsuperscript{181}. A lack of normal melatonin rhythmicity, decreased nocturnal secretion of melatonin, and phase advance in melatonin circadian rhythms have also been described in patients with schizophrenia \textsuperscript{167,179,182}. Additionally, pineal calcification in computed tomography has been demonstrated in patients with...
schizophrenia, and this structural change has been found associated with cortical atrophy \textsuperscript{183}. Mainly, the clinical importance of the relationship schizophrenia and melatonergic dysfunction might come from the fact that impairment of sleep-wake cycles which are in close relationship with melatonin phase are associated with schizophrenia symptoms such as somatic complaints, anxiety, depression, and paranoia. Further support has shown that sleep-deprived schizophrenia patients might exhibit increased psychotic symptoms \textsuperscript{167}. Moreover, the circadian rhythm of dopamine is dependent on melatonin \textsuperscript{184}. A combination of sleep disruption and circadian rhythms disturbance may lead to the elevation of dopamine activity in the brain, both directly and through phase advance in melatonin circadian rhythm \textsuperscript{167}. Preclinical studies have been also demonstrated that melatonergic receptor agonism may prevent the increase of glutamate release in prefrontal cortex \textsuperscript{185}, which has been suggested in the pathophysiology of schizophrenia, particularly of cognitive symptoms of the disorder \textsuperscript{186}. Because of its significance in the pathogenesis of schizophrenia, melatonin has become a therapeutic target for researchers. It has been shown that melatonin agonists are efficacious agents for schizophrenia-associated sleep disorders and drug-related tardive dyskinesia \textsuperscript{187,188}. Moreover, its improving effects on behavioral deficits via reducing brain oxidative stress have been shown in an animal model of schizophrenia \textsuperscript{189}.

The relationship between clock genes and schizophrenia is another undiscovered area for scientists. Few studies have been conducted to show linking circadian clock gene polymorphisms in schizophrenia to date. Takao et al. identified the Clock 311C/T polymorphism, which is associated with higher dopaminergic neurotransmission in the SCN in patients with schizophrenia \textsuperscript{14}. These results were confirmed in another study conducted in a Chinese schizophrenic population \textsuperscript{190}. \textit{Period-1} mRNA expression in the temporal lobe of post-mortem subjects with schizophrenia was found down-regulated when compared with healthy controls \textsuperscript{191}. In addition, disrupted diurnal rhythms of the \textit{Per-1}, \textit{Per-2}, \textit{Per-3}, \textit{Npas-2} and phase delay in the expression of \textit{Per-2} have been reported in white blood cells of patients with schizophrenia \textsuperscript{192}. More recently, the absence of rhythmic expression of \textit{Cry-1} and \textit{Per-2} was determined in the fibroblasts of patients with schizophrenia compared with cells obtained from healthy controls.\textsuperscript{193} Pinacho et al. reported decreased levels of CSNK1\textepsilon{} protein levels in the prefrontal cortex of patients with schizophrenia \textsuperscript{194}. However, due to the small sample sizes of the available studies, the association between schizophrenia and clock genes still needs to be clarified with further studies with larger populations.

The stress-vulnerability model for schizophrenia was first proposed in the 1970s and has been further developed since that time \textsuperscript{195,196}. Thus, the HPA axis has been one of the most attractive research targets to understand the pathophysiology of schizophrenia for decades. Increased cortisol levels have been determined in patients with schizophrenia and even in individuals at high risk for schizophrenia compared with controls \textsuperscript{197-199}. However, mean baseline cortisol level measurements in schizophrenia are not consistent in the literature \textsuperscript{200}. Nevertheless, blunted cortisol levels in response to stressors are much more consistent findings, regardless of disease stage, chronicity, and treatment condition \textsuperscript{201}. To conclude, despite it being widely accepted that sleep and circadian disorders have an important role in the etiopathogenesis of schizophrenia, well-designed and comprehensive clinical studies are still needed to explicate the genetic and neurobiologic underpinnings.
Other Psychiatric Disorders

Anxiety disorders are seen as the most frequent type of psychiatric disorders with a lifetime prevalence of 29% in the general population. Sleep disturbance is a common feature of anxiety disorders and is included in the symptom criteria for several anxiety disorders such as post-traumatic stress disorder and generalized anxiety disorder. The presence of sleep disturbances has been reported as 74% in patients with anxiety disorders. However, MDD as a frequent comorbid condition in anxiety disorders is a confounder in understanding the relationship of sleep disturbances and anxiety disorders. Studies related to generalized anxiety disorder have reported decreased total sleep time, increased sleep-onset latency, and alterations in non-REM sleep architecture, whereas findings of REM sleep and sleep efficiency are inconsistent. Patients with panic disorder frequently have both sleep disorder and/or another anxiety disorder because they could have nocturnal panic attacks, which usually occur in Stage-2 or Stage-3 of non-REM sleep, as well as decreased sleep efficiency, total sleep time, and increased sleep onset latency. Although sleep disturbances, including REM sleep-related nightmares, have been investigated in post-traumatic stress disorder, conclusions are not consistent. There is no significant difference in sleep architecture in social anxiety disorder. In an animal model, Cry-1 and Cry-2 gene protein deficiencies led to behavioral alterations characterized by an abnormally high level of anxiety. Akiyama et al. suggested that period-1 mRNA levels reduced after anti-anxiety treatment in the mouse cerebellum. Cry-2 expression was determined reduced in the hippocampus in another animal study. Furthermore, a polymorphism in BMAL-2rs2306073 has been found associated with social phobia.

Obsessive-compulsive disorder (OCD) is another debilitating disorder that is segregated from the anxiety disorders category in the DSM-5. Although sleep disturbances have been reported including decreased total sleep time, alterations in REM and non-REM sleep architecture are less clear. Certain chronotypes have been found as predictors of OCD symptoms in adults, and circadian rhythm disorders have been found as predictors of treatment outcomes. To the best of our knowledge, the role of circadian rhythms disruption in all anxiety disorders, including OCD, has yet to go beyond showing sleep disturbance; comprehensive research is warranted in the context of chronobiologic mechanisms of anxiety disorder pathology.

(Table 3 to be inserted here)

Conclusion

The circadian system is responsible for the temporal organization of physiologic functions, and disruptions can have marked functional influences on the living organism. As the role of chronobiologic systems in both physical and mental health have become better understood, research into neurobiologic mechanisms of circadian rhythms has been expanded. Mood, cognition, and behavior have complex relationships with biologic rhythms, and the vast majority of mental disorders are reciprocally associated...
with impaired circadian biology. Extensive research has shown that impaired circadian mechanisms could lead to psychiatric entities, whereas they may be an outcome of mental disturbances. Impaired HPA axis function and melatonin homeostasis are the most consistent findings in mental disorders. Independent from sleep disorders, the circadian system has a distinct role in homeostatic processes, whose impairment has an impact in emotion regulation, cognition, behavior, and, most importantly, neural plasticity, all of which are often disrupted in psychiatric phenotypes. There is some evidence suggesting that circadian rhythm genes are associated with psychiatric disorders; however, the specificity and causality of these associations have yet to be made clear. In our opinion, we are a long way from establishing a robust causative link between circadian rhythms disruption and phenotypic complexity of psychiatric disorders. A decent translational approach to the findings of animal models would likely result in a clearer understanding of pathophysiologic implications of the circadian system. Further support from continued and integrated investigations of these issues may promote a deeper appreciation of the contribution of circadian disturbances to the pathophysiology of psychiatric illnesses and related psychopathology, and will hopefully yield improved therapeutic strategies for their treatment.

**Conflicts of interest**
The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

**Funding**
The authors received no specific grant from any funding agency, commercial or not-for-profit sectors for the research, authorship, and/ or publication of this article.

**Author contributions**
Dr. Simge Seren Kirlioglu (SSK) and Dr. Yasin Hasan Balcioglu (YHB) conceived the presented idea and contributed equally to concept development and designing of the article. The screening of the literature was performed by SSK. Overall, while SSK drafted the article, both authors were involved in reviewing and revising the article and contributed to the intellectual content. SSK generated the tables, while YHB did the figure. YHB also performed a critical review and made necessary corrections. Both SSK and YHB approved the final version of the manuscript prior to submission.
References

1. Foster RG, Kreitzman L. *Rhythms of Life: The Biological Clocks That Control the Daily Lives of Every Living Thing*. New Haven (Connecticut): Yale University Press; 2005.

2. Halberg F, Cornélissen G, Katinas G, Syutkina E V, Sothern RB, Zaslavskaya R, et al. Transdisciplinary unifying implications of circadian findings in the 1950s. *J Circadian Rhythms*. 2003;1:2.

3. Konopka RJ, Benzer S. Clock mutants of Drosophila melanogaster. *Proc Natl Acad Sci U S A*. 1971;68:2112-2116.

4. Bargiello TA, Jackson FR, Young MW. Restoration of circadian behavioural rhythms by gene transfer in Drosophila. *Nature*. 1984;312:752-754.

