Esmirtazapine for the Treatment of Chronic Primary Insomnia: A Randomized Long-Term Safety Study in Elderly Outpatients

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Objectives: This 52-week, double-blind, randomized, Phase 3 study evaluated the long-term safety of esmirtazapine 1.5 mg and 3.0 mg in elderly outpatients (aged ≥65 years) with insomnia. Methods: Participants were randomized to receive esmirtazapine 1.5 mg or 3.0 mg administered once nightly. Safety and tolerability (primary objectives) were assessed via adverse event (AE) reporting, routine clinical measurements [vital signs; electrocardiogram (ECG); laboratory parameters], and residual-effects assessments. Total sleep time (TST), wake time after sleep onset (WASO), and sleep latency (SL) were assessed (secondary objectives).

Results: Of 259 randomized participants, 153 completed treatment. AEs and serious AEs were reported by 89.8% and 7.0%, respectively, of 1.5 mg recipients, and 88.5% and 3.8%, respectively, of 3.0 mg recipients. Discontinuations due to AEs were reported in 16.4% and 18.3% of participants receiving esmirtazapine 1.5 mg and 3.0 mg, respectively. The most frequent AEs (>10%) were nasopharyngitis, somnolence, dizziness, headache, dry mouth, weight increase, and fatigue. One participant died; the death was judged unrelated to treatment. Elevated eosinophil counts were noted, but not considered clinically significant. No remarkable or clinically relevant changes in laboratory parameters, vital signs, or ECG were observed. There was no evidence of residual effects; alertness at awakening increased by a median of 17 (1.5 mg) and 15 (3.0 mg) points from baseline, respectively, and ability to work/function by 12 points (both groups; all \( p<0.0001 \)). Improvements from baseline in TST, WASO, and SL were observed. Conclusions: Esmirtazapine was reasonably tolerated in elderly outpatients with insomnia. No significant safety signals were observed.

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Key Words: Elderly, Esmirtazapine, Insomnia, Outpatients, Safety.

Introduction

Insomnia is a highly prevalent disorder, characterized by either non-restorative sleep or difficulties initiating or maintaining sleep,¹ which affects up to approximately 20% of adults in the United States (U.S.).² Compromised sleep quality impacts on the daytime functioning of affected individuals, especially in social and occupational domains, leading to impaired overall quality of life and an increased risk of psychiatric comorbidities.³,⁴ The prevalence of insomnia increases with age, with estimates ranging from 20–50% in the elderly population,⁶ which makes elderly adults the most common users of hypnotic drugs.⁷ Insomnia in the elderly is associated with affective disorders, increased healthcare utilization,⁹ higher risk of accidents and falls,¹⁰ and cognitive impairment that can be misdiagnosed as dementia.¹¹ Currently, first-line pharmacotherapy for insomnia relies on the use of positive allosteric modulators of gamma-aminobutyric acid type A (GABA_\(A_\)) receptors, which induce sedation through the potentiation of the inhibitory signaling of GABA_\(A_\) receptors.¹²,¹³ Although effective in promoting sleep, GABA_\(A_\) receptor modulators elicit global inhibitory effects in the central nervous system that make them liable to producing a variety of side effects, including rebound insomnia, memory failure, and abuse potential.¹¹,¹³ Despite being the primary treatments for insomnia in elderly patients, GABA_\(A_\) receptor modulators may not be an ideal treatment for this patient population, as they are associated with impairment of motor function and coordination in elderly patients, along with increased rates of falls and motor vehicle crashes.¹⁴,¹⁵

Safety concerns associated with GABA_\(A_\) modulators prompt-
ed the clinical development of sleep-promoting compounds that facilitate sleep via more specific and targeted mechanisms of action, and this was anticipated to translate into an improved side-effects profile. For example, orexin receptor antagonists (ORAs) have been developed to promote sleep by blocking physiological effects of the orexin compounds, which are key mediators in the maintenance of arousal and vigilance.16 The ORA suvorexant has demonstrated sleep-promoting effects with minimal residual next-day somnolence or psychomotor impairment in clinical studies, including in elderly insomnia patients.7,17-20 Other pharmacological treatments for insomnia include selective antagonists of histamine H1 and α1-adrenergic receptors, antidepressants and antipsychotic drugs, which are typically prescribed off-label, melatonin receptor agonists, and over-the-counter antihistamines.11 Non-pharmacological therapies, such as cognitive behavioral therapy, have also shown efficacy as an alternative to pharmacotherapy.21

Treatment with low doses of sedating antidepressants may be a promising approach for those patients who are more likely to experience sleep-maintenance insomnia, such as the elderly. Antidepressants have relatively discrete activity and act by modulating neurotransmitters, such as histamine, serotonin, and norepinephrine.22 To date, low doses (3 and 6 mg) of the antidepressant doxepin (SilenorTM, Currax Pharmaceuticals LLC, Morristown, NJ, USA), a modulator of the histamine H1 receptor, have been approved in the U.S. for the treatment of insomnia characterized by difficulties with sleep maintenance.23 At low doses, doxepin demonstrated efficacy in improving sleep maintenance and was associated with a favorable safety profile in both younger (aged 18–64 years)24 and older (aged ≥65 years) adults.25 Several other antidepressants have been prescribed off-label in the management of insomnia, but the scientific evidence supporting their use is weak.26

Mirtazapine is an antidepressant with sleep-promoting properties27,28 that acts via antagonism of histamine H1 receptors, 5-HT1, and 5-HT3 serotonergic receptors, and adrenergic α2-autoreceptors.29 It is a racemic mixture of S(+) and R(-) enantiomers, with the former characterized by a shorter half-life compared with the latter. The maleic salt of S(+) mirtazapine, referred to as esmirtazapine, was advanced to the clinical phase of action in an investigational therapy for insomnia based on the consideration that the shorter half-life of esmirtazapine (10 h) relative to the racemic mixture (range: 20 to 40 h)30 would diminish risk of the emergence of associated side effects, in particular residual next-day effects. In clinical studies, acute (2-day) and 2- and 6-week treatment with esmirtazapine in non-elderly participants (aged 18 to <65 years) with insomnia demonstrated significant improvements in objective and subjective sleep measures at doses of 1.5 mg, 3.0 mg, and 4.5 mg relative to placebo, and was generally well tolerated, with minimal residual daytime effects31-34 and no signs of rebound insomnia.31,32

Given the burden of insomnia and the high prevalence of sleep-aid use in the geriatric population, the current study was conducted in elderly outpatients (aged ≥65 years) with chronic primary insomnia. The primary objective was to investigate the safety and tolerability of long-term (52-week) treatment with esmirtazapine at doses of 1.5 mg and 3.0 mg. The secondary objective was to collect exploratory efficacy data with respect to sleep improvement.

