An estimated one-third of the general population is affected by insomnia, and this number is increasing due to more stressful working conditions and the progressive aging of society. However, current treatment of insomnia with hypnotics, gamma-aminobutyric acid A (GABA_A) receptor modulators, induces various side effects, including cognitive impairment, motor disturbance, dependence, tolerance, hangover, and rebound insomnia. Ramelteon (Rozerem; Takeda Pharmaceutical Company Limited, Osaka, Japan) is an orally active, highly selective melatonin MT_1/MT_2 receptor agonist. Unlike the sedative hypnotics that target GABA_A receptor complexes, ramelteon is a chronohypnotic that acts on the melatonin MT_1 and MT_2 receptors, which are primarily located in the suprachiasmatic nucleus, the body’s “master clock.” As such, ramelteon possesses the first new therapeutic mechanism of action for a prescription insomnia medication in over three decades. Ramelteon has demonstrated sleep-promoting effects in clinical trials, and coupled with its favorable safety profile and lack of abuse potential or dependence, this chronohypnotic provides an important treatment option for insomnia.

Introduction

Most adults have experienced insomnia or sleeplessness at one time or another in their lives. An estimated one-third of the general population is affected by insomnia, and 10–15% have chronic insomnia [1]. Although most of us know what insomnia is and how we feel and perform after one or more sleepless nights, few seek medical advice. Many people remain unaware of the behavioral and medical options available to treat insomnia. Insomnia affects all age groups. The incidence increases with age, and among older adults, insomnia affects women more often than men [2].

The first-generation drugs for insomnia were barbiturates such as pentobarbital and phenobarbital, which were often used as sedative hypnotics/anxiolytics before benzodiazepines largely came to be used for these purposes. However, barbiturates have a high abuse potential, and overdose can cause unconsciousness and even death due to respiratory suppression [3–5].

Benzodiazepines, the second generation of hypnotics, such as triazolam, lorazepam, and estazolam, have been used for treatment of insomnia, because benzodiazepines have a low potential for abuse and low danger of lethal overdose [3,4,6]. However, these hypnotics produce several side effects including cognitive impairment, psychomotor impairment, dependence, tolerance, hangover, rebound insomnia, and so on [7–10]. Then benzodiazepine receptor agonists, the third generation of hypnotics with non-benzodiazepine chemical structures, such as zolpidem, zopiclone, and zaleplon, were developed to maintain sleep-inducing action and reduce side effects such as amnesia and motor dysfunction [11–14]. It has been reported that there is no tolerance during treatment and no or limited rebound insomnia after therapy discontinuation in the long-term use of zaleplon, eszopiclone, and modified release formulation of zolpidem [11,15,16]. However, at higher doses, non-benzodiazepine hypnotics also cause similar side effects as those caused by benzodiazepines, although the severity of side effects differs among specific drugs [11–13,17,18–20]. Benzodiazepine receptor agonists, including non-benzodiazepines, have abuse and dependence potential [18,21–23]. Furthermore, sleep produced by these
agents is electrophysiologically different from that of naturally occurring physiological sleep. Benzodiazepine receptor agonists reduce rapid eye movement (REM) sleep and increase stage 2 sleep [16,24–27]. Newer non-benzodiazepine hypnotics, including zopiclone, zaleplon, and zolpidem, also have been shown to decrease REM sleep in the first half of the night, although the effects were milder than those of benzodiazepines such as triazolam and temazepam [25,28,29]. Also, electroencephalographic (EEG) power spectral analyses revealed that there is a decrease in low-frequency (0.25–10.0 Hz) activity and increase in high-frequency activity in non-REM sleep [24,25,30].

Since its discovery in 1958 [31], melatonin has been shown to play a vital role in the regulation of circadian rhythms, including the sleep–wake cycle [32–34]. The cyclic nature of melatonin’s production by the pineal gland is controlled by neuronal output from the suprachiasmatic nucleus (SCN) of the hypothalamus. With regard to sleep in humans, melatonin production is concurrent with nocturnal sleep; the increase in endogenous melatonin levels in the evening correlates with the onset of self-reported evening sleepiness [35,36] and an increase in sleep propensity [37].

The identification and cloning of melatonin receptors has increased our understanding of melatonin’s role in sleep. In 1998, the nomenclature committee of the International Union of Basic and Clinical Pharmacology classified melatonin receptors as MT₁, MT₂, and MT₃. The MT₁ receptor (formerly ML₁A or Mel₁A) and MT₂ receptor (formerly ML₁B or Mel₁B) were initially defined as high-affinity binding sites (picomolar affinity) in chick and mammalian brains and retina [38–42]. MT₁ receptor mRNA has been detected in the SCN, and studies using an MT₁ receptor knockout mouse indicate that this receptor mediates the acute inhibition of SCN firing by melatonin [33]. MT₂ receptor mRNA has also been detected in the SCN, and the activity at this receptor has been associated with the phase-shifting effects of melatonin on circadian rhythms [32,43–45]. The MT₃ binding site (formerly ML₂) was initially defined as a low-affinity binding site (nanomolar affinity) in mammalian brains and peripheral organs [46], and it has recently been characterized as a melatonin-sensitive form of quinone reductase 2 [47,48]. MT₃ has a profile that is completely different from that of the MT₁ and MT₂ receptors, and it is not likely to be involved in the sleep–wake cycle [49].