5. Reddy P, Zehring WA, Wheeler DA, Pirrotta V, Hadfield C, Hall JC, et al. Molecular analysis of the period locus in Drosophila melanogaster and identification of a transcript involved in biological rhythms. *Cell*. 1984;38:701-710.

6. Huang R-C. The discoveries of molecular mechanisms for the circadian rhythm: The 2017 Nobel Prize in Physiology or Medicine. *Biomed J*. 2018;41:5-8.

7. Crosby EC, Woodburne RT. The mammalian midbrain and isthmus regions. Part II. The fiber connections. C. The hypothalamo-tegmental pathways. *J Comp Neurol*. 1951;94:1-32.

8. Sollars PJ, Pickard GE. The Neurobiology of Circadian Rhythms. *Psychiatr Clin North Am*. 2015;38:645-665.

9. Zhu L, Zee PC. Circadian Rhythm Sleep Disorders. *Neurol Clin*. 2012;30:1167-1191.

10. Gillette M. Chronobiology: Biological Timing in Health and Disease. *Prog Mol Biol Transl Sci*. 2013;376.

11. Dijk DJ, Duffy JF, Czeisler CA. Contribution of circadian physiology and sleep homeostasis to age-related changes in human sleep. *Chronobiol Int*. 2000;17:285-311.

12. Panda S. Circadian physiology of metabolism. *Science (80- ).* 2016;354:1008-1015.

13. Baron KG, Reid KJ. Circadian misalignment and health. *Int Rev Psychiatry*. 2014;26:139-154.

14. Takao T, Tachikawa H, Kawanishi Y, Mizukami K, Asada T. CLOCK gene T3111C polymorphism is associated with Japanese schizophrenics: A preliminary study. *Eur Neuropsychopharmacol*. 2007;17:273-276.

15. Benedetti F, Serretti A, Colombo C, Barbini B, Lorenzi C, Campori E, et al. Influence ofCLOCK gene polymorphism on circadian mood fluctuation and illness recurrence in bipolar depression. *Am J Med Genet*. 2003;123B:23-26.

16. Lee KY, Song JY, Kim SH, Kim SC, Joo E-J, Ahn YM, et al. Association between CLOCK
3111T/C and preferred circadian phase in Korean patients with bipolar disorder. *Prog Neuro-Psychopharmacology Biol Psychiatry*. 2010;34:1196-1201.

17. Jones SG, Benca RM. Circadian disruption in psychiatric disorders. *Sleep Med Clin*. 2015;10:481-493.

18. Wulff K, Gatti S, Wettstein JG, Foster RG. Sleep and circadian rhythm disruption in psychiatric and neurodegenerative disease. *Nat Rev Neurosci*. 2010;11:589-599.

19. Hardin PE, Hall JC, Rosbash M. Feedback of the Drosophila period gene product on circadian cycling of its messenger RNA levels. *Nature*. 1990;343:536-540.

20. Gekakis N, Saez L, Delahaye-Brown AM, Myers MP, Sehgal A, Young MW, et al. Isolation of timeless by PER protein interaction: defective interaction between timeless protein and long-period mutant PERL. *Science*. 1995;270:811-815.

21. Allada R, White NE, So WV, Hall JC, Rosbash M. A Mutant Drosophila Homolog of Mammalian Clock Disrupts Circadian Rhythms and Transcription of period and timeless. *Cell*. 1998;93:791-804.

22. Price JL, Blau J, Rothenfluh A, Abodeely M, Kloss B, Young MW. double-time Is a Novel Drosophila Clock Gene that Regulates PERIOD Protein Accumulation. *Cell*. 1998;94:83-95.

23. Kloss B, Price JL, Saez L, Blau J, Rothenfluh A, Wesley CS, et al. The Drosophila Clock Gene double-time Encodes a Protein Closely Related to Human Casein Kinase Iε. *Cell*. 1998;94:97-107.

24. Lowrey PL, Shimomura K, Antoch MP, Yamazaki S, Zemenides PD, Ralph MR, et al. Positional syntenic cloning and functional characterization of the mammalian circadian mutation tau. *Science*. 2000;288:483-492.

25. Mohawk JA, Green CB, Takahashi JS. Central and Peripheral Circadian Clocks in Mammals. *Annu Rev Neurosci*. 2012;35:445-462.

26. Challet E. Keeping circadian time with hormones. *Diabetes, Obes Metab*. 2015;17:76-83.

27. Colwell CS. Linking neural activity and molecular oscillations in the SCN. *Nat Rev Neurosci*. 2011;12:553-569.

28. Hankins MW, Peirson SN, Foster RG. Melanopsin: an exciting photopigment. *Trends Neurosci*. 2008;31:27-36.

29. Asher G, Sassone-Corsi P. Time for Food: The Intimate Interplay between Nutrition, Metabolism, and the Circadian Clock. *Cell*. 2015;161:84-92.

30. Bujs FN, Cazarez F, Basualdo MC, Scheer FAJL, Perusquía M, Centurion D, et al. The suprachiasmatic nucleus is part of a neural feedback circuit adapting blood pressure response. *Neuroscience*. 2014;266:197-207.
38. Sabbar M, Dhissi-Benayha O, Benazzouz A, Lakhdar-Ghazal N. Circadian Clock Protein Content and Daily Rhythm of Locomotor Activity Are Altered after Chronic Exposure to Lead in Rat. *Front Behav Neurosci*. 2017;11:178.

39. Zhang T, Huang L, Zhang L, Tan M, Pu M, Pickard GE, et al. ON and OFF retinal ganglion cells differentially regulate serotoninergic and GABAergic activity in the dorsal raphe nucleus. *Sci Rep*. 2016;6:26060.

40. Robillard R, Carpenter JS, Feilds K-L, Hermens DF, White D, Naismith SL, et al. Parallel Changes in Mood and Melatonin Rhythm Following an Adjunctive Multimodal Chronobiological Intervention With Agomelatine in People With Depression: A Proof of Concept Open Label Study. *Front Psychiatry*. 2018;9:624.

41. Duval F, Mokrani M-C, Erb A, Gonzalez opera F, Calleja C, Paris V. Relationship between chronobiological thyrotropin and prolactin responses to protirelin (TRH) and suicidal behavior in depressed patients. *Psychoneuroendocrinology*. 2017;85:100-109.

42. Robillard R, Carpenter JS, Rogers NL, Fares S, Grierson AB, Hermens DF, et al. Circadian rhythms and psychiatric profiles in young adults with unipolar depressive disorders. *Transl Psychiatry*. 2018;8:213.

43. Buttgereit F, Smolen JS, Coogan AN, Cajochen C. Clocking in: chronobiology in rheumatoid arthritis. *Nat Rev Rheumatol*. 2015;11:349-356.

44. Saetung S, Nimitphong H, Siwasaranond N, Manodpitipong A, Crowley SJ, Hood MM, et al. Eveningness Is Associated With Greater Depressive Symptoms in Type 2 Diabetes Patients: A Study in Two Different Ethnic Cohorts. *Behav Sleep Med*. 2019;17:291-301.

45. Charrier A, Olliac B, Roubertoux P, Tordjman S, Charrier A, Olliac B, et al. Clock Genes and Altered Sleep–Wake Rhythms: Their Role in the Development of Psychiatric Disorders. *Int J Mol Sci*. 2017;18:938.

46. Barandas R, Landgraf D, McCarthy MJ, Welsh DK. Circadian Clocks as Modulators of Metabolic Comorbidity in Psychiatric Disorders. *Curr Psychiatry Rep*. 2015;17:98.

47. Guilding C, Piggins HD. Challenging the omnipotence of the suprachiasmatic timekeeper: are circadian oscillators present throughout the mammalian brain? *Ear J Neurosci*. 2007;25:3195-3216.

48. Kalsbeek A, Palm IF, La Fleur SE, Scheer F AJL, Perreau-Lenz S, Ruiter M, et al. SCN Outputs and the Hypothalamic Balance of Life. *J Biol Rhythms*. 2006;21:458-469.

49. Buijs RM, Guzmán Ruiz MA, Méndez Hernández R, Rodríguez Cortés B. The suprachiasmatic nucleus; a responsive clock regulating homeostasis by daily changing the setpoints of physiological parameters. *Auton Neurosci*. 2019;218:43-50.

50. Shiiya T, Nakazato M, Mizuta M, Date Y, Mondal MS, Tanaka M, et al. Plasma Ghrelin Levels in Lean and Obese Humans and the Effect of Glucose on Ghrelin Secretion. *J Clin Endocrinol Metab*. 2002;87:240-244.

51. Shimazu T, Minokoshi Y. Systemic Glucoregulation by Glucose-Sensing Neurons in the Ventromedial Hypothalamic Nucleus (VMH). *J Endocr Soc*. 2017;1:449-459.

52. La Fleur SE, Kalsbeek A, Wortel J, Buijs RM. Polysynaptic neural pathways between the hypothalamus, including the suprachiasmatic nucleus, and the liver. *Brain Res*. 2000;871:50-56.

53. Buijs RM, Guzmán Ruiz MA, Méndez Hernández R, Rodríguez Cortés B. The suprachiasmatic nucleus; a responsive clock regulating homeostasis by daily changing the setpoints of physiological parameters. *Auton Neurosci*. 2019;218:43-50.