**Methods**

**Study design and treatments**

This Phase 3 study (Merck & Co., Inc., Kenilworth, NJ, USA, Protocol Number 176005; NCT00561574) was part of the esmirtazapine clinical development program. Since this study was conducted, the sponsor (Merck Sharp & Dohme Corp., a subsidiary of Merck & Co., Inc.) discontinued further development of esmirtazapine. The study was conducted between January 2008 and February 2010 in 32 centers across Canada, Denmark, Finland, Germany, the Netherlands, Sweden, and the U.S. The trial was carried out in compliance with the ethical principles of the Declaration of Helsinki and in accordance with the International Conference on Harmonisation guidelines for Good Clinical Practice, as well as applicable local regulatory requirements. The study was approved by the appropriate Institutional Ethics Committee or Review Board. Participants were required to sign an informed consent form after receiving an explanation of the scope and nature of the investigation.

This was a randomized, double-blind, parallel-group (assignment to two doses of esmirtazapine), multisite study undertaken in elderly participants over a period of 52 weeks. The study design comprised a 14-day screening process during which participants’ eligibility for the study was ascertained based on prespecified inclusion criteria (Fig. 1). This was followed by randomization (Day 1) using an Interactive Response (telephone) system, which assigned a 6-digit randomization number to eligible participants. Participants were randomized to receive either 1.5 mg or 3.0 mg esmirtazapine (manufactured by NV Organon, Oss, The Netherlands, a subsidiary of Merck & Co., Inc.) before bedtime for a treatment duration of 364 days (52 weeks). Participant visits occurred at the end of Weeks 1, 2, and 4 during the first month of the trial, and every 4 weeks thereafter. A follow-up visit was scheduled 1 week after study completion or premature discontinuation. A telephone call was scheduled 30 days after the last dose to follow-up on any adverse events (AEs) occurring after the follow-up visit.
Participants
Eligible participants were aged $\geq 65$ years at screening and had a normal bedtime within the 21:00–01:00-hours range, with variations not exceeding 2 h for five out of seven nights. Participants were required to demonstrate the ability to independently complete a sleep questionnaire in the week preceding randomization using the LogPad, a handheld electronic data-recording device that captures the sleep diary with respect to subjective drug-efficacy parameters and measures of drug residual effects. Further, participants had a documented diagnosis of chronic primary insomnia according to the Diagnostic and Statistical Manual of Mental Disorders, 4th Edition, Text Revision (DSM-IV-TR) criteria for primary insomnia (DSM-IV-TR 307.42) with a duration of $\geq$ 1 month, and met the following criteria based on medical or sleep history for at least 3 nights per week for at least 1 month: total sleep time (TST) $\leq 6.5$ h; wake time after sleep onset (WASO) $\geq 60$ min; and sleep latency (SL) $\geq 30$ min.

Participants with other sleep disorders based on DSM-IV-TR criteria, those presenting with sleep disturbances caused by significant medical or DSM-IV-TR psychiatric illness, and those who were currently, or within the last 2 years had been, diagnosed or treated for depression were excluded. Participants with signs of dementia or other serious cognitive impairment, defined as a score of <26 on the Mini-Mental State Examination, were also excluded.

Safety and tolerability evaluations
The safety and tolerability analyses (primary objective) were performed in the all-participants-treated population, which was defined as all participants who received at least one dose of trial medication. Recording of AEs was performed from signing the informed consent form until 30 days after end of treatment. Clinical examinations evaluating vital signs and body weight were conducted at screening (Day -14) and at every visit, including the 7-day follow-up visit. Physical examinations were scheduled at screening (Day -14) and Week 52. In addition, electrocardiogram (ECG) was assessed at Days -14 and -7 and Weeks 2, 24, and 52, and laboratory parameters, including hematology, biochemistry, and urinalysis, were evaluated at screening and Weeks 2, 4, 12, 24, 36, and 52.

Residual effects (special safety measures) were assessed based on a sleep diary completed weekly on the LogPad from Day 1 until Week 52. The sleep diary captured participants’ ratings of alertness at awakening, energy level, and ability to work/function during the day on visual analog scales (VAS) from 0–100 mm. Participants also reported their tendency to nap during the day (“Yes/No”) and indicated the napping time per day (in h and min).

Efficacy evaluations
Exploratory efficacy evaluations (secondary objective) were performed in the intent-to-treat population of all randomized participants who received at least one dose of trial medication and had at least one recorded post-baseline efficacy measurement on the weekly sleep diary. Efficacy measures included TST, WASO, SL, number of awakenings, and sleep quality based on weekly sleep diaries completed from Day 1 until Week 52. In addition, the extent of sleep difficulties was assessed on Day -7 and Weeks 2, 4, 12, 24, 36, and 52 using the insomnia severity index (ISI) based on participant ratings. The investigator global rating (IGR) scale was used by the clinician to rate insomnia symptom severity (IGR-severity (IGR-S)) and the therapeutic effect of the treatment regimen. These evaluations were completed at Day -7 (IGR-S score only) and Weeks 2, 4, 12, 24, 36, and 52.

