Indiplon in the management of insomnia

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Abstract: Indiplon is a novel pyrazolopyrimidine, nonbenzodiazepine γ-aminobutyric acid (GABA) agonist studied for the treatment of insomnia. This article reviews the chemistry, pharmacology, clinical pharmacokinetics, drug interactions, clinical trials, safety, tolerability, contraindications, use in special populations, and dosing of indiplon. OVID, International Pharmaceutical Abstracts (IPA), and PubMed databases were searched (1966 to February 2009) for the keywords indiplon, NBI-34060, and insomnia. References of key articles were also reviewed to identify additional publications. Only English language articles were selected for review. Indiplon has been shown to have high affinity and selectivity for the GABAA₂₁ receptor subunit associated with sedation. In clinical studies, indiplon has demonstrated efficacy in improving latency to sleep onset, latency to persistent sleep, total sleep time, wake time after sleep onset, number of awakenings after sleep onset, and overall sleep quality when compared to placebo. Indiplon has a favorable safety profile with limited rebound insomnia and no tolerance. Neurocrine Biosciences, Incorporated received an Approvable Letter from the United States Food and Drug Administration in December 2007 for the indiplon IR 5 mg and 10 mg capsules based on meeting three additional requirements. At the time of this writing, indiplon remains unapproved.

Keywords: indiplon, insomnia, NBI-34060

Introduction

The diagnostic criteria for primary insomnia, from the Diagnostic and Statistical Manual of Mental Disorders fourth edition, text revision (DSM-IV-TR), includes difficulty falling asleep, intermittent wakefulness during sleep and/or nonrestorative sleep (poor quality sleep) for at least one month.¹ From the 2005 Sleep in America Poll, the symptoms reported at least a few nights a week with insomnia were waking up feeling unrefreshed (38%), awake a lot at night (32%), difficulty falling asleep (21%), and woke up too early and could not get back to sleep (21%).² Clinically, patients can be impacted in their social and occupational functioning. Daytime consequences such as fatigue, decreased attention, decreased energy, along with lack of motivation, and deteriorated mood may occur. Approximately one-third of adult Americans experience symptoms of insomnia intermittently, while 10%–15% report having chronic insomnia.³,⁴

Treatment options for insomnia include cognitive behavioral therapy and/or pharmacologic therapy.⁵,⁷ The pharmacologic classes prescribed for insomnia include benzodiazepines, nonbenzodiazepine sedative hypnotics, melatonin receptor agonists, antidepressants with sedating properties, and sedating antihistamines. Indiplon is a...
nonbenzodiazepine sedative hypnotic with more selectivity for γ-aminobutyric acid (GABA) receptors with the α₁ subunit.9

Neurocrine Biosciences, Incorporated (Inc) (San Diego, CA, USA) has researched immediate (IR) and modified release (MR) formulations of indiplon. On May 15, 2006, the US Food and Drug Administration (FDA) granted an Approvable Letter for indiplon IR 5 mg and 10 mg capsules.10 The FDA also requested a reanalysis of certain study data to support the indications of sleep initiation and middle of the night dosing, a reanalysis of safety data for the elderly, and a supplemental pharmacokinetic/food effect trial. A New Drug Application for indiplon IR was resubmitted to the FDA on June 12, 2007. Indiplon IR received an Approvable Letter from the FDA in December 2007 for the 5 mg and 10 mg dosage forms based on meeting three additional requirements. The FDA requested a study in elderly patients, a study comparing adverse events of indiplon versus an approved product for the treatment of insomnia, and a preclinical study to evaluate its use in the third trimester of pregnancy. According to the Neurocrine Biosciences, Inc 2008 fourth quarter report, a meeting with the FDA occurred in July 2008 and the company is awaiting a final version of those minutes to determine the next step with indiplon.11 Currently indiplon is not reported as being approved for use in any other countries.12

Indiplon MR 15 mg tablets received a Not Approvable Letter in May 2006.10 The not approvable status involved indiplon MR doses >15 mg/d and insufficient sleep maintenance data. The FDA requested a long-term safety and efficacy trial for adults and the development of a separate MR dose for the elderly. At this time, the MR 15 mg dose has not been resubmitted to the FDA.

This review will focus on the chemistry, pharmacology, clinical pharmacokinetics, drug interactions, clinical trials, adverse reactions, contraindications, use in special populations, and dosing of indiplon in the treatment of primary insomnia.