54. Shiiya T, Nakazato M, Mizuta M, Date Y, Mondal MS, Tanaka M, et al. Plasma Ghrelin Levels in Lean and Obese Humans and the Effect of Glucose on Ghrelin Secretion. *J Clin Endocrinol Metab*. 2002;87:240-244.

55. Kalsbeek A, Fliers E, Romijn JA, La Fleur SE, Wortel J, Bakker O, et al. The Suprachiasmatic Nucleus Generates the Diurnal Changes in Plasma Leptin Levels. *Endocrinology*. 2001;142:2677-2685.

56. Kalsbeek A, van der Spek R, Lei J, Endert E, Buijs RM, Fliers E. Circadian rhythms in the hypothalamo–pituitary–adrenal (HPA) axis. *Mol Cell Endocrinol*. 2012;349:20-29.
Van De Kar LD, Lorens SA. Differential serotonergic innervation of individual hypothalamic nuclei and other forebrain regions by the dorsal and median midbrain raphe nuclei. *Brain Res.* 1979;162:45-54.

Malek ZS, Sage D, Pévet P, Raison S. Daily Rhythm of Tryptophan Hydroxylase-2 Messenger Ribonucleic Acid within Raphe Neurons Is Induced by Corticoid Daily Surge and Modulated by Enhanced Locomotor Activity. *Endocrinology.* 2007;148:5165-5172.

Buijs FN, León-Mercado L, Guzmán-Ruiz M, Guerrero-Vargas NN, Romo-Nava F, Buijs RM. The Circadian System: A Regulatory Feedback Network of Periphery and Brain. *Physiology.* 2016;31:170-181.

Pontes ALB de, Engelberth RCGJ, Nascimento E da S, Cavalcante JC, Costa MSM de O, Pinato L, et al. Serotonin and circadian rhythms. *Psychol Neurosci.* 2010;3:217-228.

Zaki NFW, Spence DW, BaHammam AS, Pandi-Perumal SR, Cardinali DP, Brown GM. Chronobiological theories of mood disorder. *Eur Arch Psychiatry Clin Neurosci.* 2018;268:107-118.

Pévet P. Melatonin: From Seasonal to Circadian Signal. *J Neuroendocrinol.* 2003;15:422-426.

Reiter RJ. The melatonin rhythm: both a clock and a calendar. *Experientia.* 1993;49:654-664.

Cagnacci A, Elliott JA, Yen SS. Melatonin: a major regulator of the circadian rhythm of core temperature in humans. *J Clin Endocrinol Metab.* 1992;75:447-452.

Chenu F, El Mansari M, Blier P. Electrophysiological Effects of Repeated Administration of Agomelatine on the Dopamine, Norepinephrine, and Serotonin Systems in the Rat Brain. *Neuropsychopharmacology.* 2013;38:275-284.

Kalsbeek A, Cutrerera R., van Heerikhuize J., van der Vliet J, Buijs R. GABA release from suprachiasmatic nucleus terminals is necessary for the light-induced inhibition of nocturnal melatonin release in the rat. *Neuroscience.* 1999;91:453-461.

Khaldy H, León J, Escames G, Bikjdaouene L, García JJ, Acuña-Castroviejo D. Circadian Rhythms of Dopamine and Dihydroxyphenyl Acetic Acid in the Mouse Striatum: Effects of Pinealectomy and of Melatonin Treatment. *Neuroendocrinology.* 2002;75:201-208.

Uz T, Akhisaroglu M, Ahmed R, Manev H. The Pineal Gland is Critical for Circadian Period1 Expression in the Striatum and for Circadian Cocaine Sensitization in Mice. *Neuropsychopharmacology.* 2003;28:2117-2123.

Macchi MM, Bruce JN. Human pineal physiology and functional significance of melatonin. *Front Neuroendocrinol.* 2004;25:177-195.

Burton R. *The Anatomy of Melancholy.* London: Oxford University Press; 1621.

Richter CP. *Biological Clocks in Medicine and Psychiatry.* Springfield, Illinois: Charles C. Thomas.; 1965.

Souètre E, Salvati E, Belougou J-L., Pringuey D, Candito M, Krebs B, et al. Circadian rhythms in depression and recovery: Evidence for blunted amplitude as the main chronobiological abnormality. *Psychiatry Res.* 1989;28:263-278.

Hall P, Spear FG, Stirland D. Diurnal variation of subjective mood in depressive states. *Psychiatr Q.* 1964;38:529-536.

Wehr TA, Wirz-Justice A. Circadian rhythm mechanisms in affective illness and in antidepressant drug action. *Pharmacopsychia.* 1982;15:31-39.

Zung WWK, Green RL. Seasonal Variation of Suicide and Depression. *Arch Gen Psychiatry.* 1974;30:89.

Milstein V, Small JG, Shelbourne D, Small IF. Manic depressive illness: onset, diurnal temperature and season of birth. *Dis Nerv Syst.* 1976;37:373-375.

Eastwood MR, Peacocke J. Seasonal Patterns of Suicide, Depression and Electroconvulsive
Frangos E, Athanassenas G, Tsitourides S, Psilolignos P, Robos A, Katsanou N, et al. Seasonality of the episodes of recurrent affective psychoses: Possible prophylactic interventions. *J Affect Disord.* 1980;2:239-247.

Berkol TD, Seren Kirlioglu S, Balcioglu YH, Ustun N, Islam S, Ozylidirim I. Comparison of sociodemographic and clinical characteristics of bipolar patients with and without seasonal patterns. *Anadolu Psikiyatr Derg.* 2017;18:571-576.

Faedda GL, Tondo L, Teicher MH, Baldessarini RJ, Gelbard HA, Floris GF. Seasonal Mood Disorders: Patterns of Seasonal Recurrence in Mania and Depression. *Arch Gen Psychiatry.* 1993;50:17-23.

Iasevoli F, Avvisati L, Gilardi V, Latte G, Prinzivalli E, de Berardis D, et al. Chronobiology of Mood Disorders. *Melatonin, Neuroprotective Agents Antidepressant Ther.* 2016:273-295.

McClung CA. How Might Circadian Rhythms Control Mood? Let Me Count the Ways... *Biol Psychiatry.* 2013;74:242-249.

Wehr TA, Sack D, Rosenthal N, Duncan W, Gillin JC. Circadian rhythm disturbances in manic-depressive illness. *Fed Proc.* 1983;42:2809-2814.

Wirz-Justice A, Pühringer W, Hole G. Sleep deprivation and clomipramine in endogenous depression. *Lancet.* 1976;308:912.

Wirz-Justice A, Tobler I, Kafka MS, Nahe D, Marangos PJ, Borbély AA, et al. Sleep deprivation: Effects on circadian rhythms of rat brain neurotransmitter receptors. *Psychiatry Res.* 1981;5:67-76.

Simon RD. Shift work disorder: clinical assessment and treatment strategies. *J Clin Psychiatry.* 2012;73:e20.

Vargas I, Garland SN, Kloss JD, Perlis ML. Insomnia and psychiatric disorders. *Sleep Heal.* 2019:373-389.

Vadnie CA, McClung CA. Circadian Rhythm Disturbances in Mood Disorders: Insights into the Role of the Supraocular Nucleus. *Neural Plast.* 2017;2017:1-28.

Stubbs B, Wu Y-T, Prina AM, Leng Y, Cosco TD. A population study of the association between sleep disturbance and suicidal behaviour in people with mental illness. *J Psychiatr Res.* 2016;82:149-154.

Monteleone P, Maj M. The circadian basis of mood disorders: Recent developments and treatment implications. *Eur Neuropsychopharmacol.* 2008;18:701-711.

Kupfer DJ, Foster FG, Coble P, McPartland RJ, Ulrich RF. The application of EEG sleep for the differential diagnosis of affective disorders. *Am J Psychiatry.* 1978;135:69-74.

Kupfer DJ, Foster FG. Interval between onset of sleep and rapid-eye-movement sleep as an indicator of depression. *Lancet.* 1972;300:684-686.

Kupfer DJ, Ulrich RF, Coble PA, Jarrett DB, Grochocinski V, Domann J, et al. Application of automated REM and slow wave sleep analysis: II. Testing the assumptions of the two-process model of sleep regulation in normal and depressed subjects. *Psychiatry Res.* 1984;13:335-343.

Rush AJ, Erman MK, Giles DE, Schlesser MA, Carpenter G, Vasavada N, et al. Polysomnographic Findings in Recently Drug-Free and Clinically Remitted Depressed Patients. *Arch Gen Psychiatry.* 1986;43:878-884.

Giles DE, Jarrett RB, Roffwarg HP, Rush AJ. Reduced rapid eye movement latency. A predictor of recurrence in depression. *Neuropsychopharmacology.* 1987;1:33-39.

Pillai V, Kalmbach DA, Ciesla JA. A meta-analysis of electroencephalographic sleep in depression: Evidence for genetic biomarkers. *Biol Psychiatry.* 2011;70:912-919.