Statistical methods
Safety and efficacy parameters were evaluated using summary and descriptive statistics, respectively, with changes from...
baseline where appropriate. Continuous variables were analyzed by summary statistics. Paired t-statistics were used for testing the change from baseline by visit and by week as an exploratory analysis for efficacy endpoints. The ratings on the weekly sleep diary collected at Day 1 were used as baseline. No primary or secondary hypotheses for the testing of either efficacy or safety were performed. Categorical endpoints were summarized by treatment group, and summary statistics included sample size and percent for each response category. The denominator for percentage calculations was the total number of participants with non-missing values within each treatment group in that specific time point.

Data from weekly sleep diaries were analyzed using the last observation carried forward (LOCF) approach; sensitivity analyses were conducted based on observed case (OC) data. As the primary objective of this study was to assess long-term safety, the efficacy analysis is presented descriptively and with p values as exploratory analyses.

The planned number of participants was based on the need to collect long-term safety data in elderly patients. Assuming a discontinuation rate of 50–80%, a sample size of 250 participants was considered sufficient. A sample size of 130 participants in each group would have an effect size of 1.1 to detect a difference in means of 86 min for TST duration, assuming the common standard deviation was 80 min using a two-sample t-test with a 5% two-sided significance level.

**Results**

**Participants**

Of the 348 participants screened for this trial, 259 were randomized and treated, with 128 participants allocated to the 1.5-mg esmirtazapine treatment group and 131 participants assigned to the 3.0-mg treatment group (Fig. 2). A total of 153 participants (59.1%) completed the study, while 106 participants discontinued prematurely: 48 participants in the 1.5-mg group and 58 participants in the 3.0-mg group. A total of 173 participants were exposed to at least 6 months of treatment, including 82 (62.6%) in the 3.0-mg group and 91 (71.1%) in the 1.5-mg group. AEs were the most common reason for discontinuation [21 participants (16.4%) in the 1.5-mg group and 24 participants (18.3%) in the 3.0-mg group]. Insufficient therapeutic effect was the next most common reason for discontinuation and was more prevalent in the 3.0-mg group [21 participants (16.0%)] than the 1.5-mg group [11 participants (8.6%)].

The majority of participants were female (62.5%), white (97.7%), and of non-Hispanic/Latino descent (97.7%), with a mean (standard deviation) age of 70.9 (±5.2) years, and a mean body mass index (BMI) of 26.9 (±3.8) kg/m². The most reported BMI category was overweight (43.2%) (Table 1).

**Safety and tolerability**

A total of 231 participants experienced at least one AE during the in-treatment period, including 115 participants (89.8%) in the 1.5-mg group and 116 participants (88.5%) in the 3.0-mg group (Table 2). AEs judged by the investigators to be treatment-related were more frequently reported in the 3.0-mg group [88 participants (67.2%)] than the 1.5-mg group [74 participants (57.8%)]. The most frequent AEs, with an incidence of >10%, were nasopharyngitis, somnolence, dizziness, headache, dry mouth, weight increase, and fatigue. Somnolence (26.7% vs. 11.7%), dizziness (14.5% vs. 9.4%), dry mouth (10.7% vs. 7.8%), and weight increase (15.3% vs. 8.6%) occurred more commonly in the 3.0-mg group vs. the 1.5-mg group. The majority of cases of somnolence, dizziness, dry mouth, weight increase, and fatigue.

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**Figure 2.** Participant disposition. *participants took at least one dose of trial medication, †information as per End of Trial Form, ‡one participant in the 1.5-mg group discontinued due to lack of compliance with study diary. AE: adverse event, SAE: serious AE.
increase, and fatigue were considered treatment-related by the investigator (Table 2). Approximately 10% of the participants in both the 1.5-mg and 3.0-mg groups experienced AEs of severe intensity, including one event of dizziness and one event of fatigue in the 1.5-mg treatment group, and one event of somnolence, one event of fatigue, and two events of dry mouth in the 3.0-mg group, which were considered probably or possi-

| Table 1. Participant demographics at screening |
|------------------------------------------------|
| **Variables** | **Esmirtazapine 1.5 mg** (n=128) | **Esmirtazapine 3.0 mg** (n=131) | **Total** (n=259) |
| Age (years) | | | |
| Median (range) | 69.0 (65.0–86.0) | 70.0 (65.0–87.0) | 69.0 (65.0–87.0) |
| Mean (SD) | 70.7 (5.2) | 71.1 (5.1) | 70.9 (5.2) |
| Height (cm) | | | |
| Median (range) | 165.0 (139.7–198.1) | 166.4 (144.0–188.0) | 166.0 (139.7–198.1) |
| Mean (SD) | 167.3 (10.5) | 167.4 (9.7) | 167.4 (10.1) |
| Weight (kg) | | | |
| Median (range) | 73.7 (48.6–124.3) | 75.0 (45.3–107.0) | 73.9 (45.3–124.3) |
| Mean (SD) | 76.1 (15.8) | 75.5 (13.2) | 75.8 (14.5) |
| Gender, n (%) | | | |
| Female | 83 (64.8) | 79 (60.3) | 162 (62.5) |
| Male | 45 (35.2) | 52 (39.7) | 97 (37.5) |
| Race, n (%) | | | |
| Black or African-American | 4 (3.1) | 2 (1.5) | 6 (2.3) |
| White | 124 (96.9) | 129 (98.5) | 253 (97.7) |
| Ethnicity, n (%) | | | |
| Hispanic or Latino | 2 (1.6) | 4 (3.1) | 6 (2.3) |
| Not Hispanic or Latino | 126 (98.4) | 127 (96.9) | 253 (97.7) |
| BMI (kg/m²) | | | |
| Median (range) | 26.3 (17.9–35.8) | 26.6 (19.2–35.4) | 26.5 (17.9–35.8) |
| Mean (SD) | 27.0 (4.0) | 26.9 (3.7) | 26.9 (3.8) |
| BMI category, n (%) | | | |
| Underweight (BMI<18.5) | 1 (0.8) | 0 (0.0) | 1 (0.4) |
| Normal range (18.5≤BMI<25) | 43 (33.6) | 43 (32.8) | 86 (33.2) |
| Overweight (25≤BMI<30) | 55 (43.0) | 57 (43.5) | 112 (43.2) |
| Obese (BMI≥30) | 29 (22.7) | 31 (23.7) | 60 (23.2) |

BMI: body mass index, SD: standard deviation

bly drug-related.