The notion that direct administration of exogenous melatonin could promote sleep has been dampened by inconsistent efficacy results in clinical trials [50–53]. According to the 2005 National Institutes of Health (NIH) State-of-the-Science Conference Statement on Manifestations and Management of Chronic Insomnia in Adults, melatonin’s effect on sleep promotion is questionable based on information from meta-analyses and consensus statements. Because melatonin is not regulated by the FDA, preparations vary in their melatonin content, making comparisons across studies difficult. Although melatonin appears to be effective for the treatment of circadian rhythm disorders [54–56], little consistent evidence exists for its efficacy in the treatment of insomnia. The limited efficacy of melatonin for insomnia may be partly attributed to its short half-life [57,58]. A controlled-release melatonin has been shown to improve initiation of sleep and increase sleep efficiency and total sleep time in clinical trials in elderly people with insomnia [59,60]. These finding suggests that a high-affinity MT₁/MT₂ receptor agonist with a longer half-life than that of melatonin might be a useful therapy for sleep disorders in this population. Recently, there has been a focus on the development of hypnotic agents that selectively target melatonin receptors [61–63].

Ramelteon ((5)-N-[2-(1,6,7,8-tetrahydro-2H-inden-5-yl)[2-(1,6,7,8-tetrahydro-2H-inden-5-yl)ethyl]propionamide] is a melatonin receptor agonist that shows high selectivity for the MT₁/MT₂ receptors [64]. Figure 1 compares the chemical structures of melatonin and ramelteon. Ramelteon has a half-life of 0.83–1.90 h (much longer than that of melatonin [58]) and undergoes a rapid, high first-pass metabolism [65,66]. In clinical trials, ramelteon has demonstrated sleep-promoting effects, with no next-day residual effects, rebound insomnia, or withdrawal effects, making it the first available nonscheduled prescription insomnia medication [16,67–69]. This review summarizes the preclinical data, clinical efficacy, and safety profile of ramelteon.

Preclinical Pharmacology

Neurochemical Effects

High Affinity for MT₁/MT₂ Receptors

In vitro studies have demonstrated that ramelteon is a potent and highly selective MT₁/MT₂ receptor agonist (Table 1) [70]. In studies of Chinese hamster ovary (CHO) cells expressing human melatonin receptors, ramelteon exhibited a 6-fold higher affinity for the MT₁ receptor than that by melatonin [70]. Similarly, ramelteon had a higher affinity for the MT₂ receptor than that of melatonin. The dissociation equilibrium constant (Kₐ) and maximal number of binding sites (Bₘₐₓ) values of ramelteon were 15.0 ± 3.0 pM and 555 ± 114 fM/mg protein, respectively, for the MT₁ receptor, and 328 ± 12 pM and 133 ± 2 fM/mg protein, respectively, for the MT₂ receptor. The affinities of ramelteon for the MT₁ and MT₂ receptors were comparable to those of 2-iodomelatonin. N-acetyl-5-HT, a precursor of melatonin with high affinity for the MT₁ binding site, showed
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Figure 1 Comparison of chemical structures of melatonin and ramelteon, (S)-N-[2-(1,6,7,8-tetrahydrosso-2H-indeno-5,4-b]furan-8-yl)ethyl]propionamide.

Table 1 Receptor binding characteristics

|                | Affinity (K_i) for human MT_1 receptor expressed in CHO cells | Affinity (K_i) for human MT_2 receptor expressed in CHO cells | Affinity (K_i) for MT_3 binding site in hamster brain |
|----------------|---------------------------------------------------------------|---------------------------------------------------------------|-------------------------------------------------------|
| Ramelteon      | 14 ± 0.5 pM                                                   | 112 ± 5 pM                                                   | 2650 ± 180 nM                                         |
| Melatonin      | 80.7 ± 2.1 pM                                                | 383 ± 5 pM                                                   | 24.1 ± 0.5 nM                                         |
| 2-iodomelatonin| 13.1 ± 0.3 pM                                                | 188 ± 4 pM                                                   | 0.964 ± 0.015 nM                                      |
| N-acetyl-5-HT  | 81,300 ± 6000 pM                                             | 3,640,000 ± 30,000 pM                                        | 15.7 ± 2.8 nM                                         |
| Prazosin       | >2,730,000 pM                                                 | >5,370,000 pM                                                | 6.16 ± 0.46 nM                                        |

Each value represents the mean and standard error [70].

very low affinity for the MT_1 and MT_2 receptors. Prazosin, an α_1 antagonist with high affinity for the MT_3 binding site [71,72], showed negligible affinity for the MT_1 and MT_2 receptors [70].

In studies of Syrian hamster brain, the affinity of ramelteon for the MT_3 binding site was extremely weak (K_i: 2.65 µM) relative to melatonin and 2-iodomelatonin; the values being 24.1 nM and 0.924 nM, respectively [70]. The affinity of ramelteon for the MT_3 binding site was 1/110 that of melatonin and 1/2750 that of 2-iodomelatonin. N-acetyl-5-HT and prazosin also showed high affinities for the MT_3 binding site [70], as reported previously [38,71].