Chemistry and pharmacology

The GABA_A receptor is a pentameric molecule composed of a combination of one or more specific subunit types. Although 19 different subunits are known to exist,12 the majority of GABA_A receptors in the central nervous system consist of α₁–α₃, β₁–β₃ and γ₁–γ₃ subunits.13 The interaction of benzodiazepines with multiple GABA_A receptor subunits containing α₁–α₃ subunits is thought to elicit the variety of effects seen with these agents such as anxiolysis, amnesia, muscle relaxation, sedation, and anticonvulsant activity.13,14 The α₁-receptor subunit has been associated with inducing sedation.13,14 The theoretical advantage of having a selective α₁ subunit agonist is that sedating effects are achieved while avoiding other effects thought to be mediated by the other α subunits to which benzodiazepines bind.

In contrast to benzodiazepines, the nonbenzodiazepine sedative hypnotics (ie, zolpidem, eszopiclone, zopiclone, zaleplon) are more selective for the GABA_A receptors with the α₁-receptor subunit.6 Indiplon, a new pyrazolopyrimidine sedative hypnotic, has been shown to have high affinity and selectivity for the α₁ subunit associated with sedation.8,9 Petroski and colleagues15 showed indiplon to be at least nine times more selective for α₁ compared to α₂, α₃, and α₅ subunits.15 This study further showed that the degree of selectivity for α₁ over the α₂ and α₃ subunits, was greater for indiplon as compared to zolpidem, zopiclone, and zaleplon. The α₂ subunit is involved in producing anxiolytic effects.13 The 9.3 times lower affinity of indiplon to the α₂ vs α₁ subunit suggests it does not have significant anxiolytic properties.15

Continual use of benzodiazepines such as triazolam has been associated with the development of tolerance during intermediate and long-term use.16 Although tolerance has been reported less with agents such as zolpidem,16 it remains important to evaluate these parameters for agents used for inducing sleep. Jochelson and colleagues17 studied 30 healthy males, aged 22–41, to identify pharmacokinetic or pharmacodynamic tolerance associated with indiplon. Study subjects were prescribed indiplon 10 mg, 30 mg, or 45 mg once daily for 14 consecutive days. The pharmacokinetic and pharmacodynamic analysis showed similar results on day 1 as compared to day 14 concluding no tolerance was associated with indiplon. Moscovitch and colleagues18 also reported no evidence of tolerance in a six-month elderly study. The mean number of indiplon IR doses throughout the study was 22 per month. Indiplon’s high selectivity for the α₁ subunit, as well as a short half-life may theoretically contribute to its lack of tolerance.15

Clinical pharmacokinetics

Indiplon is extensively metabolized in the liver to two major inactive metabolites.19 The CYP3A4/5 enzyme primarily accounts for the formation of N-desmethyl-indiplon (60%–70%) while carboxylesterase enzymes form N-desacetyl indiplon (30%–40%). A secondary inactive metabolite, N-desmethyl-N-desacetyl-indiplon, is produced
by the N-demethylation of N-desacetyl-indiplon as well as the N-deacetylation of N-desmethyl-indiplon. Less than 1% of indiplon is eliminated in the urine or feces as unchanged drug.

Animal data show indiplon attains a peak plasma level (Tmax) in 30 minutes and has a half-life of 60 minutes. A human trial by Rogowski and colleagues administered indiplon IR 15 mg orally to all subjects and revealed the Tmax in males was 0.73 hours with a half-life of 1.97 hours. In female subjects, the Tmax was 0.82 hours with a half-life of 1.71 hours. Although this trial included only 24 subjects, no statistically significant gender differences were found in regards to indiplon’s pharmacokinetics. Jochelson and colleagues evaluated 12 adults aged 18–45 and compared them to 13 elderly subjects aged 65–79. After one oral dose of indiplon IR 15 mg, the mean Tmax in the younger adult population was 2.3 hours compared to 2.7 hours in the elderly group. The mean half-life of indiplon was 1.5 hours in the young adults and 1.8 hours in the elderly group. Statistical evaluation of this data determined no significant differences exist between the young and elderly populations. This study included similar numbers of males and females in each group and also determined no differences in gender exist in regards to indiplon pharmacokinetics. The carboxylesterase metabolic pathway, which is less affected by age and not known to have gender differences, is one of the proposed reasons for similar pharmacokinetics seen across these populations.