One participant in the 1.5-mg treatment group died due to narcotic intoxication. The autopsy report indicated a history of asthma, heroin abuse, recent depression, and remote hand injuries. Toxicological studies were positive for ethanol and morphine. Based on these findings, the cause of death was described as morphine toxicity with arteriosclerotic cardiovascular disease listed as a contributing condition. The death was judged as unlikely to be related to study drug by the investigator and the sponsor.

During the in-treatment period, a total of 13 serious AEs (SAEs) were reported in nine participants (7.0%) in the 1.5-mg group, and seven SAEs were reported in five participants (3.8%) in the 3.0-mg group. In the 1.5-mg group, one participant with a history of arterial hypertension experienced two SAEs (cardiac failure and hypertensive heart disease), which the investigator judged to be possibly drug-related, whereas the sponsor deemed these SAEs unlikely to be related to the study drug. Another participant in the 1.5-mg group experienced three SAEs (contusion, dizziness, and fall) that were judged by the investigator as unlikely to be related to the study drug; however, the sponsor considered these events to be possibly related to the study drug. The remaining SAEs were considered by both the sponsor and the investigator to be either not related or unlikely related to the study drug. Except for the deceased participant, and one participant who recovered from cardiac events but also experienced intervertebral disc protrusion and recovered with sequelae (paraparesis of both legs), all participants in the 1.5-mg and 3.0-mg groups recovered or were still recovering from their SAEs at trial completion. During the post-treatment period (2 days after the 30-day follow-up period), one participant treated with 3.0 mg esmirtazapine reported an SAE (hernia) that was not considered to be treatment-related.

There were no clinically meaningful changes in hematology parameters over the course of the study, with the exception of notable increases from baseline in both the percentage of eosinophils, which increased by a median of 16.7% and 20.4% with esmirtazapine 1.5 mg and 3.0 mg, respectively, and absolute eosinophil counts, which increased by a median of 20.0% and 14.3% with esmirtazapine 1.5 mg and 3.0 mg, respectively, over the course of the study. However, none of the abnormally high values were considered to be clinically significant.

Although there were no clinically significant changes in mean or median values for biochemical parameters between baseline and the last measurement, there were some differences observed among treatment groups: higher median percentag-

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2.0%), total bilirubin (1.5 mg: 4.9%; 3.0 mg: 6.3%), and glucose (1.5 mg: 3.9%; 3.0 mg: 10.3%), and lower for gamma-glutamyl transferase (1.5 mg: 5.3%; 3.0 mg: 0.0%). There were no clinically relevant changes in urinalysis, vital signs, or ECG parameters.

Residual effects

Based on the daytime functioning questionnaire in the weekly

Table 2. Safety profile of esmirtazapine

| | Esmirtazapine 1.5 mg | Esmirtazapine 3.0 mg |
|---|----------------|----------------|
| (n=128; n, %) | (n=131; n, %) |
| Overall safety profile | | |
| Participants with AEs | 115 (89.8) | 116 (88.5) |
| Deaths ¹ | 1 (0.8) | 0 (0.0) |
| Participants with SAEs ² | 9 (7.0) | 5 (3.8) |
| Participants who discontinued due to AEs | 21 (16.4) | 24 (18.3) |
| Participants with drug-related AEs ³ | 74 (57.8) | 88 (67.2) |
| Participants with severe AEs | 13 (10.2) | 14 (10.7) |

| AEs occurring in ≥2.5% of participants in any treatment group | Related | Total | Related | Total |
|---|---|---|---|---|
| Somnolence | 15 (11.7) | 15 (11.7) | 33 (25.2) | 35 (26.7) |
| Weight increased | 11 (8.6) | 11 (8.6) | 18 (13.7) | 20 (15.3) |
| Dizziness | 8 (6.3) | 12 (9.4) | 14 (10.7) | 19 (14.5) |
| Nasopharyngitis | 0 (0.0) | 21 (16.4) | 0 (0.0) | 17 (13.0) |
| Fatigue | 15 (11.7) | 17 (13.3) | 11 (8.4) | 15 (11.5) |
| Dry mouth | 9 (7.0) | 10 (7.8) | 12 (9.2) | 14 (10.7) |
| Headache | 7 (5.5) | 13 (10.2) | 3 (2.3) | 12 (9.2) |
| Increased appetite | 5 (3.9) | 5 (3.9) | 12 (9.2) | 12 (9.2) |
| Back pain | 0 (0.0) | 8 (6.3) | 0 (0.0) | 8 (6.1) |
| Hypertension | 0 (0.0) | 3 (2.3) | 1 (0.8) | 11 (8.4) |
| Urinary tract infection | 0 (0.0) | 2 (1.6) | 1 (0.8) | 10 (7.6) |
| Diarrhea | 1 (0.8) | 7 (5.5) | 4 (3.1) | 9 (6.9) |
| Arthritis | 0 (0.0) | 6 (4.7) | 1 (0.8) | 9 (6.9) |
| Restless legs syndrome | 3 (2.3) | 3 (2.3) | 6 (4.6) | 8 (6.1) |
| Influenza | 0 (0.0) | 4 (3.1) | 0 (0.0) | 5 (3.8) |
| Nightmare | 3 (2.3) | 3 (2.3) | 5 (3.8) | 6 (4.6) |
| Nausea | 1 (0.8) | 3 (2.3) | 3 (2.3) | 6 (4.6) |
| Pain in extremity | 0 (0.0) | 2 (1.6) | 1 (0.8) | 6 (4.6) |
| Upper respiratory tract infection | 0 (0.0) | 3 (2.3) | 0 (0.0) | 4 (3.1) |
| Insomnia | 0 (0.0) | 4 (3.1) | 1 (0.8) | 4 (3.1) |
| Oropharyngeal pain | 0 (0.0) | 3 (2.3) | 0 (0.0) | 4 (3.1) |
| Sinusitis | 0 (0.0) | 4 (3.1) | 0 (0.0) | 3 (2.3) |
| Constipation | 1 (0.8) | 2 (1.6) | 4 (3.1) | 4 (3.1) |
| Hepatic enzyme increased | 0 (0.0) | 0 (0.0) | 2 (1.5) | 4 (3.1) |
| Cough | 1 (0.8) | 8 (6.3) | 0 (0.0) | 3 (2.3) |
| Sedation | 4 (3.1) | 4 (3.1) | 2 (1.5) | 2 (1.5) |
| Abnormal dreams | 6 (4.7) | 6 (4.7) | 1 (0.8) | 1 (0.8) |
| Upper abdominal pain | 2 (1.6) | 4 (3.1) | 0 (0.0) | 1 (0.8) |