Agonistic Effect on MT_1/MT_2 Receptors

Melatonin receptors, when activated, inhibit adenylate cyclase, which leads to a decrease in cAMP [73,74]. Ramelteon demonstrated this agonist activity at both MT_1 and MT_2 receptors. In CHO cells expressing the human melatonin receptors, ramelteon inhibited forskolin-stimulated cAMP production in a concentration-dependent manner. [70] The half maximal inhibitory concentration (IC_{50}) values of ramelteon and melatonin for the inhibition of cAMP production in cells expressing MT_1 receptors were 21.2 and 77.8 pM, respectively, and in cells expressing MT_2 receptors, were 53.4 and 904.0 pM, respectively (Table 2) [70]. Taken together, ramelteon’s affinity and activity at MT_1/MT_2 receptors and its low affinity for the MT_3 binding site make it a potentially more suitable sleep-promoting agent than exogenous melatonin.
Effects on Other Receptors and Enzymes

To help differentiate ramelteon from sedative hypnotics that target the GABA<sub>A</sub>-receptor complex, investigators assessed ramelteon’s affinity to GABA and a wide variety of other CNS binding sites [70]. As expected, ramelteon did not show significant inhibition of binding (≥50%) to 131 G-protein-coupled receptors, transporters, and ion channels, including monoamine receptors, opioid receptors, central benzodiazepine receptors, and dopamine transporters tested at 10 μM [70]. Moreover, ramelteon showed no effect on enzyme activity of 54 tested enzymes at 10–1000 μM [70]. Melatonin caused no significant inhibition of the binding of almost all receptors, except for 5-HT<sub>1A</sub> receptor, for which the K<sub>i</sub> value was 5.6 μM. Ramelteon’s negligible affinity to GABA, serotonin, acetylcholine, glutamate, noradrenaline, opioid, histamine, and dopamine receptors is noteworthy, as ancillary activity at these receptors may result in unwanted secondary or residual effects.

Sleep-Promoting Effects

To study the sleep-promoting effects of ramelteon in cats and monkeys using polysomnography (PSG), a chamber was constructed to allow animals to move freely while being attached to electroencephalogram (EEG) cables [49]. In freely moving cats, the effect of ramelteon on sleep was stronger and lasted longer than the effect of exogenous melatonin (Fig. 2) [49]. Ramelteon (0.001, 0.01, and 0.1 mg/kg, p.o.) was shown to promote sleep, as evidenced by decreases in the percentage of time spent awake and increases in the percentage of time spent in slow wave sleep (SWS) and REM sleep. The median latency to sleep onset (time to first SWS lasting more than 1 min) was 24 min in cats given ramelteon 0.1 mg/kg compared with 60 min in cats given vehicle control. In contrast, melatonin resulted in a mild sleep-promoting effect; melatonin (0.001–1.000 mg/kg, p.o.) showed no statistically significant effect on sleep latency, and the duration of the sleep-promoting action of the highest dose of melatonin (2 h) was shorter than that of ramelteon (6 h).

Ramelteon was shown to cause sleep promotion in freely moving cats. However, it was not enough for starting clinical trial, because the experimental conditions were artificial and the effect of ramelteon was on daytime sleep in cats. It was critical for the project to confirm pharmacological efficacy in primates, focusing on the effect on nighttime sleep, which is much more close to the clinical use of ramelteon. Since there was no evaluation system to assess PSG in freely moving monkeys, the authors developed the PSG assay system through a try and error process [75]. In the study, young adult female crab-eating macaques (Macaca fascicularis), implanted with electrodes for EEG, electrooculogram (EOG), and electromyogram (EMG) recording, were used, well habituated to the recording chamber located in a soundproof, electrically shielded room that was maintained under conditions similar to those of the home cage under freely moving unrestrained condition. Sleep architecture in the unrestrained monkeys was similar to that of humans, which was also confirmed by recent reports [76].

Ramelteon (0.03 and 0.3 mg/kg, p.o.) showed sleep-promoting effects, as evidenced by statistically significant increases in total duration of sleep and decreases in latencies to sleep onset (time to the first consecutive 3 min of sleep stage) of light sleep (sleep stages 1 and 2) and SWS (sleep stages 3 and 4) (Figs. 3–5) [75]. Melatonin (0.3 mg/kg, p.o.) showed a statistically significant reduction in latency to onset of light sleep but not of SWS and had no statistically significant effect on sleep duration [75]. The higher doses of melatonin (1 and 3 mg/kg) had no statistically significant effect on the latencies to each sleep stage or sleep duration. The benzodiazepine receptor agonist zolpidem (1–30 mg/kg, p.o.) showed no statistically significant effects on sleep latencies to each sleep stage or sleep duration [75]. With regard to observable behavior, monkeys given zolpidem 30 mg/kg displayed sedation and myorelaxation, whereas monkeys given either ramelteon or melatonin at any doses showed no apparent motor dysfunction (Fig. 5) [75]. As zolpidem showed marked sedation including myorelaxation at 100 mg/kg, p.o., the authors did not assess the sleep-inducing action at this dose. They suggested that, if they conduct the study at 100 mg/kg, it might cause more apparent effect. Therefore, zolpidem might cause sleep-inducing action at doses exerting marked sedation in monkeys. A number of studies showed that zolpidem causes potent reduction of sleep latency in humans [11,14,77], suggesting that there might be a species difference in sleep induction by zolpidem. Ramelteon significantly shortened sleep latency and increased total sleep time in monkeys. Similar clinical effects have been confirmed in clinical studies [69,78], suggesting that the preclinical study using monkeys was valid for the prediction of clinical efficacy in humans.