Rotenberg VS, Shamir E, Barak Y, Indursky P, Kayumov L, Mark M. REM sleep latency and wakefulness in the first sleep cycle as markers of major depression: A controlled study vs. schizophrenia and normal controls. *Prog Neuro-Psychopharmacology Biol Psychiatry.* 2002;26:1211-1215.
99. Steiger A, Kimura M. Wake and sleep EEG provide biomarkers in depression. *J Psychiatr Res*. 2010;44:242-252.

100. Monteleone P, Martiadis V, Maj M. Circadian rhythms and treatment implications in depression. *Prog Neuro-Psychopharmacology Biol Psychiatry*. 2011;35:1569-1574.

101. De Berardis D, Orsolini L, Serroni N, Girinelli G, Iasevoli F, Tomasetti C, et al. The role of melatonin in mood disorders. *ChronoPhysiology Ther*. 2015;65-75.

102. Emens J, Lewy A, Kinzie JM, Arntz D, Rough J. Circadian misalignment in major depressive disorder. *Psychiatry Res*. 2009;168:259-261.

103. Shafii M, MacMillan DR, Key MP, Derrick AM, Kaufman N, Nahinsky ID. Nocturnal Serum Melatonin Profile in Major Depression in Children and Adolescents. *Arch Gen Psychiatry*. 1996;53:1009.

104. Hasler BP, Buysse DJ, Kupfer DJ, Germain A. Phase relationships between core body temperature, melatonin, and sleep are associated with depression severity: Further evidence for circadian misalignment in non-seasonal depression. *Psychiatry Res*. 2010;178:205-207.

105. Cagnacci A, Kräuchi K, Wirz-Justice A, Volpe A. Homeostatic versus Circadian Effects of Melatonin on Core Body Temperature in Humans. *J Biol Rhythms*. 1997;12:509-517.

106. Kishi T, Kitajima T, Ikeda M, Yamanouchi Y, Kinoshita Y, Kawashima K, et al. CLOCK may Predict the Response to Fluvoxamine Treatment in Japanese Major Depressive Disorder Patients. *NeuroMolecular Med*. 2009;11:53-57.

107. Wirz-Justice A. Biological rhythm disturbances in mood disorders. *Int Clin Psychopharmacol*. 2006;21:S11-S15.

108. Soria V, Martínez-Amorós Ê, Escaramís G, Valero I, Pérez-Egea R, García C, et al. Differential Association of Circadian Genes with Mood Disorders: CRY1 and NPAS2 are Associated with Unipolar Major Depression and CLOCK and VIP with Bipolar Disorder. *Neuropsychopharmacology*. 2010;35:1279-1289.

109. Lavebratt C, Sjöholm LK, Partonen T, Schalling M, Forsell Y. PER2 variation is associated with depression vulnerability. *Am J Med Genet Part B Neuropsychiatr Genet*. 2010;153:570-581.

110. Kennaway DJ. Clock genes at the heart of depression. *J Psychopharmacol*. 2010;24:5-14.

111. Etain B, Milhiet V, Bellivier F, Leboyer M. Genetics of circadian rhythms and mood spectrum disorders. *Eur Neuropsychopharmacol*. 2011;21:S676-S682.

112. Melhuish Beaupre L, Brown GM, Kennedy JL. Circadian genes in major depressive disorder. *World J Biol Psychiatry*. September 2018;1-11.

113. Herbert J, Goodyer IM, Grossman AB, Hastings MH, de Kloet ER, Lightman SL, et al. Do Corticosteroids Damage the Brain? *J Neuroendocrinol*. 2006;18:393-411.

114. Dumbell R, Matveeva O, Oster H. Circadian clocks, stress, and immunity. *Front Endocrinol (Lausanne)*. 2016;7:37.

115. Carroll BJ, Martin FI, Davies B. Resistance to suppression by dexamethasone of plasma 11-O.H.C.S. levels in severe depressive illness. *Br Med J*. 1968;3:285-287.

116. Keller J, Gomez R, Williams G, Lembke A, Lazzeroni L, Murphy GM, et al. HPA axis in major depression: cortisol, clinical symptomatology and genetic variation predict cognition. *Mol Psychiatry*. 2017;22:527-536.

117. Gold PW. The organization of the stress system and its dysregulation in depressive illness. *Mol Psychiatry*. 2015;20:32-47.

118. Goodyer IM, Herbert J, Altham PME. Adrenal steroid secretion and major depression in 8- to 16-year-olds. III. Influence of cortisol/DHEA ratio at presentation on subsequent rates of disappointing life events and persistent major depression. *Psychol Med*. 1998;28:265-273.

119. Gallagher P, Young A. Cortisol/DHEA ratios in depression [2]. *Neuropsychopharmacology*. 2002;26:410.
120. Markopoulou K, Papadopoulos A, Juruena MF, Poon L, Pariante CM, Cleare AJ. The ratio of cortisol/DHEA in treatment resistant depression. Psychoneuroendocrinology. 2009;34:19-26.
121. Pariante CM, Lightman SL. The HPA axis in major depression: classical theories and new developments. Trends Neurosci. 2008;31:464-468.
122. Carvalho LA, Garner BA, Dew T, Fazakerley H, Pariante CM. Antidepressants, but not antipsychotics, modulate GR function in human whole blood: An insight into molecular mechanisms. Eur Neuropsychopharmacol. 2010;20:379-387.
123. Raison CL, Miller AH. Is Depression an Inflammatory Disorder? Curr Psychiatry Rep. 2011;13:467-475.
124. Pasco JA, Nicholson GC, Williams LJ, Jacka FN, Henry MJ, Kotowicz MA, et al. Association of high-sensitivity C-reactive protein with de novo major depression. Br J Psychiatry. 2010;197:372-377.
125. Postal M, Appenzeller S. The importance of cytokines and autoantibodies in depression. Autoimmun Rev. 2015;14:30-35.
126. Felger JC, Lotrich FE. Inflammatory cytokines in depression: Neurobiological mechanisms and therapeutic implications. Neuroscience. 2013;246:199-229.
127. Narasimamurthy R, Hatori M, Nayak SK, Liu F, Panda S, Verma IM. Circadian clock protein cryptochrome regulates the expression of proinflammatory cytokines. Proc Natl Acad Sci. 2012;109:12662-12667.
128. Imeri L, Opp MR. How (and why) the immune system makes us sleep. Nat Rev Neurosci. 2009;10:199-210.
129. Irwin MR, Olmstead R, Carroll JE. Sleep Disturbance, Sleep Duration, and Inflammation: A Systematic Review and Meta-Analysis of Cohort Studies and Experimental Sleep Deprivation. Biol Psychiatry. 2016;80:40-52.
130. Plante DT, Winkelman JW. Sleep Disturbance in Bipolar Disorder: Therapeutic Implications. Am J Psychiatry. 2008;165:830-843.
131. Harvey AG. Sleep and circadian rhythms in bipolar disorder: Seeking synchrony, harmony, and regulation. Am J Psychiatry. 2008;165:820-829.
132. Dallaspezia S, Benedetti F. Sleep in other psychiatric disorders. In: Oxford Textbook of Sleep Disorders. Oxford: Oxford University Press; 2017:451.
133. Rocha PMB, Neves FS, Corrêa H. Significant sleep disturbances in euthymic bipolar patients. Compr Psychiatry. 2013;54:1003-1008.
134. Geoffroy PA, Scott J, Boudebesse C, Lajnèf M, Henry C, Leboyer M, et al. Sleep in patients with remitted bipolar disorders: a meta-analysis of actigraphy studies. Acta Psychiatr Scand. 2015;131:89-99.
135. Kaplan KA, McGlinchey EL, Soehner A, Gershon A, Talbot LS, Eidelman P, et al. Hypersomnia subtypes, sleep and relapse in bipolar disorder. Psychol Med. 2015;45:1751-1763.
136. Boudebesse C, Geoffroy PA, Bellivier F, Henry C, Folkard S, Leboyer M, et al. Correlations between objective and subjective sleep and circadian markers in remitted patients with bipolar disorder. Chronobiol Int. 2014;31:698-704.
137. Sylvia LG, Chang WC, Kamali M, Tohen M, Kinrys G, Deckersbach T, et al. Sleep disturbance may impact treatment outcome in bipolar disorder: A preliminary investigation in the context of a large comparative effectiveness trial. J Affect Disord. 2018;225:563-568.
138. Harvey AG, Soehner AM, Kaplan KA, Hein K, Lee J, Kanady J, et al. Treating insomnia improves mood state, sleep, and functioning in bipolar disorder: A pilot randomized controlled trial. J Consult Clin Psychol. 2015;83:564-577.
139. Kanady JC, Soehnera AM, Harvey AG. A Retrospective Examination of Sleep Disturbance across the Course of Bipolar Disorder. J sleep Disord Ther. 2015;4.
140. Ng TH, Chung K-F, Ho FY-Y, Yeung W-F, Yung K-P, Lam T-H. Sleep–wake disturbance in
interepisode bipolar disorder and high-risk individuals: A systematic review and meta-analysis. 
*Sleep Med Rev.* 2015;20:46-58.