One participant had an SAE 2 days after the 30-day follow-up period; this participant is not counted in the table. ¹irrespective of time point of death, ²including the fatal SAE, ³relationship specified by the investigator as "Definite," "Probable," "Possible." AE: adverse event, SAE: serious AE.
Instead, improvement was noted with an increase from baseline for alertness at awakening, level of energy, and ability to work/function. At end of treatment, the alertness score at awakening had increased from baseline by a median of 17 and 15 points for the 1.5-mg and 3.0-mg groups, respectively (based on LOCF data). Energy level score increased by a median of 13 (1.5-mg) and 11 (3.0-mg) points from baseline, and ability to work/function by 12 points (both dose groups). There was no significant change in nap time at end of treatment compared with baseline in either dose group.

Efficacy

Exploratory efficacy measures were determined by comparison of subjective sleep parameters (TST, WASO, and SL) recorded on Day 1 with those recorded at end of treatment (Table 4, Supplementary Table 2 in the online-only Data Supplement). Mean changes from baseline in weekly TST, WASO, and SL based on the primary analysis (using LOCF) and sensitivity analysis (based on OC data) are shown in Fig. 3 and Supplementary Fig. 1 (in the online-only Data Supplement), respectively, and median changes from baseline are described in the text. Results were consistent in both the OC and LOCF analyses for all sleep efficacy measures evaluated; therefore, the LOCF data are described below while the corresponding OC data are presented in the online-only Data Supplement.

**Table 3.** Summary statistics for residual effects parameters (all participants-treated population; last observation carried forward)

| Statistic                      | Change from baseline to end of treatment |
|--------------------------------|------------------------------------------|
|                                | Esmirtazapine 1.5 mg (n=128) | Esmirtazapine 3.0 mg (n=131) |
| Alertness at awakening         |                                |                                |
| n                              | 123 | 128 |
| Median (range)                 | 17 (-71 to 97) | 15 (-46 to 88) |
| Mean (SD)                      | 18.3 (29.2) | 16.2 (24.9) |
| p-value                        | <0.0001 | <0.0001 |
| Feeling full of energy the past 7 days |                                |                                |
| n                              | 123 | 128 |
| Median (range)                 | 13 (-87 to 95) | 11 (-58 to 83) |
| Mean (SD)                      | 16.1 (27.7) | 13.6 (24.3) |
| p-value                        | <0.0001 | <0.0001 |
| VAS rating on ability to work/function |                                |                                |
| n                              | 123 | 128 |
| Median (range)                 | 12 (-77 to 94) | 12 (-50 to 88) |
| Mean (SD)                      | 14.7 (26.7) | 14.6 (23.1) |
| p-value                        | <0.0001 | <0.0001 |
| Total nap time (min)           |                                |                                |
| n                              | 25  | 25  |
| Median (range)                 | -1 (-45.0 to 40.0) | 5 (-120.0 to 30.0) |
| Mean (SD)                      | -2.8 (16.7) | -0.9 (27.4) |
| p-value                        | 0.4174 | 0.8737 |

SD: standard deviation, VAS: visual analog scale

**Table 4.** Summary statistics of SL, WASO, and TST at baseline and end of treatment (intent-to-treat population; last observation carried forward)

| Sleep characteristic (min) | Time point | Esmirtazapine 1.5 mg (n=128) | Esmirtazapine 3.0 mg (n=129) |
|---------------------------|------------|-------------------------------|-------------------------------|
|                           |            | Mean (SD)                     | Mean (SD)                     |
|                           |            | Median (range)                | Median (range)                |
| Sleep duration            |            |                                |                                |
| TST                       | n          | 127                           | 127                           |
|                           | Baseline   | 294.0 (71.0)                 | 293.5 (67.3)                 |
|                           | n          | 124                           | 124                           |
|                           | End of treatment | 381.0 (82.5)  | 380.4 (99.1) |
|                           |            | 300.0 (150.0 to 666.0)        | 300.0 (0.0 to 436.0)          |
| Sleep maintenance         |            |                                |                                |
| WASO                      | n          | 127                           | 127                           |
|                           | Baseline   | 109.4 (86.4)                 | 102.4 (71.0)                 |
|                           | n          | 124                           | 124                           |
|                           | End of treatment | 47.5 (53.9)  | 52.1 (67.4) |
|                           |            | 30.0 (0.0 to 360.0)           | 25.0 (0.0 to 360.0)           |
| Sleep induction           |            |                                |                                |
| SL                        | n          | 127                           | 127                           |
|                           | Baseline   | 88.6 (72.5)                  | 91.4 (77.7)                  |
|                           | n          | 124                           | 124                           |
|                           | End of treatment | 60.8 (81.4)  | 69.3 (85.7) |
|                           |            | 30.0 (5.0 to 445.0)           | 30.0 (3.0 to 450.0)           |

SD: standard deviation, SL: sleep latency, TST: total sleep time, WASO: wake time after sleep onset
At baseline, the median TST per night was 300.0 min for the 1.5-mg and 3.0-mg dose groups (Table 4). The TST improved during the 52-week treatment period for both groups, and at the end of treatment, both groups reported a TST time of 390.0 min. The median change in TST from baseline to end of treatment was 90.0 min in both dose groups. The median subjective WASO at baseline was 90.0 min for both dose groups. There was an overall decline from baseline in WASO with both doses of esmirtazapine, and this decrease was more pronounced in the 1.5-mg group than the 3.0-mg group. At end of treatment, the median WASO was 30.0 min and 25.0 min for participants receiving esmirtazapine 1.5 mg and 3.0 mg, respectively. The median decrease from baseline in WASO at end of treatment was 50.0 min and 38.0 min for the 1.5-mg and 3.0-mg groups, respectively. The median SL per night tended to decrease over the study, from 60 min and 70 min at baseline for participants receiving esmirtazapine 1.5 mg and 3.0 mg, respectively, to 30 min in both dose groups at end of treatment. The median decrease from baseline was 25.0 min and 17.5 min in the 1.5-mg and 3.0-mg groups, respectively.