Unlike the zolpidem, which increased EEG fast waves (≥14 Hz) during sleep, EEG spectra and a fast Fourier transform (FFT) after ramelteon and melatonin administration were indistinguishable from those of naturally occurring physiological sleep (Fig. 6) [75]. EEG power spectral analyses in humans have shown that treatment with benzodiazepines or zolpidem caused a significant reduction of low-frequency activity (0.25–10.0 Hz) and a significant increase in high-frequency activity in non-REM sleep observed in healthy young subjects.
Figure 2  (A) Effects of ramelteon on sleep and wakefulness in freely moving cats. Each value shows the mean (with standard error) percentage of time spent in the stages of wakefulness, SWS, or REM sleep during each block of 2 h after drug administration. Eight of 14 cats were randomly used in each dose group. *P ≤ 0.05, **P ≤ 0.01, compared with the vehicle-treated control (ANOVA). #P ≤ 0.05, ##P ≤ 0.01, compared with the vehicle-treated control (paired t-test with Holm’s correction) [49]. (B) Effects of melatonin on sleep and wakefulness in freely moving cats. Each value shows the mean (with standard error) percentage of time spent in the stages of wakefulness, SWS, or REM sleep during each block of 2 h after drug administration. Eight of 14 cats were randomly used in each dose group. *P ≤ 0.05, **P ≤ 0.01, compared with the vehicle-treated control (ANOVA). #P ≤ 0.05, ##P ≤ 0.01, compared with the vehicle-treated control (paired t-test with Holm’s correction) [49].

Therefore, sleep induced by benzodiazepine receptor agonists may be qualitatively different from natural sleep in terms of EEG findings. EEG spectral analyses showed that sleep induced by ramelteon and melatonin was indistinguishable from that of vehicle-treated control [75], implying that MT₁/MT₂ receptor agonists may promote physiological sleep in animals.

Circadian Reentrainment

To assess the ability of ramelteon to shift the circadian rhythm, a study of running-wheel activity in rats was conducted [79]. After an abrupt 8-h advance of the light–dark cycle, the circadian rhythm of running-wheel activity of rats was gradually resynchronized to the new
light–dark cycle. Rats treated with ramelteon (0.1 and 1.0 mg/kg, p.o.) took less time to resynchronize to the new light–dark cycle compared with rats that received vehicle control (Fig. 7). Melatonin 10 mg/kg, p.o., also accelerated reentrainment of running-wheel activity rhythm following the phase advance [79]. These results indicate that ramelteon was closely mimicking the central actions of melatonin as a circadian rhythm signal, not simply acting as a hypnotic.

The MT$_2$ receptor had been shown to be important for the phase-shifting effects of melatonin in the SCN [32,43–45]. Ramelteon induced phase advance by application at zeitgeber time (ZT) 10 but not at ZT 6 in SCN slice in rats. The phase-advancing effect was inhibited by an MT$_2$ receptor antagonist, 4-phenyl-2-acetamidotetraline [80], suggesting that MT$_2$ receptor might be involved in the phase-advancing effect of ramelteon.

Learning and Memory and Motor Function

On the Morris water-maze task for spatial learning and memory, rats treated with ramelteon (3–30 mg/kg, p.o.) and melatonin (10–100 mg/kg, p.o.) showed no statistically significant differences from vehicle control in time.
to find the submerged platform (Fig. 8) or the number of crossings of the area where the platform had been located during training [79]. In contrast, diazepam (3–30 mg/kg, p.o.) and triazolam (0.1–1 mg/kg p.o.) resulted in statistically significant delays in platform reaching times and reductions in the number of platform crossings, indicating an adverse effect on learning and memory function [79].

On the delayed matching to position task for memory and attention, rats treated with ramelteon (3–30 mg/kg, p.o.) and melatonin (10–100 mg/kg, p.o.) showed no impairment on task performance [79]. In contrast, high doses of diazepam (30 mg/kg, p.o.) and triazolam (1 and 3 mg/kg, p.o.) resulted in statistically significant reductions in the number of correct responses across delays [79].
On the conditioned place-preference test for rewarding property, rats treated with ramelteon (3–30 mg/kg, p.o.) or melatonin (10–100 mg/kg, p.o.) showed no preference in the drug-associated compartment, indicating a low potential for reinforcing behavior (Fig. 11) [79]. In contrast, morphine (1 mg/kg, s.c.), triazolam (0.5 mg/kg, p.o.), and diazepam (5 mg/kg, p.o.) resulted in statistically significant increases in the time spent in the drug-associated compartment [79], consistent with reports of the rewarding properties that may occur with the use of benzodiazepines [22,82].

On a task to determine if ramelteon produces benzodiazepine receptor agonist-like discriminative stimulus effects, the monkeys were trained to reliably discriminate between subcutaneous injections of vehicle and the benzodiazepine midazolam using a standard 2-lever procedure for shock avoidance [83]. Ramelteon was then administered intravenously in lieu of midazolam at doses of 0.32, 1.0, 3.2, 5.6, and 10.0 mg/kg. After receiving ramelteon, the monkeys responded primarily on the vehicle-associated lever versus the midazolam-associated lever, indicating that these animals did not generalize ramelteon to midazolam. In a similar study, monkeys dependent on diazepam were trained to reliably discriminate between subcutaneous injections of vehicle and flumazenil (benzodiazepine antagonist) using a standard 2-lever procedure for food presentation [83]. Ramelteon was then administered intravenously at doses of 3.2, 5.6, and 10.0 mg/kg 15 min prior to increasing doses of flumazenil. The percentage of responses on the flumazenil-associated lever was similar between ramelteon 10 mg/kg and vehicle control, indicating that ramelteon did not attenuate the effect of flumazenil.