A trial by Bozigian and colleagues further evaluated the pharmacokinetics of indiplon 10 mg, 15 mg, 20 mg, and 30 mg daily in 24 young healthy males. A dose-dependent linear increase in area under the curve (AUC) as well as a proportional dose-dependent increase in maximum concentration was evident. The Tmax was about one hour for all doses and the half-life varied from 2.59–3.82 hours.

Drug interactions

Ketoconazole, troleandomycin, and erythromycin are inhibitors of the CYP3A4 enzyme and have been shown to decrease the metabolism of indiplon. Ketoconazole increased the plasma AUC by 2.4-fold while erythromycin increased indiplon’s plasma AUC by 1.25-fold. The alternative route of carboxylesterase metabolism may explain why the change in plasma AUC due to a highly potent inhibitor like ketoconazole is limited to a 2.4-fold increase. Indiplon was also studied in conjunction with sertraline and paroxetine and no changes in indiplon pharmacokinetics were seen, nor were there any pharmacodynamic changes in regards to tests of psychomotor function and alertness.

Concomitant use of rifampin, a CYP3A4 inducer, has been shown to reduce the plasma AUC of indiplon by 70%.

Indiplon is a weak inhibitor of CYP450 enzymes, and is not expected to cause any clinically significant inhibition of other medications. Abel and colleagues studied indiplon use in conjunction with aminophylline, digoxin, and warfarin and revealed indiplon did not affect the pharmacokinetics of any of the drugs studied. Indiplon does not induce the CYP1A2 or CYP3A4 enzymes and thus is not expected to cause any significant interaction through enzyme induction. Interactions with the carboxylesterase metabolism pathway are not expected, as no currently marketed drugs are known to inhibit these enzymes.

Berkowitz and colleagues studied the interaction of ethanol with indiplon. Ten male subjects were randomized to indiplon 10 mg, ethanol 0.7 mg/ml, or both. Indiplon’s peak plasma level at one hour and half-life were not affected by ethanol. The Digital Symbol Substitution Test (DSST) and Symbol Copying Test (SCT) were utilized to evaluate performance. DSST involves measuring cognitive function by giving a subject 90 seconds to use a nine digit code key and substitute the correct symbol for each digit shown on the test. A lower score on the DSST indicates more impairment. The SCT assesses the motor component of the DSST by asking the subject to copy the same numbers used in the DSST. Based on the DSST and SCT, a slight decrease in performance was evident when indiplon and ethanol were combined, however no effect on sedation or reaction time was seen. Different metabolic pathways easily explain the lack of pharmacokinetic interaction, however the minimal pharmacodynamic interaction is not as well defined. The lack of interaction of indiplon with the α₁ subunit has been proposed as a possible reason for the lack of a pharmacodynamic interaction with ethanol.

Clinical studies

Indiplon IR efficacy and safety was evaluated in 10 randomized, double-blind, placebo controlled trials and one open label extension trial. The efficacy of indiplon MR was evaluated in four randomized, double-blind, placebo controlled trials. Trial data is summarized in Table 1 for indiplon IR in adults, Table 2 for indiplon IR in the elderly, and Table 3 for indiplon MR. Table 4 provides a summary of the clinical efficacy terms used in the indiplon studies. It should be noted that Neurocrine Biosciences, Inc sponsored all the studies and many of the studies are only available in abstract form.
Immediate release indiplon