The median ISI total score at baseline was 18 for both the 1.5-mg and 3.0-mg esmirtazapine treatment groups. An improvement in ISI score from baseline to endpoint was observed for both treatment groups. At Week 52 (Visit 18), the median change from baseline in ISI was -9 and -11 for the 1.5-mg and 3.0-mg treatment groups, respectively; at end of treatment, the median change from baseline in ISI was -8 for both treatment groups. Changes from baseline in ISI total score by visit/week are shown in Table 5 (OC data). Similarly, a significant improvement in IGR-S was observed at end of treatment relative to baseline in both treatment groups, indicating a lower severity of insomnia. The median change from baseline to end of treatment for IGR-S was -2 in both treatment groups. The response rate, defined as the percentage of participants with “marked” or “moderate” improvement, increased over time in both treatment groups, from 46.8% at Week 2 to 59.8% at end of study for participants receiving the 1.5-mg dose, and from 61.7% to 64.3%, respectively, for participants receiving the 3.0-mg dose.

Overall, the magnitude of change in the subjective efficacy variables was generally consistent throughout the 52 weeks of treatment, and no apparent difference between the 1.5-mg and 3.0-mg treatment groups was observed.

**Discussion**

The current study examined the long-term safety of esmirtazapine in elderly patients with a diagnosis of chronic insomnia. The efficacy assessments were exploratory in nature, as no placebo comparator was included. The results indicate reasonable tolerability of both investigated doses of esmirtazapine (1.5 mg and 3.0 mg) administered before bedtime for the long-term treatment of elderly patients with insomnia. The exploratory efficacy data demonstrated a marked improvement from baseline in subjective sleep diary measures (i.e., TST, WASO, SL). Similarly, improvements from baseline were also noted in the ISI and IGR-S. These improvements from base-
line were sustained over the 52-week treatment period and did not exhibit apparent differences between the two dose groups. The percentage of participants completing the current study (59.1%) was comparable with the completion rate (65.4%) of a 1-year, open-label, extension phase of two randomized trials of zaleplon.35

The safety of 1.5-mg, 3.0-mg, and 4.5-mg doses of esmirtazapine was previously examined for a shorter duration of treatment and in a younger population of adult patients with insomnia (aged ≤65 years).31,32,34 In these studies, esmirtazapine was well tolerated with minimal residual effects. The incidence of AEs and SAEs was higher in the current study relative to previously published studies of esmirtazapine treatment in non-elderly patients with primary insomnia.31,32 In a 2-week study in non-elderly patients, 25.5% and 32.8% of patients had AEs, and 1.5% and 0.8% had SAEs, with esmirtazapine 1.5 mg and 3.0 mg, respectively. In a 6-week study in non-elderly patients, 35.3% of patients had AEs following esmirtazapine 3.0 mg administration. No SAEs were reported with esmirtazapine 3.0-mg in the 6-week study. However, cross-study comparisons should be interpreted with caution due to the differences in study design and participant population. Consistent with previous findings from 2- and 6-week studies of esmirtazapine treatment in non-elderly patients, the current study results indicate that somnolence was among the most commonly reported AEs. This effect did not seem to interfere with overall daily functioning as assessed by the weekly diary, given the sustained improvements in scores of alertness at awakening, energy level, and the ability to work/function throughout the treatment period as compared with baseline. Increase in body weight was also a frequently recorded AE (8.6% and 15.3% in the 1.5-mg and 3.0-mg groups, respectively). This is not surprising, given that this side effect has been previously associated with mirtazapine treatment for depression36,37 and was attributed to its mechanism of action, which is based on the blockade of H1, and 5-HT2, and 5-HT3 receptors.29

The fall rate observed in this study was relatively low: three participants (2.3%) receiving a dose of 1.5 mg esmirtazapine reported a fall. While treatment of insomnia symptoms in elderly patients frequently leads to the prescription of GABA_A receptor modulators,30 this population is more sensitive to the AEs of such therapeutic agents: it is reported that one in seven elderly patients treated with benzodiazepines will have an AE.30 One study reported that 40% of elderly patients admitted for emergency care after a fall had been prescribed benzodiazepines and Z-drugs.40 Furthermore, compared with other times, it is reported that the fall risk for nursing-home residents is greater (odds ratio 3.79; 95% confidence interval 1.10–13.00) within 24 h of benzodiazepine administration.41 Increased caution when prescribing Z-drugs for long-term treatment of insomnia in the geriatric population is, therefore, recommended.42 The relatively low fall rate observed for esmirtazapine in