On an operant task to assess a reinforcing effect, rhesus monkeys were free to self-administer intravenous ramelteon [84]. In the 2-h testing periods, there were no significant differences in the number of self-administrations (fixed ratio of 50) of ramelteon at doses ranging from 0.025–0.400 mg/kg per infusion relative to vehicle control [84]. In contrast, pentobarbital at doses of 0.5 and 1.0 mg/kg per infusion resulted in statistically significant increases in the number of self-administrations. When the monkeys were free to self-administer drug (fixed ratio of 5) around the clock, the mean numbers of self-administrations of ramelteon were comparable to those of vehicle, further indicating that ramelteon showed no positive reinforcing effect. With regard to observable behavior, no gross behavioral changes were noted in monkeys treated with ramelteon or vehicle.
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Figure 6  Typical electroencephalographic (EEG) spectra and fast Fourier transform (FFT) analysis in freely moving monkey when treated with ramelteon (0.3 mg/kg, p.o.), melatonin (1 mg/kg, p.o.), and zolpidem (30 mg/kg, p.o.). Typical samples were selected during non-REM sleep. EEG, electro-oculogram (EOG) and electromyographic (EMG) activities were recorded [103].

Figure 7  Effects of ramelteon on reentrainment of running-wheel activity rhythm following an 8-hour phase advance. Running-wheel activity in the subjective night as a percentage of total activity in 24 hours is shown with standard error. Ramelteon or vehicle was administered 5–30 min before lights out of the new light-dark cycle for 14 days starting the day of the phase-shift. Arrows and black bars show the phase-shift and the vehicle or ramelteon treatment periods [36].

control. In monkeys treated with pentobarbital, hyporeactivity, slowed motion, and/or ataxia was noted.

In a long-term study to assess physical dependence liability, monkeys received ramelteon (10 mg/kg, i.g. catheter) daily for 1 year [83]. During weeks 14, 27, and 40, the treatment was temporally discontinued for 5 days to assess the effects of treatment discontinuation on operant behavior (conditioned lever pressing to obtain food and avoid shock) and observable clinical signs (such as yawning and grooming) using a paradigm sensitive to benzodiazepine dependence. Daily treatment with ramelteon had no overall effect on monkey’s behavior to obtain food or avoid shock. Operant response rates in individual monkeys did not systematically change over the course of the study. In addition, there were no apparent changes in body weight, motor activity, posture, or behavior observed during the periods of treatment or after discontinuation of treatment.
Figure 8  Effects of ramelteon, melatonin, diazepam, and triazolam on the Morris water maze task in rats. Each value shows the mean time to find the platform submerged in the water (A, B, C, D). *P ≤ 0.025, compared with the respective vehicle control group [one-tailed Williams test, 36].

Figure 9  Percentage of mice failing the rota-rod performance test after administration of ramelteon (A), melatonin (B), N-acetyl-5-HT (C), or diazepam (D) at the indicated doses. Twelve mice were used in each group. **P ≤ 0.01, compared with the control group treated with vehicle (chi-square test with Holm's correction) [81].
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Clinical Efficacy and Safety

Metabolism

Ramelteon is metabolized primarily via oxidation to hydroxyl and carbonyl derivatives, with secondary metabolism producing glucuronide conjugates [65]. Ring-opening biotransformation results in the formation of M-I, and hydroxylation of the carboxy-metabolite, M-III, results in the formation of M-IV (Fig. 12) [65]. The major metabolite of ramelteon in serum is the monohydroxylated metabolite, M-II. Cytochrome P450 (CYP)1A2 is the major isozyme involved in the hepatic metabolism of ramelteon; the CYP2C subfamily and CYP3A4 isozymes are also involved to a minor degree. The major metabolite in serum was M-II, having exposure approximately 39-fold greater than that of ramelteon at the 16-mg dose. The half-life of ramelteon ranges from 0.83 to 1.90 h, depending on the dose [65], which is considerably longer than that of melatonin. This might be regarded as a decisive advantage for the use of ramelteon as a sleep promoter, because the relatively poor efficiency of melatonin is presumed to be partly due to its short half-life [57,58].

M-II is a major metabolite and is physiologically active, and its affinity for chick Mel1a/Mel1c receptors is 0.675 nM, one-fifth to one-tenth the binding affinity of ramelteon, for the human MT1 and MT2 receptors, and it is 17- to 25-fold less potent than ramelteon in in vitro functional assays [70]. M-II also showed a potent sleep-promoting action in freely moving cats [49]. M-II had no significant affinities for other receptors or various enzyme activities, as those of ramelteon, suggesting that M-II is also an MT1/MT2 receptor selective agonist [70]. However, the levels of other metabolites are low, and also the affinities for Mel1a/Mel1c receptors are negligible, as demonstrated by K_i values of 36.0 nM for M-III and >236 nM for both M-I and M-IV. Thus, M-II may contribute to pharmacological effects of ramelteon.

Figure 10 Percentage of mice failing the rota-rod performance test after administration of diazepam alone or diazepam in combination with ramelteon (A), melatonin (B), N-acetyl-5-HT (C) at the indicated doses. Twelve mice were used in each group. *P ≤ 0.05 **P ≤ 0.01, compared with the control group treated with vehicle (chi-square test with Holm’s correction) [53].

Figure 11 Effect of ramelteon on place preference in rats. Increase in time spent in the compound-associated compartment in the conditioned place-preference tests in rats in the experiment 1 (A and C) and experiment 2 (B). The numbers of rats used are shown in parentheses. Vehicle groups in panels A and B are identical. *P ≤ 0.05, compared with the value at the preconditioning phase (paired t-test) [79].
Sleep-Promoting Effects

In randomized, double-blind, placebo-controlled clinical trials, the sleep-promoting effects of the short- and long-term use of ramelteon have been demonstrated in patients with chronic insomnia as well as in healthy subjects subjected to a first-night-effect model of transient insomnia. Overnight monitoring with PSG (EEG, EOG, and EMG) was used to objectively evaluate the ability of ramelteon to reduce latency to persistent sleep, maintain sleep, and increase the duration of sleep. Patients’ subjective assessments of sleep were also evaluated with a post-sleep questionnaire. Latency to persistent sleep (defined as time in minutes to the first of 20 consecutive epochs of sleep on PSG) [78,85] or patient-reported sleep latency was the primary measures in these trials.