141. Robillard R, Naismith SL, Rogers NL, Scott EM, Ip TKC, Hermens DF, et al. Sleep-Wake cycle and melatonin rhythms in adolescents and young adults with mood disorders: Comparison of unipolar and bipolar phenotypes. 
*Eur Psychiatry.* 2013;28:412-416.

142. Nováková M, Praško J, Látalová K, Sládek M, Sumová A. The circadian system of patients with bipolar disorder differs in episodes of mania and depression. 
*Bipolar Disord.* 2015;17:303-314.

143. Kennedy SH, Tighe S, McVey G, Brown GM. Melatonin and cortisol “switches” during mania, depression, and euthymia in a drug-free bipolar patient. 
*J Nerv Ment Dis.* 1989;177:300-303.

144. Lewy AJ, Wehr TA, Goodwin FK. Plasma melatonin in manic-depressive illness. 
*Catech Basic Clin Front.* January 1979:1173-1175.

145. Lewy AJ, Wehr TA, Goodwin FK, Newsome DA, Rosenthal NE. Manic-Depressive Patients May Be Supersensitive To Light. 
*Lancet.* 1981;317:383-384.

146. Livianos L, Sierra P, Arques S, García A, Rojo L. Is melatonin an adjunctive stabilizer? 
*Psychiatry Clin Neurosci.* 2012;66:82-83.

147. Geoffroy PA. Clock Genes and Light Signaling Alterations in Bipolar Disorder: When the Biological Clock Is Off. 
*Biol Psychiatry.* 2018;84:775-777.

148. McCarthy MJ, Welsh DK. Cellular Circadian Clocks in Mood Disorders. 
*J Biol Rhythms.* 2012;27:339-352.

149. Kripke DF, Nievergelt CM, Joo EJ, Shekhtman T, Kelsoe JR. Circadian polymorphisms associated with affective disorders. 
*J Circadian Rhythms.* 2009;7:2.

150. McVey G, Brown GM. Melatonin and cortisol “switches” during mania, depression, and euthymia in a drug-free bipolar patient. 
*Catech Basic Clin Front.* January 1979:1173-1175.

151. Palme K, Dmitrzak-Weglarz M, Maciukiewicz M, Wilkosc M, Leszczynska-Rodziewicz A, Zarembo D, et al. Suicidal behavior in the context of disrupted rhythmicity in bipolar disorder—Data from an association study of suicide attempts with clock genes. 
*Psychiatry Res.* 2015;226:517-520.

152. Pawlak J, Dmitrzak-Weglarz M, Maciukiewicz M, Wilkosc M, Leszczynska-Rodziewicz A, Zarembo D, et al. Suicidal behavior in the context of disrupted rhythmicity in bipolar disorder—Data from an association study of suicide attempts with clock genes. 
*Psychiatry Res.* 2015;226:517-520.

153. Belvederi Murri M, Prestia D, Mondelli V, Pariente C, Patti S, Olivieri B, et al. The HPA axis in bipolar disorder: Systematic review and meta-analysis. 
*Psychoneuroendocrinology.* 2016;63:327-342.
160. Girshkin L, Matheson SL, Shepherd AM, Green MJ. Morning cortisol levels in schizophrenia and bipolar disorder: A meta-analysis. Psychoneuroendocrinology. 2014;49:187-206.

161. Modabbernia A, Taslimi S, Brietzke E, Ashrafi M. Cytokine Alterations in Bipolar Disorder: A Meta-Analysis of 30 Studies. Biol Psychiatry. 2013;74:15-25.

162. Brietzke E, Stertz L, Fernandes BS, Kauer-Sant’Anna M, Mascarenhas M, Escosteguy Vargas A, et al. Comparison of cytokine levels in depressed, manic and euthymic patients with bipolar disorder. J Affect Disord. 2009;116:214-217.

163. Bai Y-M, Su T-P, Li C-T, Tsai S-J, Chen M-H, Tu P-C, et al. Comparison of pro-inflammatory cytokines among patients with bipolar disorder and unipolar depression and normal controls. Bipolar Disord. 2015;17:269-277.

164. Peirson SN, Foster RG. Sleep and Circadian Rhythm Disruption in Psychosis. Circadian Med. May 2015:271-282.

165. Wulff K, Dijk D-J, Middleton B, Foster RG, Joyce EM. Sleep and circadian rhythm disruption in schizophrenia. Br J Psychiatry. 2012;200:308-316.

166. Yates NJ. Schizophrenia: The role of sleep and circadian rhythms in regulating dopamine and psychosis. Rev Neurosci. 2016;27:669-687.

167. Fasshauer D, Sutton RB, Brunger AT, Jahn R, Vikman J, Molnár Z, et al. Conserved structural features of the synaptic fusion complex: SNARE proteins reclassified as Q- and R-SNAREs. Proc Natl Acad Sci. 1998;95:15781-15786.

168. Oliver PL, Davies KE. Interaction between environmental and genetic factors modulates schizophrenic endophenotypes in the Snap-25 mouse mutant blind-drunk. Hum Mol Genet. 2009;18:4576-4589.

169. Lamont EW, Coutu DL, Cermakian N, Boivin DB. Circadian rhythms and clock genes in psychotic disorders. Isr J Psychiatry Relat Sci. 2010;47:27-35.

170. Cosgrave J, Wulff K, Gehman P. Sleep, circadian rhythms, and schizophrenia. Curr Opin Psychiatry. 2018;31:176-182.

171. Anderson G, Maes M. Melatonin: an overlooked factor in schizophrenia and in the inhibition of anti-psychotic side effects. Metab Brain Dis. 2012;27:113-119.
180. Olcese JM, Cao C, Mori T, Mamcarz MB, Maxwell A, Runfeldt MJ, et al. Protection against cognitive deficits and markers of neurodegeneration by long-term oral administration of melatonin in a transgenic model of Alzheimer disease. *J Pineal Res.* 2009;47:82-96.

181. Galván-Arrieta T, Trueta C, Cercós MG, Valdés-Tovar M, Alarcón S, Oikawa J, et al. The role of melatonin in the neurodevelopmental etiology of schizophrenia: A study in human olfactory neuronal precursors. *J Pineal Res.* 2017;63:e12421.

182. Rao ML, Gross G, Strebel B, Halaris A, Huber G, Bräunig P, et al. Circadian rhythm of tryptophan, serotonin, melatonin, and pituitary hormones in schizophrenia. *Biol Psychiatry.* 1994;35:151-163.

183. Sandyk R, Kay SR. The Relationship of Pineal Calcification to Cortical Atrophy in Schizophrenia. *Int J Neurosci.* 1991;57:179-191.

184. Witkovsky P. Dopamine and retinal function. *Doc Ophthalmol.* 2004;108:17-39.

185. Tardito D, Milanese M, Bonifacino T, Musazzi L, Grilli M, Mallei A, et al. Blockade of stress-induced increase of glutamate release in the rat prefrontal/frontal cortex by agomelatine involves synergy between melatonergic and 5-HT2Creceptor-dependent pathways. *BMC Neurosci.* 2010;11:1-4.

186. Lewis DA, Moghaddam B. Cognitive dysfunction in schizophrenia: Convergence of γ-aminobutyric acid and glutamate alterations. *Arch Neurol.* 2006;63:1372-1376.

187. Gorfine T, Assaf Y, Goshen-Gottstein Y, Yeshurun Y, Zisapel N. Sleep-anticipating effects of melatonin in the human brain. *Neuroimage.* 2006;31:410-418.

188. Shamir E, Barak Y, Shalman I, Laudon M, Zisapel N, Tarrasch R, et al. Melatonin Treatment for Tardive Dyskinesia. *Arch Gen Psychiatry.* 2001;58:1049.

189. Onaolapo AY, Aina OA, Onaolapo OJ. Melatonin attenuates behavioural deficits and reduces brain oxidative stress in a rodent model of schizophrenia. *Biomed Pharmacother.* 2017;92:373-383.

190. Zhang J, Liao G, Liu C, Sun L, Liu Y, Wang Y, et al. The association of CLOCK gene T3111C polymorphism and hPER3 gene 54-nucleotide repeat polymorphism with Chinese Han people schizophrenics. *Mol Biol Rep.* 2011;38:349-354.

191. Aston C, Jiang L, Sokolov BP. Microarray analysis of postmortem temporal cortex from patients with schizophrenia. *J Neurosci Res.* 2004;77:858-866.

192. Sun H-Q, Li S-X, Chen F-B, Zhang Y, Li P, Jin M, et al. Diurnal neurobiological alterations after exposure to clozapine in first-episode schizophrenia patients. *Psychoneuroendocrinology.* 2016;64:108-116.

193. Johansson A-S, Owe-Larsson B, Hetta J, Lundkvist GB. Altered circadian clock gene expression in patients with schizophrenia. *Schizopr Res.* 2016;174:17-23.

194. Pinacho R, Villalmanzo N, Meana JJ, Ferrer I, Berengueras A, Haro JM, et al. Altered CSNK1E, FABP4 and NEFH protein levels in the dorsolateral prefrontal cortex in schizophrenia. *Schizopr Res.* 2016;177:88-97.