### Table 5. Change from baseline for ISI total score by visit (intent-to-treat population: observed cases)

| Statistic | Change in total ISI score from baseline |
|-----------|----------------------------------------|
|           | Esmirtazapine 1.5 mg (n=128) | Esmirtazapine 3.0 mg (n=129) |
| Visit 5 (week 2) | | |
| n         | 118 | 117 |
| Median (range) | -4 (-21 to 6) | -5 (-20 to 7) |
| Mean (SD) | -4.7 (5.0) | -5.6 (5.4) |
| p-value | <0.0001 | <0.0001 |
| Visit 6 (week 4) | | |
| n         | 110 | 112 |
| Median (range) | -7 (-17 to 4) | -7 (-21 to 6) |
| Mean (SD) | -6.5 (4.9) | -7.0 (5.7) |
| p-value | <0.0001 | <0.0001 |
| Visit 8 (week 12) | | |
| n         | 110 | 100 |
| Median (range) | -7 (-19 to 5) | -9 (-22 to 6) |
| Mean (SD) | -7.7 (6.0) | -8.6 (5.9) |
| p-value | <0.0001 | <0.0001 |
| Visit 11 (week 24) | | |
| n         | 96 | 91 |
| Median (range) | -9 (-21 to 4) | -9 (-25 to 7) |
| Mean (SD) | -8.5 (5.9) | -8.8 (6.4) |
| p-value | <0.0001 | <0.0001 |
| Visit 14 (week 36) | | |
| n         | 81 | 77 |
| Median (range) | -10 (-20 to 2) | -9 (-22 to 4) |
| Mean (SD) | -9.4 (5.5) | -8.9 (5.9) |
| p-value | <0.0001 | <0.0001 |
| Visit 18 (week 52) | | |
| n         | 70 | 68 |
| Median (range) | -9 (-22 to 6) | -11 (-21 to 4) |
| Mean (SD) | -8.9 (6.4) | -10.3 (5.8) |
| p-value | <0.0001 | <0.0001 |
| Endpoint | | |
| n         | 122 | 123 |
| Median (range) | -8 (-22 to 6) | -8 (-25 to 7) |
| Mean (SD) | -7.5 (6.6) | -7.5 (6.9) |
| p-value | <0.0001 | <0.0001 |

*p-value: performed for change from baseline within treatment based on t-statistics, baseline: the last observation before study treatment. ISI: insomnia severity index, SD: standard deviation
the current study is of interest; however, study limitations should be taken into consideration.

Discontinuation due to insufficient therapeutic effect appeared to occur more frequently in the 3.0-mg group than the 1.5-mg group. Although the small sample size limits the ability to draw conclusions regarding this finding, and a definitive explanation is therefore unclear, greater residual effects at the higher dose, which may have diminished participants’ interpretations of efficacy, may have contributed. This may be supported by the observation that approximately twice the number of participants in the 3.0-mg group reported somnolence and dizziness compared with the 1.5-mg group. Lack of compliance with taking medication per protocol could be another contributing factor; however, cumulative compliance in this study was 95.8% for the 3.0-mg group and 97.6% for the 1.5-mg group, and is unlikely to fully explain the difference.

The observation that both doses of esmirtazapine showed no apparent deleterious effects on measures of alertness at awakening, energy level, and ability to work/function (as captured with the sleep diary) is consistent with data from previous studies that have assessed the potential for next-day hangover effects from esmirtazapine in non-elderly adults with insomnia.31,32 Following a 2-week treatment with esmirtazapine (1.5 mg, 3.0 mg, and 4.5 mg), there was no evidence of residual effects in the esmirtazapine groups.31 A 6-week treatment with esmirtazapine also demonstrated minimal residual next-day effect, as evaluated by a digit symbol substitution test, VAS score for morning alertness, and the Bond-Lader rating scale.32 In another study that evaluated effects of an acute, 2-day treatment of esmirtazapine in patients with insomnia, a decrease in energy level and ability to work/function was experienced after the first night compared with placebo; however, this effect was not reported after the second night.34 In the current study, both treatment groups showed improvement from baseline in alertness at awakening and energy levels, as assessed by the weekly diary, and both esmirtazapine doses provided similar median changes from baseline for both parameters. Sedation has been reported for mirtazapine,44 although no evidence of next-day sedation was observed among patients with major depressive disorder treated with mirtazapine (45 mg) as compared with those treated with fluoxetine (A Winokur, personal communication). A study assessing the effects of esmirtazapine 1.5 mg and 4.5 mg on next-day driving performance showed that single and repeated administration of esmirtazapine 1.5 mg had no clinically meaningful effects on driving performance. With a single dose of esmirtazapine 4.5 mg, driving impairment was observed; however, this resolved following repeated dosing.35 A lack of prominent morning sedation observed with esmirtazapine in this study may be of interest to psychiatrists.

Previous reports on the efficacy of esmirtazapine in non-elderly adult patients with primary insomnia have indicated sustained beneficial effects on objective and subjective patient-reported measures of sleep onset and maintenance over the study period.31,32,34 However, the examined treatment durations did not exceed 6 weeks, which does not reflect the long-term course of therapy that might be required when treating patients with chronic insomnia. The results of the present study suggest that esmirtazapine has enduring sleep-promoting effects in elderly patients over a period of 52 weeks, and that these effects did not appear to be accompanied by the development of tolerance. This is supported by the marked improvements in TST, WASO, SL, ISI, and IGR-S values compared with baseline at each of the assessments conducted throughout the study.

The apparent sleep-promoting properties of esmirtazapine should be considered in the context of long-term efficacy data of other pharmacological treatments for insomnia. While benzodiazepines have established short-term benefits with regard to improvement in sleep initiation and maintenance, their long-term efficacy has not been established.26 As such, they are recommended for short-term use, with continued use believed to be associated with tolerance.26 Long-term use of non-benzodiazepines has been more rigorously tested in clinical trials. For example, in a randomized, double-blind, placebo-controlled study, nighttime use of eszopiclone in patients (aged 21–69 years) with chronic insomnia demonstrated efficacy over a 6-month period.44 Similar to the sustained improvements in TST, WASO, and SL over the year-long period of the current study, the authors reported that the median SL and WASO decreased from baseline by 30 min and 39 min, respectively, and median TST increased from baseline by 83 min after 6 months.44 Similarly, zolpidem extended-release (12.5 mg) was well tolerated in non-elderly adults (aged 18–64 years) and improved sleep onset and maintenance parameters compared with placebo in a 6-month randomized, double-blind, placebo-controlled, multicenter study.45 In elderly patients (aged 59–95 years) with insomnia, nightly zaleplon (5 mg and 10 mg) was associated with sustained improvements in sleep onset, sleep duration, and number of awakenings in a year-long, open-label extension study.45 However, it should be noted that relatively few studies have reported the systematic monitoring of TST, WASO, and SL parameters over a comparably long time period, and this, combined with differences in trial design, limits the ability to draw direct comparisons with other therapeutics. The current study highlights the need for systematic assessment of long-term treatment effects of drugs for insomnia.