Transient Insomnia

Ramelteon’s ability to improve transient insomnia was assessed in two clinical trials that utilized a first-night-effect model of transient insomnia. In Roth et al.’s trial of 375 healthy subjects unfamiliar with a sleep laboratory environment (i.e., experiencing transient insomnia), ramelteon at doses of 16 mg and 64 mg produced statistically significant decreases in latency to persistent sleep and increases in total sleep time, as measured with PSG [85]. The mean latency to persistent sleep was 14.1 and 15.5 min in the ramelteon 16-mg and 64-mg groups, respectively, compared with 24.6 min in the placebo group (P ≤ 0.001 for both groups vs. placebo). According to a post-sleep questionnaire, subjects in the 16-mg group reported statistically significant improvements in subjective sleep latency (P = 0.013; no P-value reported for subtotal sleep time). Ramelteon was not associated with next-day residual psychomotor impairment, as measured by the Digit Symbol Substitution Test (DSST). However, the use of the ramelteon 64-mg dose was associated with small but statistically significant reductions in subjective next-day levels of alertness and ability to concentrate relative to placebo. Zammit et al. conducted a similar trial of 289 healthy subjects that showed that latency to persistent sleep, as measured with PSG, was 12.2 min in the ramelteon 8-mg group and 14.8 min in the ramelteon 16-mg group compared with 19.7 min in the placebo group; the 8-mg dose reached statistical significance (P = 0.004) [68]. The mean total sleep time was 436.8 min and 433.1 min in the 8-mg and 16-mg groups, respectively, compared with 419.7 min in the placebo group (P ≤ 0.05 for both groups vs. placebo). Subjective measures of sleep latency and total sleep time were improved, but the differences from placebo were not statistically significant. In this trial, ramelteon had no significant effects on the DSST and subjective levels of alertness and ability to concentrate as well as on other next-day residual effect measures including memory recall tests and visual analog scales (VAS) for mood and feeling.

Chronic Insomnia

Patients with a history of chronic insomnia (primary insomnia as defined by Diagnostic and Statistical Manual
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Figure 13 Effects of ramelteon on sleep latency in patients with chronic insomnia under placebo-controlled, double-blind, randomized, 5-period crossover study. All data are shown as the least square means. ***P ≤ 0.001, compared with placebo [78].

of Mental Disorders (DSM)-IV-TR for at least 3 months) were also evaluated in clinical trials. Figure 13 illustrates the differences from placebo in latency to persistent sleep by dose of ramelteon in these trials. In Erman et al.’s multiple-dose cross-over study of 107 adults with chronic insomnia, a 2-night administration of ramelteon resulted in statistically significant reductions in latency to persistent sleep (P ≤ 0.001) and increases in total sleep time (P ≤ 0.05), as measured with PSG, at each dose tested (4–32 mg) [78]. Interestingly, all doses of ramelteon produced similar reductions in latency to persistent sleep (13.4–14.8 min more than placebo), with no apparent dose-dependent effects, and this wide therapeutic window was seen across studies. Patients who received ramelteon 16 mg reported significant reductions in subjective sleep latency (P ≤ 0.05); no other statistically significant effects in subjective sleep ratings were observed. Ramelteon had no statistically significant effect on next-day residual effect measures including the DSST, memory recall tests, and patient-reported levels of alertness and ability to concentrate. However, ramelteon had no significant effect on the mean wake time after sleep onset (WASO).

In Zammit et al.’s extended trial in 405 patients with chronic insomnia, ramelteon 8 mg and 16 mg resulted in improvements in latency to persistent sleep and total sleep time over 5 weeks of nightly treatment, as measured objectively by PSG and subjectively by the post-sleep questionnaire (Fig. 14) [69]. Ramelteon had no significant effects on WASO. Ramelteon showed no statistically significant effect on the next-day DSST performance. Patients in the 8-mg ramelteon group demonstrated a small, but statistically significant, decrease in the mean score compared with placebo on the immediate memory recall test at week 3 and the delayed memory recall test at week 1. At other time points, no significant differences between ramelteon and placebo were found on memory function tests. Subjective levels of alertness and ability to concentrate were similar between the ramelteon groups and placebo with two exceptions: patients in the ramelteon 8-mg group reported statistically significant improvements in the ability to concentrate at week 1 and the level of alertness at week 5. This trial also included a 2-day placebo runout period that showed that the discontinuation of ramelteon treatment was not associated with either rebound insomnia or withdrawal effects, as assessed with the Tyer Benzodiazepine Withdrawal Symptom Questionnaire (BWSQ).
The BWSQ is a questionnaire that solicits specific information on 20 symptoms commonly experienced during withdrawal from benzodiazepine receptor agonists. In Roth et al.’s trial of ramelteon in 100 elderly patients (≥65 years) with chronic insomnia, a 2-night administration of either ramelteon 4 mg or ramelteon 8 mg produced statistically significant reductions in latency to persistent sleep and increases in total sleep time, as measured with PSG [86]. The mean latency to persistent sleep was 28.7 min and 30.8 min in the 4-mg and 8-mg groups, respectively, compared with 38.4 min in the placebo group (P ≤ 0.01 for both groups vs. placebo). The mean total sleep time was 359.4 min and 362.0 min in the 4-mg and 8-mg groups, respectively, compared with 350.4 min in the placebo group (P ≤ 0.05 for both groups vs. placebo). According to the post-sleep questionnaire, patients who received ramelteon 4 mg reported statistically significant reductions in subjective sleep latency (P ≤ 0.05). There were no next-day residual effects with either dose of ramelteon, as measured with the DSST, memory recall tests, and patient-reported levels of alertness or ability to concentrate.