Adult studies

Efficacy of indiplon IR was evaluated in patients meeting DSM-IV criteria for primary insomnia with nocturnal awakenings. Patients received indiplon IR 10 mg, 20 mg, or placebo. This was a post-bedtime dose study and most patients took the medication between the hours of 12:00 AM and 2:00 AM. The mean number of doses taken per week was 3.9 ± 1.9, 3.7 ± 1.7, and 3.7 ± 1.9 for the 10 mg, 20 mg, and placebo groups, respectively. All study outcomes were subjectively measured. The primary outcome was patientrated latency to sleep onset (LSO) post-dose in which the four-week average was decreased to mean ± standard error of mean (SEM) values of 36.5 ± 1.8 minutes (p = 0.0023) in the 10 mg group, 34.4 ± 1.7 minutes (p < 0.0001) in the 20 mg group, and 45.2 ± 2.2 minutes in the placebo group. Average total sleep time (TST) improved significantly at 253 ± 6.8 minutes in the 10 mg group (p = 0.0099) and 278 ± 6.5 minutes in the 20 mg group (p < 0.0001), compared to 229 ± 6.6 minutes in the placebo group. Baseline values for the wake time after sleep onset (WASO) were 27.5 ± 4.6 minutes for the placebo group, 31.7 ± 4.7 minutes for the 10 mg group, and 37.7 ± 4.6 minutes for the 20 mg group. Only in the 20 mg group was the four-week average WASO significant with values of 24.4 ± 3.5, 16.2 ± 3.6, and 12 ± 3.5 minutes (p = 0.0122) in the placebo, 10 mg, and 20 mg groups, respectively. The number of awakenings after sleep-onset (NAASO) was significantly reduced only in the 20 mg group (p = 0.0125). Overall sleep quality was improved in both indiplon treatment groups compared to placebo (10 mg, p = 0.0007 and 20 mg, p < 0.0001). Next day residual effects as rated on the Visual Analog Scale for sleepiness (VAS-S) were significantly better for both the 10 mg and 20 mg groups as indicated by a 10 mm or greater improvement in the VAS-S for next day alertness compared to placebo (p < 0.05). VAS-S is a scale where study subjects rate their level of alertness or sleepiness on a VAS which ranges from very alert to very sleepy.

Walsh and colleagues examined indiplon IR 10 mg or 20 mg versus placebo over five weeks in patients with primary insomnia of at least three months duration. Results from week 1 for polysomnography (PSG) measuring latency to persistent sleep (LPS) demonstrated significant improvement in the 10 mg and 20 mg groups versus placebo. The values for LPS at week 1 were 27.7 minutes for the 10 mg group (p < 0.01) and 27.1 minutes for the 20 mg group (p < 0.05), in contrast to 36.9 minutes for the placebo group. At week 5 of the study, LPS values remained significant at 29.2 minutes (p < 0.01) for the 10 mg group and 24.8 minutes (p < 0.05) for the 20 mg group, versus 40.1 minutes in the placebo group. TST (no values reported) was significantly better than placebo only during week 1 in the 10 mg group (p < 0.02). At week 1, the LSO was significantly reduced compared to placebo with values of 49.8 (p < 0.0001), 44.4 (p < 0.003), and 72.5 minutes in the indiplon 10 mg, 20 mg, and placebo groups, respectively. The LSO remained significantly better than placebo in both active treatments at week 5 (p < 0.02; no values reported). Sleep quality was only significantly better during week 1 in the 10 mg and 20 mg groups versus

### Table 1 Clinical efficacy trials of indiplon IR in adults

<table>
<thead>
<tr>
<th>Author (year)</th>
<th>N</th>
<th>Age range in years (Mean)</th>
<th>Duration</th>
<th>Dose (mg)</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Farber (2008)</td>
<td>35</td>
<td>18–45 (32.4)</td>
<td>One night per arm</td>
<td>indiplon 10, 20 placebo zolpidem 10 zopiclone 7.5</td>
<td>No change in DSST, SCT, VAS-S for indiplon at 4 and 6 hours post-dose. ↑ VAS-S for zolpidem at 4 hours. ↑ VAS-S for zopiclone at 4 and 6 hours. No change in DSST and SCT for zolpidem and zopiclone at 4 and 6 hours.</td>
</tr>
<tr>
<td>Roth (2007)</td>
<td>264</td>
<td>18–64 (46)</td>
<td>Four weeks</td>
<td>indiplon 10, 20 placebo</td>
<td>↓ LSO, ↓ swASO (20 mg only), ↓ sNAASO (20 mg only), ↑ TST, ↑ sleep quality</td>
</tr>
<tr>
<td>Walsh (2004)</td>
<td>194</td>
<td>Range NR (40.2)</td>
<td>Five weeks</td>
<td>indiplon 10, 20 placebo</td>
<td>↓ LPS, ↓ LSO, ↑ TST (10 mg, week 1 only), ↑ sleep quality (week 1 only)</td>
</tr>
<tr>
<td>Rosenberg (2007)</td>
<td>593</td>
<td>21–64 (32)</td>
<td>One night</td>
<td>indiplon 10, 20 placebo</td>
<td>↓ LPS, ↓ TST, ↓ NAASO (20 mg only), ↓ WASO (20 mg only), ↑ sleep quality</td>
</tr>
<tr>
<td>Roth (2003)</td>
<td>228</td>
<td>18–59 (NR)</td>
<td>One night</td>
<td>indiplon 15, 30 placebo</td>
<td>↓ LPS, ↓ LSO, No change TST</td>
</tr>
<tr>
<td>Scharf (2007)</td>
<td>702</td>
<td>21–64 (46)</td>
<td>Three months</td>
<td>indiplon 10, 20 placebo</td>
<td>↓ LSO, ↑ TST, ↓ swASO, ↓ sNAASO, ↑ sleep quality</td>
</tr>
</tbody>
</table>