Several limitations should be considered when interpreting the findings of this study. The study design did not include a non-active (placebo) treatment arm and participants were only blinded to the dose of treatment they received, which limits ef-
ficacy evaluation. The contribution of factors other than drug activity to the frequency of AEs and reported therapeutic effect cannot, therefore, be excluded. Also, the sample size of the study was not determined by power analysis for detecting a significant difference at a fixed, Type I error rate. As a result, there was no formal hypothesis formulation and testing in this study. The study was only designed to assess within-group statistical differences relative to baseline values—not differences between the dose groups. Although there were no apparent differences between the two dose groups with regard to exploratory efficacy outcomes, dose-response measurements were not collected for this study, so only a tentative statement regarding the lack of apparent dose-response effects of esmirtazapine in improving insomnia sleep parameters long-term can be made. Finally, co-occurrence of insomnia with anxiety and/or depression is prevalent in the elderly population, and patients with such comorbidities were excluded from this study. This limits the generalizability of the findings and warrants further research efforts in patients with comorbid insomnia to establish the safety and efficacy of esmirtazapine in these subpopulations.

Our findings suggest that esmirtazapine doses of 1.5 mg and 3.0 mg nightly are reasonably tolerated in elderly outpatients with insomnia as long-term treatment (up to 1 year) without evidence of significant residual effects. Despite the lack of placebo and the limited power of the analysis, the persistent improvements in sleep parameters throughout this year-long study in elderly patients are encouraging, especially when considering tolerance issues associated with the long-term use of many hypnotics.

Supplementary Materials

The online-only Data Supplement is available with this article at https://doi.org/10.13078/jsm.190032.

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Conflicts of Interest

NI-M, QC, and AP are current or former employees of Merck Sharp & Dohme Corp., a subsidiary of Merck & Co., Inc., Kenilworth, NJ, USA, and may own stock and/or stock options in Merck & Co., Inc., Kenilworth, NJ, USA. AW received research funding and personal fees from Merck Sharp & Dohme Corp., a subsidiary of Merck & Co., Inc., Kenilworth, NJ, USA for the conduct of the study at UConn Health, Farmington, CT, USA.

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Supplementary Table 1. Summary statistics for residual effects parameters (all-participants-treated population; observed case)

| Statistic                              | Change from baseline to end of treatment |
|----------------------------------------|------------------------------------------|
|                                        | Esmirtazapine 1.5 mg (n=128) | Esmirtazapine 3.0 mg (n=131) |
| Alertness at awakening                 |                                          |
| n                                      | 123                                      | 128                                      |
| Median (range)                         | 17 (-71 to 97)                          | 15 (-46 to 88)                          |
| Mean (SD)                              | 18.3 (29.2)                             | 16.2 (24.9)                             |
| p-value                                | <0.0001                                  | <0.0001                                  |
| Feeling full of energy the past 7 days |                                          |
| n                                      | 123                                      | 128                                      |
| Median (range)                         | 13 (-87 to 95)                          | 11 (-58 to 83)                          |
| Mean (SD)                              | 16.1 (27.7)                             | 13.6 (24.3)                             |
| p-value                                | <0.0001                                  | <0.0001                                  |
| VAS rating on ability to work/function |                                          |
| n                                      | 123                                      | 128                                      |
| Median (range)                         | 12 (-77 to 94)                          | 12 (-50 to 88)                          |
| Mean (SD)                              | 14.7 (26.7)                             | 14.6 (23.1)                             |
| p-value                                | <0.0001                                  | <0.0001                                  |
| Total nap time (min)                   |                                          |
| n                                      | 25                                       | 25                                       |
| Median (range)                         | -1 (-45.0 to 40.0)                      | 5 (-120.0 to 30.0)                      |
| Mean (SD)                              | -2.8 (16.7)                             | -0.9 (27.4)                             |
| p-value                                | 0.4174                                   | 0.8737                                   |

SD: standard deviation, VAS: visual analog scale
### Supplementary Table 2. Summary statistics of SL, WASO, number of enquiries, and TST at baseline and end of treatment (intent-to-treat population; observed case)

| Sleep characteristic | Time point | Esmirtazapine 1.5 mg (n=128) | Esmirtazapine 3.0 mg (n=129) |
|----------------------|------------|-----------------------------|-----------------------------|
|                      |            | Mean (SD) | Median (range) | Mean (SD) | Median (range) |
| Sleep duration       |            |           |               |           |               |
| TST                  | Baseline   | 294.0 (71.0) | 300.0 (150.0 to 666.0) | 293.5 (67.3) | 300.0 (0.0 to 436.0) |
|                      | End of treatment | 381.0 (82.5) | 390.0 (120.0 to 510.0) | 380.4 (99.1) | 390.0 (6.0 to 660.0) |
| Sleep maintenance    | WASO       | 109.4 (86.4) | 90.0 (0.0 to 660.0) | 102.4 (71.0) | 90.0 (13.0 to 540.0) |
|                      | End of treatment | 47.5 (53.9) | 30.0 (0.0 to 360.0) | 52.1 (67.4) | 25.0 (0.0 to 360.0) |
| Sleep induction      | SL         | 88.6 (72.5) | 60.0 (9.0 to 480.0) | 91.4 (77.7) | 70.0 (4.0 to 36.0) |
|                      | End of treatment | 60.8 (81.4) | 30.0 (5.0 to 445.0) | 69.3 (85.7) | 30.0 (3.0 to 450.0) |

SD: standard deviation, SL: sleep latency, TST: total sleep time, WASO: wake time after sleep onset
Supplementary Figure 1. Mean change from baseline for weekly TST (A), WASO (B), and SL (C) in the intent-to-treat population (observed case). SL: sleep latency, TST: total sleep time, WASO: wake time after sleep onset.