A large 5-week outpatient trial in 829 older adult patients (≥65 years) with chronic insomnia was conducted to specifically assess subjective ratings of sleep [87]. In this trial, nightly administration of either ramelteon 4 mg or ramelteon 8 mg produced reductions in patient-reported sleep latency and increases in patient-reported total sleep time at week 1, and these improvements were sustained throughout the study. At week 1, subjective sleep latency was 70.2 min in the 4-mg group and 70.2 min in the 8-mg group compared with 78.5 min in the placebo group (P ≤ 0.01 for both doses vs. placebo). At week 1, subjective total sleep time was 324.6 min in the 4-mg group and 321.1 min in the 8-mg group compared with 313.9 min in the placebo group (P ≤ 0.01 for 4 mg vs. placebo, P = 0.055 for 8 mg vs. placebo). Ramelteon did not show either rebound insomnia or withdrawal effects, according to subjective sleep latency and BWSQ results during the 7-day placebo runout period.

In the longest trial of ramelteon to date, 1213 patients with chronic insomnia took open-label ramelteon (8 mg or 16 mg) nightly for 1 year [88]. Patient-reported sleep was improved by the first week of treatment and was sustained throughout the remainder of the study. Improvements in sleep latency from baseline at month 1 were 34.0% and 35.1% with the 8-mg and 16-mg dose, respectively, and continued to improve through month 6 (44.7% and 49.1%) and month 12 (50.3% and 52.1%). Improvements from baseline in patient-reported total sleep time at month 1 (15.2% and 16.9%), month 6 (21.6% and 22.7%), and month 12 (25.5% and 23.9%) were also reported with ramelteon 8-mg and 16-mg doses, respectively. There was no evidence of rebound insomnia with ramelteon during the 3-day placebo runout period.

Considering the clinical data thus far, ramelteon has consistently demonstrated significant sleep-promoting effects, as evidenced by reductions in sleep latency and increases in sleep duration, among patients with chronic insomnia and subjects with transient insomnia, during both short-term and long-term treatment, in both the sleep laboratory and outpatient setting by both objective PSG and subjective reports. The efficacy by subjective reports [87] was weaker than that by objective PSG measurement [69,78]. The efficacy was less than that of sedative hypnotics [11,15,2008]. However, the subjective efficacy of ramelteon was more prominent in subjective reports to questionnaire done the next morning in the sleep laboratory than that in subjective reports done at home, possibly suggesting that compliance is more important for subjective evaluation of melatonin receptor agonists. In 2005, ramelteon was approved in the United States for the treatment of insomnia characterized by a difficulty in sleep onset. Current data suggest that ramelteon has no significant effects on WASO, and the efficacy for sleep maintenance insomnia has not been confirmed. Further studies are required to clarify the effect on WASO and search for a possible medication for sleep maintenance insomnia, taking into consideration the dosage of ramelteon and appropriate patient population.

**Safety**

Ramelteon was generally well tolerated across the clinical trials. Most adverse events were considered mild or moderate in severity, and very low incidences of adverse events during ramelteon administration were observed. Only three events were noted to occur at an incidence of ≥2% with ramelteon 8 mg (the recommended therapeutic dose) vs. placebo: somnolence (5% vs. 3%), dizziness (5% vs. 3%), and fatigue (4% vs. 2%).

In the 1-year trial of ramelteon in 1213 adults with chronic insomnia, the overall incidence of adverse events was similar at 6 months and 1 year [89]. The adverse events were predominantly mild or moderate. At 1 year, the adverse events most frequently reported with ramelteon 8 mg and 16 mg, respectively, were nasopharyngitis (10.5% and 14.9%), somnolence (9.5% and 8.1%), upper respiratory tract infection (7.6% and 11.1%), headache (1.9% and 13.5%), and sinusitis (1.9% and 7.8%). Of 38 subjects (3.1%) reporting a serious adverse event, only three adverse events were possibly treatment related. There were no clinically meaningful changes in vital signs, physical examinations, clinical...
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Figure 15 Effects of ramelteon on patient-reported sleep latency in older adults with chronic insomnia. Patients include older adults (≥65 years; n = 829) with chronic insomnia. Placebo, ramelteon 4 mg, or ramelteon 8 mg was taken nightly for 5 weeks, and patient-reported sleep data were collected in sleep diaries. *P ≤ 0.05, **P ≤ 0.01, compared with the placebo control [87].

chemistry, hematology, or urinalysis values and no electrocardiogram trends to suggest adverse effects on cardiac function over the 1 year of ramelteon treatment. No notable changes in endocrine and sexual/reproductive function were observed except for the mean free and total testosterone level, which had a slight decrease in older men that returned to normal by the final visit of the trial.

Abuse Liability

Benzodiazepine receptor agonists, which are the most commonly used prescription medications for treatment of insomnia, have been associated with a potential for abuse [18,21–23]. All benzodiazepine receptor agonists (including the newer agents zolpidem, zaleplon, and eszopiclone) are classified as schedule IV controlled substances, meaning they possess some potential for abuse and the adverse consequences of abuse (e.g., withdrawal, psychomotor impairment, and overdose).