195. Zubiñ J, Spring B. Vulnerability: A new view of schizophrenia. *J Abnorm Psychol.* 1977;86:103-126.

196. Coulon N, Brailly-Tabard S, Walter M, Tordjman S. Altered circadian patterns of salivary cortisol in individuals with schizophrenia: A critical literature review. *J Physiol.* 2016;110:439-447.

197. Singh M, Solanki RK, Bagaria B, Swami MK. Hypothalamic-Pituitary-Adrenal (HPA) axis functioning among patients with schizophrenia: A cross sectional comparative study. *African J Psychiatry (South Africa).* 2015;18:2.

198. Mittal VA, Walker EF. Minor physical anomalies and vulnerability in prodromal youth. *Schizopr Res.* 2011;129:116-121.

199. Carol EE, Mittal VA. Resting cortisol level, self-concept, and putative familial environment in adolescents at ultra high-risk for psychotic disorders. *Psychoneuroendocrinology.* 2015;57:26-
200. Bradley AJ, Dinan TG. Review: A systematic review of hypothalamic-pituitary-adrenal axis function in schizophrenia: implications for mortality. *J Psychopharmacol*. 2010;24:91-118.

201. Zorn J V., Schür RR, Boks MP, Kahn RS, Joëls M, Vinkers CH. Cortisol stress reactivity across psychiatric disorders: A systematic review and meta-analysis. *Psychoneuroendocrinology*. 2017;77:25-36.

202. Remes O, Brayne C, van der Linde R, Lafortune L. A systematic review of reviews on the prevalence of anxiety disorders in adult populations. *Brain Behav*. 2016;6:e00497.

203. Boland EM, Ross RJ. Recent Advances in the Study of Sleep in the Anxiety Disorders, Obsessive-Compulsive Disorder, and Posttraumatic Stress Disorder. *Psychiatr Clin North Am*. 2015;38:761-776.

204. Cox RC, Olatunji BO. A systematic review of sleep disturbance in anxiety and related disorders. *J Anxiety Disord*. 2016;37:104-129.

205. Mesa F, Beidel DC, Bunnell BE. An examination of psychopathology and daily impairment in adolescents with social anxiety disorder. *PLoS One*. 2014;9.

206. Brown TM, Black B, Uhde TW. The sleep architecture of social phobia. *Biol Psychiatry*. 1994;35:420-421.

207. De Bundel D, Gangarossa G, Biever A, Bonnefont X, Valjent E. Cognitive dysfunction, elevated anxiety, and reduced cocaine response in circadian clock-deficient cryptochrome knockout mice. *Front Behav Neurosci*. 2013;7:152.

208. Akiyama M, Kiri hara T, Takahashi S, Minami Y, Yoshinobu Y, Moriya T, et al. Modulation of mPer1 gene expression by anxiolytic drugs in mouse cerebellum. *Br J Pharmacol*. 1999;128:1616-1622.

209. Griesauer I, Diao W, Ronovsky M, Elbouai I, Sartori S, Singewald N, et al. Circadian abnormalities in a mouse model of high trait anxiety and depression. *Ann Med*. 2014;46:148-154.

210. Sipilä T, Kananen L, Greco D, Donner J, Silander K, Terwilliger JD, et al. An Association Analysis of Circadian Genes in Anxiety Disorders. *Biol Psychiatry*. 2010;67:1163-1170.

211. *American Psychiatric Association. Diagnostic and Statistical Manual of Mental Disorders: DSM-5*. 5th ed. Washington, D.C.: American Psychiatric Pub; 2013.

212. Cox RC, Olatunji BO. Circadian Rhythms in Obsessive-Compulsive Disorder: Recent Findings and Recommendations for Future Research. *Curr Psychiatry Rep*. 2019;21:54.

213. Shi S, White MJ, Borsetti HM, Pendergast JS, Hida A, Ciarleglio CM, et al. Molecular analyses of circadian gene variants reveal sex-dependent links between depression and clocks. *Transl Psychiatry*. 2016;6:e748-e748.

214. Shi J, Wittke-Thompson JK, Badner JA, Hattori E, Potash JB, Willour VL, et al. Clock genes may influence bipolar disorder susceptibility and dysfunctional circadian rhythm. *Am J Med Genet Part B Neuropsychiatr Genet*. 2008;147B:1047-1055.

215. Suzuki M, Dallaseppa S, Locatelli C, Lorenzi C, Uchiyama M, Colombo C, et al. CLOCK gene variants associated with the discrepancy between subjective and objective severity in bipolar depression. *J Affect Disord*. 2017;210:14-18.

216. Benedetti F, Riccaboni R, Dallaseppa S, Locatelli C, Smeraldi E, Colombo C. Effects of CLOCK gene variants and early stress on hopelessness and suicide in bipolar depression. *Chronobiol Int*. 2015;32:1156-1161.

217. Utge SJ, Soronen P, Loukola A, Kronholm E, Ollila HM, Pirkola S, et al. Systematic Analysis of Circadian Genes in a Population-Based Sample Reveals Association of TIMELESS with Depression and Sleep Disturbance. *Reitsma PH, ed. PLoS One*. 2010;5:e9259.

218. Dmitrak-Weglarz MP, Pawlak JM, Maciukiewicz M, Moczko J, Wilkos M, Leszczynska-Rozdziewicz A, et al. Clock gene variants differentiate mood disorders. *Mol Biol Rep*. 2015;42:277-288.
219. Mansour HA, Wood J, Logue T, Chowdari K V., Dayal M, Kupfer DJ, et al. Association study of eight circadian genes with bipolar I disorder, schizoaffective disorder and schizophrenia. *Genes, Brain Behav.* 2006;5:150-157.

220. Etain B, Jamain S, Milhiet V, Lajnef M, Boudebesse C, Dumaine A, et al. Association between circadian genes, bipolar disorders and chronotypes. *Chronobiol Int.* 2014;31:807-814.

221. Hua P, Liu W, Chen D, Zhao Y, Chen L, Zhang N, et al. Cry1 and Tef gene polymorphisms are associated with major depressive disorder in the Chinese population. *J Affect Disord.* 2014;157:100-103.

222. Partonen T, Treutlein J, Alpman A, Frank J, Johansson C, Depner M, et al. Three circadian clock genes Per2, Arntl, and Npas2 contribute to winter depression. *Annu Med.* 2007;39:229-238.

223. Liu JJ, Sudic Hukic D, Forsell Y, Schalling M, Öxby U, Lavebratt C. Depression-associated ARNTL and PER2 genetic variants in psychotic disorders. *Chronobiol Int.* 2015;32:579-584.

224. Artioli P, Lorenzi C, Pirovano A, Serretti A, Benedetti F, Catalano M, et al. How do genes exert their role? Period 3 gene variants and possible influences on mood disorder phenotypes. *Eur Neuropsychopharmacol.* 2007;17:587-594.

225. Maglione JE, Nievergelt CM, Parimi N, Evans DS, Ancoli-Israel S, Stone KL, et al. Associations of PER3 and RORA Circadian Gene Polymorphisms and Depressive and Insomnia Phenotypes in Older Adults. *Am J Geriatr Psychiatry.* 2015;23:1075-1087.

226. Brasil Rocha PM, Campos SB, Neves FS, da Silva Filho HC. Genetic Association of the PERIOD3 (Per3) Clock Gene with Bipolar Disorder. *Psychiatry Investig.* 2017;14:674-680.

227. Nievergelt CM, Kripke DF, Barrett TB, Burg E, Remick RA, Sadovnick AD, et al. Suggestive evidence for association of the circadian genes PERIOD3 and ARNTL with bipolar disorder. *Am J Med Genet Part B Neuropsychiatr Genet.* 2006;141B:234-241.

228. Benedetti F, Dallapazza S, Colombo C, Pirovano A, Marino E, Smeraldi E. A length polymorphism in the circadian clock gene Per3 influences age at onset of bipolar disorder. *Neurosci Lett.* 2008;445:184-187.

229. Severino G, Manchia M, Contu P, Squassina A, Lampus S, Ardau R, et al. Association study in a Sardinian sample between bipolar disorder and the nuclear receptor REV-ERBα gene, a critical component of the circadian clock system. *Bipolar Disord.* 2009;11:215-220.

230. Kishi T, Kitajima T, Ikeda M, Yamanouchi Y, Kinoshita Y, Kawashima K, et al. Association analysis of nuclear receptor Rev-erb alpha gene (NR1D1) with mood disorders in the Japanese population. *Neurosci Res.* 2008;62:211-215.

231. Geoffroy PA, Etain B, Lajnef M, Zerdazi E-H, Brichant-Petitjean C, Heilbronner U, et al. Circadian genes and lithium response in bipolar disorders: associations with PPARGC1A (PGC-1 α ) and RORA. *Genes, Brain Behav.* 2016;15:660-668.

232. Lai Y-C, Kao C-F, Lu M-L, Chen H-C, Chen P-Y, Chen C-H, et al. Investigation of Associations between NR1D1, RORA and RORB Genes and Bipolar Disorder. Maher B, ed. *PLoS One.* 2015;10:e0121245.
237. Gałecki P, Szemraj J, Bartosz G, Bienkiewicz M, Galecka E, Florkowski A, et al. Single-nucleotide polymorphisms and mRNA expression for melatonin synthesis rate-limiting enzyme in recurrent depressive disorder. *J Pineal Res.* 2010;48:311-317.

238. Talarowska M, Szemraj J, Zajączkowska M, Gałecki P. ASMT gene expression correlates with cognitive impairment in patients with recurrent depressive disorder. *Med Sci Monit.* 2014;20:905-912.

239. Etain B, Dumaine A, Bellivier F, Pagan C, Francelle L, Goubran-Botros H, et al. Genetic and functional abnormalities of the melatonin biosynthesis pathway in patients with bipolar disorder. *Hum Mol Genet.* 2012;21:4030-4037.

240. Gałecka E, Szemraj J, Florkowski A, Galecki P, Bieńkiewicz M, Karbownik-Lewińska M, et al. Single nucleotide polymorphisms and mRNA expression for melatonin MT2 receptor in depression. *Psychiatry Res.* 2011;189:472-474.

241. Lee KY, Ahn YM, Kim SH, Kang H-G, Joo E-J. Genetic association study of CSNK1E gene in bipolar disorder and circadian characteristics. *Nord J Psychiatry.* 2018;72:599-604.

242. Matsunaga S, Ikeda M, Kishi T, Fukuo Y, Aleksic B, Yoshimura R, et al. An evaluation of polymorphisms in casein kinase 1 delta and epsilon genes in major psychiatric disorders. *Neurosci Lett.* 2012;529:66-69.

243. Kaladchibachi SA, Doble B, Anthopoulos N, Woodgett JR, Manoukian AS. Glycogen synthase kinase 3, circadian rhythms, and bipolar disorder: A molecular link in the therapeutic action of lithium. *J Circadian Rhythms.* 2007;5:3.

244. Szczepankiewicz A, Skibinska M, Hauser J, Slopien A, Leszczynska-Rodziewicz A, Kapelski P, et al. Association Analysis of the &lt;i&gt;GSK-3&lt;/i&gt; T–50C Gene Polymorphism with Schizophrenia and Bipolar Disorder. *Neuropsychobiology.* 2006;53:51-56.

245. Kupfer D, Foster FG. INTERVAL BETWEEN ONSET OF SLEEP AND RAPID-EYE-MOVEMENT SLEEP AS AN INDICATOR OF DEPRESSION. *Lancet.* 1972;300:684-686.

246. Kupfer DJ. REM latency: a psychobiologic marker for primary depressive disease. *Biol Psychiatry.* 1976;11:159-174.

247. Sitaram N, Nurnberger JI, Gershon ES, Gillin JC. Cholinergic regulation of mood and REM sleep: Potential model and marker of vulnerability to affective disorder. *Am J Psychiatry.* 1982;139:571-576.

248. Millar A, Espie CA, Scott J. The sleep of remitted bipolar outpatients: A controlled naturalistic study using actigraphy. *J Affect Disord.* 2004;80:145-153.

249. Hudson JJ, Lipinski JF, Keck PE, Aizley HG, Lukas SE, Rothschild AJ, et al. Polysomnographic Characteristics of Young Manic Patients: Comparison with Unipolar Depressed Patients and Normal Control Subjects. *Arch Gen Psychiatry.* 1992;49:378-383.

250. Linkowski P, Mendlewicz J. Sleep electroencephalogram and rhythm disturbances in mood disorders, *Curr Opin Psychiatry.* 1993;6:35-37.

251. Hudson JJ, Lipinski JF, Frankenburg FR, Grochocinski VJ, Kupfer DJ. Electroencephalographic Sleep in Mania. *Arch Gen Psychiatry.* 1988;45:267-273.

252. Gillin JC, Duncan W, Pettigrew KD, Frankel BL, Snyder F. Successful Separation of Depressed, Normal, and Insomniac Subjects by EEG Sleep Data. *Arch Gen Psychiatry.* 1979;36:85-90.

253. Lauer CJ, Wiegand M, Krieg JC. All-night electroencephalographic sleep and cranial computed tomography in depression - A study of unipolar and bipolar patients. *Eur Arch Psychiatry Clin Neurosci.* 1992;242:59-68.

254. Giles DE, Rush AJ, Roffwarg HP. Sleep parameters in bipolar I, bipolar II, and unipolar depressions. *Biol Psychiatry.* 1986;21:1340-1343.

255. Fossion P, Staner L, Drameix M, Kempenaers C, Kerkhofs M, Hubain P, et al. Does sleep EEG data distinguish between UP, BPI or BPII major depressions? An age and gender controlled study. *J Affect Disord.* 1998;49:181-187.
256. Jernajczyk W. Latency of eye movement and other REM sleep parameters in bipolar depression. *Biol Psychiatry*. 1986;21:465-472.

257. Bian Y, Liang W-Y, Yue W-H, Han X-L, Lin C, Zhang J, et al. Sleep architecture in drug naïve patients with schizophrenia: A meta-analysis. *Chinese Ment Heal J*. 2017;31:208-214.

258. Nelson JC, Davis JM. DST Studies in Psychotic Depression: A Meta-Analysis. *Am J Psychiatry*. 1997;154:1497-1503.

259. Keller J, Flores B, Gomez RG, Solvason HB, Kenna H, Williams GH, et al. Cortisol Circadian Rhythm Alterations in Psychotic Major Depression. *Biol Psychiatry*. 2006;60:275-281.

260. Belanoff JK, Kaltezan M, Sund B, Fleming Ficek SK, Schatzberg AF. Cortisol activity and cognitive changes in psychotic major depression. *Am J Psychiatry*. 2001;158:1612-1616.

261. Gold PW. The organization of the stress system and its dysregulation in depressive illness. *Mol Psychiatry*. 2014;20:32-47.

262. Pelham RW, Vaughan GM, Dandoek KL, Vaughan MK. New York: United Nations, 1977. 2. International Commission on Radiological Protection, ICRP publication 26. *J Biometeorol*. 1975;19.

263. Paparrigopoulos T. Melatonin response to atenolol administration in depression: Indication of β-adrenoceptor dysfunction in a subtype of depression. *Acta Psychiatr Scand*. 2002;106:440-445.

264. Wehr TA, Sack DA, Duncan WC, Mendelson WB, Rosenthal NE, Gillin JC, et al. Sleep and circadian rhythms in affective patients isolated from external time cues. *Psychiatry Res*. 1985;15:327-339.

265. Buckley TM, Schatzberg AF. A pilot study of the phase angle between cortisol and melatonin in major depression - A potential biomarker? *J Psychiatr Res*. 2010;44:69-74.

266. Tuunainen A, Kripke DF, Elliott JA, Assmus JD, Rex KM, Klauber MR, et al. Depression and endogenous melatonin in postmenopausal women. *J Affect Disord*. 2002;69:149-158.

267. Claustrat B, Chazot G, Brun J, Jordan D, Sassolas G. Altered sleep architecture and higher incidence of subsyndromal depression in low endogenous melatonin secretors. *Psychoneuroendocrinology*. 2010;25:S102-S108.

268. Beck‐Friis J, von Rosen D, Kjellman BF, Ljunggren J‐G, et al. Serum melatonin and urinary 6-sulfatoxymelatonin in major depression. *Psychoneuroendocrinology*. 1984;9:291-12.

269. Wetterberg L. Clinical Importance of Melatonin. *Prog Brain Res*. 1979;52:539-547.

270. Parry BL, Newton RP. Chronobiological basis of female-specific mood disorders. *Neuropsychopharmacology*. 2001;25:S102-S108.

271. Beck‐Friis J, Kjellman BF, Ljunggren J, Wetterberg L. Melatonin in relation to body measures, sex, age, season and the use of drugs in patients with major affective disorders and healthy subjects. *Psychoneuroendocrinology*. 1984;9:261-277.

272. Wetterberg L. Melatonin. *Prog Brain Res*. 1979;52:539-547.

273. Claustrat B, Chazot G, Brun J, Jordan D, Sassolas G. A chronobiological study of melatonin and cortisol secretion in depressed subjects: plasma melatonin, a biochemical marker in major depression. *Biol Psychiatry*. 1984;19:1215-1228.

274. Parry BL, Newton RP. Chronobiological basis of female-specific mood disorders. *Neuropsychopharmacology*. 2001;25:S102-S108.

275. Beck‐Friis J, Kjellman BF, Aperia B, Uden F, von Rosen D, Ljunggren J -G, et al. Serum melatonin in relation to clinical variables in patients with major depressive disorder and a hypothesis of a low melatonin syndrome. *Acta Psychiatr Scand*. 1985;71:319-330.

276. Nair NPV, Hariharasubramanian N, Pilapil C. Circadian rhythm of plasma melatonin in endogenous depression. *Prog Neuropsychopharmacol Biol Psychiatry*. 1984;8:715-718.