In a study of 14 subjects with a history of sedative hypnotic or anxiety drug abuse, ramelteon doses up to 160 mg (20 times the therapeutic dose) had no effect on a wide variety of instruments including subject ratings of mood and behavior, observer rating of mood and behavior, and assessments of behavioral and cognitive performance [67]. Figure 15 shows that ramelteon showed no statistically significant effect at any time point (0.5–24 h post dose) on representative abuse measures including “liking” scales, the DSST, and circular lights task. Ramelteon had no statistically significant effect on other questionnaire items “drug strength,” “good effects,” and “monetary street value.” Similarly, ramelteon had no statistically significant effect on word recall/recognition task, enter and recall task, and balance task performance at any dose compared with placebo. In contrast, triazolam consistently produced a dose-related effect on all these subjective and behavioral measures, consistent with its profile as a sedative drug with known abuse potential. This trial demonstrates that ramelteon does not share the pharmacologic side effect profile observed among benzodiazepine receptor agonists.

Future Perspective

Ramelteon has been shown to improve sleep disorders in patients suffering from transient and chronic insomnia without causing significant adverse events. However, it has been suggested that melatonin receptor agonists have opportunities to cure various diseases, including circadian rhythm sleep disorders (CRSD), depression, Alzheimer disease, bipolar disorder, cancer, hypertension, urinary incontinence, and so on.

Patients with CRSD often experience sleep disruption, daytime fatigue, and impaired mental and physical function. The prevalence of CRSD is unknown, but it has been estimated that 5–10% of patients referred to sleep disorder clinics have a CRSD [90]. Among blind individuals, non-24-hour sleep-wake syndrome may occur in as many as 50% [91]. Transient CRSD, such as jet lag and shift-work syndrome, may be quite common in the general population. As a pharmacologic therapy for CRSD, melatonin has shown some effectiveness [92–94], and a systematic review of 10 trials concluded that melatonin treatment (0.5–5.0 mg per day for 2–5 days) is effective for alleviating the symptoms of jet lag after travel across several time zones [95].

A number of studies have shown altered melatonin levels in depressed patients and suggested that mood disorders may be related to low melatonin levels, although the reports were not consistent [96,97]. Recently, agomelatine, a melatonin agonist, showed some efficacy in the treatment of major depressive disorder [98,99]. Agomelatine has been reported to have affinity for 5-HT2c receptors; however, the affinity was much less than that for melatonin MT1/MT2 receptor, suggesting that antidepressive effect of agomelatine might be related to the effects via activation of MT1/MT2 receptors. Furthermore, reduced melatonin levels and altered timing of melatonin secretion were observed in patients with bipolar affective disorder [100,101].

The secretion of endogenous melatonin declines with aging [102], and this may be related to the sleep-wake rhythm disturbance in elderly people [103,104]. Abnormal decreases in nocturnal melatonin have also been observed in patients with Alzheimer disease, another population prone to sleep disturbances [103–105].
Administration of melatonin to Alzheimer disease patients has been found to improve significantly sleep and circadian abnormality [106–108]. These findings suggest that melatonin may play a role in significant regulation of various physiological functions. Thus, further studies using ramelteon, a potent and selective MT<sub>1</sub> and MT<sub>2</sub> receptor agonist well tolerated in humans, to examine its clinical efficacy in these disorders would be interesting and may provide an insight into treatment options for these patients.

**Conclusion**

Ramelteon shows selective affinity for MT<sub>1</sub> and MT<sub>2</sub> receptors and acts as a full agonist. Currently, MT<sub>1</sub> and MT<sub>2</sub> receptors have been suggested to be related to sleep promotion and circadian clock such as phase advance. Ramelteon exerts potent sleep promotion in experimental animals including cats and monkeys, without causing any significant adverse effects such as learning and memory impairment, impairment of motor coordination, and drug abuse ability. The sleep induced by ramelteon cannot be distinguished from natural sleep using FFT analysis in monkeys, in contrast to the altered sleep pattern produced by current sedative hypnotics.

In clinical trials, ramelteon decreases sleep latency and increases total sleep time, without causing hangover, addiction, and withdrawal effects. It has no significant effects on psychomotor and cognitive function and does not show drug abuse potential. Taken together, the proven clinical efficacy of ramelteon combined with its safety profile positions it as a fourth-generation insomnia treatment worth considering for select patients suffering from insomnia (Table 3). Further development of ramelteon for various diseases or disorders is expected.

**Conflict of Interest**

The authors have no conflicts of interest.

**Table 3** Classification of therapeutic drugs for insomnia

| Classification                      | Drug                          | Adverse events                                                      | Sleep pattern     |
|-------------------------------------|-------------------------------|---------------------------------------------------------------------|-------------------|
| Barbiturates                        | Pentobarbital, phenobarbital  | Respiratory suppression, amnesia, loss of motor dysfunction, drug abuse, dependence | Sedative sleep    |
| Benzodiazepme receptor agonists      | Triazolam, temazepam          | Amnesia, loss of motor coordination, drug abuse, dependence         | Sedative sleep    |
| Benzodiazepme receptor agonists      | Zolpidem, zaleplon, zopiclone  | Amnesia, loss of motor coordination, drug abuse, dependence         | Sedative sleep    |
| MT<sub>1</sub>/MT<sub>2</sub> receptor agonist | Ramelteon                     | No serious adverse event                                             | Physiological sleep |

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