277. Brown R, Kocsis JH, Caroff S, Amsterdam J, Winokur A, Stokes PE, et al. Differences in nocturnal melatonin secretion between melancholic depressed patients and control subjects. *Am
277. Frazer A, Brown R, Kocsis J, Caroff S, Amsterdam J, Winokur A, et al. Patterns of melatonin rhythms in depression. *J Neural Transm Suppl*. 1986;21:269-290.

278. Fountoulakis KN, Karamouzis M, Iacovides A, Nimatoudis J, Diakogiannis J, Kaprinis G, et al. Morning and evening plasma melatonin and dexamethasone suppression test in patients with nonseasonal major depressive disorder from northern Greece (latitude 40-41.5°). *Neuropsychobiology*. 2001;44:113-117.

279. Kennedy SH, Kutcher SP, Ralevski E, Brown GM. Nocturnal melatonin and 24-hour 6-sulphatoxymelatonin levels in various phases of bipolar affective disorder. *Psychiatry Res*. 1996;63:219-222.

280. Monteleone P, Natale M, La Rocca A, Maj M. Decreased nocturnal secretion of melatonin in drug-free schizophrenics: No change after subchronic treatment with antipsychotics. *Neuropsychobiology*. 1997;36:159-163.

281. Monteleone P, Maj M, Fusco M, Kemali D, Reiter RJ. Depressed nocturnal plasma melatonin levels in drug-free paranoid schizophrenics. *Schizophr Res*. 1992;7:77-84.

282. Bersani G, Mameli M, Garavini A, Pancheri P, Nordio M. Reduction of night/day difference in melatonin blood levels as a possible disease-related index in schizophrenia. *Neuroendocrinol Lett*. 2003;24:181-184.
**Figure 1.** Flowchart of articles selected for the review.
| Gene      | Nomenclature and Protein                      | Protein function                                                                 | Associated disorder                                      |
|-----------|-----------------------------------------------|----------------------------------------------------------------------------------|---------------------------------------------------------|
| Clock     | Circadian Locomotor Output Cycles Kaput (CLOCK) | Positive regulation of \textit{period} and \textit{timeless} genes through interaction with BMAL-1 | BDD16,108,149,214–216,193,197,199,214,216,217,190,192 |
| Timeless  | Timeless homolog (TIM)                        | Negative regulation of CLOCK-BMAL-1 activity through interaction with PER and close the circadian feedback loop | BD1217,214,217,219,220                                  |
| Cry-1     | Cryptochrome-1 (CRY-1)                        | Inhibition of CLOCK-BMAL-1                                                         | BD108,208,221                                            |
| Cry-2     | Cryptochrome-2 (CRY-2)                        | Inhibition of CLOCK-BMAL-1                                                         | BD108,221                                               |
| Per-1     | Period homolog 1 (PER-1)                      | Negative regulation of CLOCK-BMAL-1 activity through interaction with CRY and close the circadian feedback loop | BD149,192                                               |
| Per-2     | Period homolog 2 (PER-2)                      | Negative regulation of CLOCK-BMAL-1 activity through interaction with CRY and close the circadian feedback loop | BD108,109,221                                            |
| Per-3     | Period homolog 3 (PER-3)                      | Seems not to have a critical role circadian rhythm. Contribute to determination of diurnal preference | BD108,219,224,225,234,230,192                           |
| Bmal-1 (or ARNTL-1) | Brain muscle ARNT like protein-1 (Aryl Hydrocarbon Receptor Nuclear Translocator like-1) (BMAL-1/ARNTL-1) | Positive regulation of \textit{period} and \textit{timeless} genes through interaction with CLOCK | BD108,217,222                                            |
| Bmal-2    | Brain muscle ARNT like protein-2              | Probably has a role in activation of CLOCK and CLOCK-controlled genes              | ANX120                                                  |
| Npas-2    | Neuronal PAS domain protein-2 (NPAS-2)        | Intrinsic enhancer for pre-mRNA splicing                                          | BD108,213,222,108,149,192                              |
| Nr1d-1 (or Rev-erb-α) | Nuclear receptor subfamily-1, group d, member 1 (or orphan nuclear receptor REV-ERB-α) (NR1D1/REV-ERB-α) | Works as nuclear hormone receptors. Compete with RORA for binding to the BMAL-1 promoter and repress the BMAL-1 | BD108,217,222,108,149,192,223,224                        |
| Rora      | Retinoid-related orphan receptor a (RORA)     | Works as nuclear hormone receptors. Compete with NR1D1 for binding to the BMAL-1 promoter and activate the BMAL-1 | BD108,217,225                                            |
| Rorb      | Retinoid-related orphan receptor b (RORB)     | Works as nuclear hormone receptors. Compete with NR1D1 for binding to the BMAL-1 promoter and activate the BMAL-1 | BD150,236                                                |
| Dbp       | D site of albumin promoter binding protein    | Being regulated by CLOCK-BMAL-1 and CRY-1. Supports the rhythmic transcription of downstream genes | BD108,214                                                |
| Asmt      | Acetylserotonin methyltransferase             | The last enzyme of the melatonin synthesis pathway                                | BD108,214,219                                           |
| Mtnr-1B   | Melatonin receptor 1b                         | G protein coupled melatonin reseptor                                             | BD140                                                   |
| Aanat     | Arylalkylamine N-acetyltransferase            | The first enzyme of the melatonin synthesis pathway                              | BD108                                                   |
| Csnk-1ε   | Casein kinase 1 epsilon (CSNK1ε)              | Phosphorylates of PER, CRY and BMAL, increases their degradation                  | BD108,217,BD124,242,242,194,242                        |
| Csnk-1δ   | Casein kinase 1 delta (CSNK1δ)                | Phosphorylates of PER, CRY and BMAL, increases their degradation Regulation circadian period length | BD149,242,192                                           |
| GSK-3β    | Glycogen synthase kinase-3β (GSK-3β)          | Regulation circadian period length                                                | BD143,244                                                |

*CLOCK T3111C polymorphism*
Table 2. Main alterations of sleep architecture in psychiatric disorders

| Disorder | Major alterations |
|----------|-------------------|
| MDD      | Shortened latency of the initial REM sleep, prolonged first REM period, increased total REM time, increased REM density and proportion of REM sleep, decreased non-REM sleep | 95–97,245,246 |
| BD       | **Euthymia**: Increased REM density and proportion of REM sleep, longer sleep onset latency and sleep duration, lower sleep efficiency | 133,134,247,248 |
|          | **Mania**: Shortened REM sleep latency, increased REM activity and REM density, reduced total sleep time | 249–251 |
|          | **Depression**: More fragmented REM sleep periods, shortened REM sleep latency | 252,253 |
|          | Longer sleep latency, increased proportion of REM sleep, trend toward higher percentage of awakenings in bipolar depression than in unipolar depression | 254–256 |
| SCH      | **Comparison to healthy control**: Reduced total sleep time, longer sleep onset latency, lower sleep efficiency and REM latency, increased REM density, decreased total REM time, decreased non-REM stage-3 and stage-4 | 175 |
|          | **Medication naive patients**: reduced total sleep time, lower sleep efficiency, increased REM latency, decreased stage-4 of non-REM sleep, increased stage-1 of non-REM | 257 |
|          | Duration of illness has not an effect on polysomnography parameters | 175 |
| ANX      | **Generalized anxiety disorder**: reduced total sleep time, longer sleep onset latency, alterations in non-REM sleep architecture, inconsistent findings for REM sleep architecture and sleep efficiency | 204 |
|          | **Panic disorder**: decreased sleep efficiency and total sleep time, longer sleep onset latency, REM and non-REM sleep architecture findings are less clear | 204 |
|          | **Post-traumatic stress disorder**: reduced total sleep time, longer sleep onset latency, variations in REM sleep | |
| OCD      | Reduced total sleep time, increased wake after sleep onset, inconsistent findings for REM and non-REM sleep architectures | 204 |
Table 3. Summary of consistent findings on the alterations of two major neurohumoral systems regulating circadian rhythm in psychiatric disorders

| Diagnosis | HPA Axis | Neurohumoral System |
|-----------|----------|---------------------|
| MDD       | Elevated baseline cortisol levels, disruption on the dexamethasone suppression test results \(^{115,116,258-261}\) increased cortisol/ DHEA ratio \(^{118-120}\) | Lower nocturnal melatonin levels, delayed melatonin secretion onset and offset \(^{102,262-278}\) |
| BD        | Increased cortisol and ACTH levels in manic phase Findings about HPA axis abnormalities are seen both depressive and euthymic phase, it is preferred to evaluate them as state and trait marker due to clinical variations \(^{158}\) | Higher melatonin levels in manic phase at the daytime \(^{142}\) Findings about nocturnal melatonin levels among BD phases are inconsistent \(^{72,144,145,279}\) |
| SCH       | Baseline cortisol levels are inconsistent Blunted cortisol stress response \(^{201}\) | Lower nocturnal melatonin levels \(^{280,281}\), phase advance in melatonin rhythm \(^{182}\), the absence of melatonin rhythmicity \(^{282}\) |