Notes: *Abstract-only data; solution; designated as primary endpoint.*

**Abbreviations:** DSST, digit symbol substitution test; LSO, latency to sleep onset; LPS, latency to persistent sleep; NAASO, number of awakenings after sleep onset; NR, not reported; s, subjective or self; SCT, symbol copying test; TST, total sleep time; VAS-S, visual analog scale of sleepiness; WASO, wake after sleep onset.
placebo (p < 0.03; no values reported). No difference in next day residual effects was noted between the placebo and indiplon groups.

The efficacy of indiplon was also examined in two studies using a transient model of insomnia. The first study evaluated the efficacy of indiplon IR 10 mg and 20 mg doses in healthy volunteers using an experimental model of transient insomnia induced by sleeping in an unfamiliar place and a two-hour earlier bedtime. LPS values as measured by PSG were significantly decreased in the 10 mg group to 21.2 ± 1.5 minutes and in the 20 mg group to 16.8 ± 1.1 minutes, compared to the placebo group at 33.1 ± 2.5 minutes (p < 0.0001). TST as measured by PSG was significantly increased in both the 10 mg and 20 mg groups compared to placebo with mean values of 414.5 ± 3.9 minutes (p = 0.0044), 423.5 ± 3.1 minutes (p < 0.0001), and 402.9 ± 3.9 minutes, respectively. NAASO as measured by PSG was significantly decreased only in the 20 mg group and the median values were 7 in the 10 mg group (p = 0.0583), 6 in the 20 mg group (p = 0.0084), and 8 in the placebo group. WASO as measured by PSG was significantly decreased only in the 20 mg group with mean values of 49.9 ± 2.9 in the placebo group, 48.7 ± 3.5 (p = 0.2540) in the 10 mg group, and 42.5 ± 2.8 minutes (p = 0.0091) in the 20 mg group. Subjective sleep quality was significantly improved in both the 10 mg (p = 0.0182) and 20 mg groups compared to placebo (p < 0.0001). No differences in next day residual effects were noted in either treatment group versus placebo. The second study which used the experimental transient insomnia regimen included healthy volunteers using indiplon 15 mg or 30 mg solution, versus placebo. PSG measured LPS was significantly decreased with mean values of 17.5 minutes in the 15 mg group and 16.2 minutes in the 30 mg group, compared to 34.1 minutes for placebo (p < 0.001). Significant improvement in subjective LSO was also noted with mean values of 15.8, 15.4, and 31.1 minutes for the 15 mg, 30 mg, and placebo groups, respectively (p < 0.001). No improvement in the PSG, TST, or subjectively measured TST was found in the indiplon groups versus placebo. Evaluation of next day effects found no difference between the indiplon and placebo groups.

Only one study has examined the long-term efficacy of indiplon. Scharf and colleagues evaluated indiplon IR 10 mg, 20 mg, or placebo in patients who met DSM-IV criteria for primary insomnia of at least three months duration. In the three-month study, all of the primary and secondary outcomes were subjectively measured through patient sleep diaries. At one month, mean LSO was significantly improved to 34 ± 1.3 minutes for 10 mg (p < 0.0001) and 33.0 ± 1.3 minutes for 20 mg (p < 0.0001), compared to 48.7 ± 1.9 minutes for placebo. Significant efficacy was maintained for all three months for LSO in the active treatment groups compared to placebo. Mean TST was 327.5 ± 4 minutes at one month in the placebo group compared to 362.8 ± 3.9 minutes (p < 0.0001) in the 10 mg group, and 372.1 ± 4 minutes (p < 0.0001) in the 20 mg group, and efficacy was statistically maintained for three months. WASO was 61.1 ± 2.5, 49.7 ± 2.4 (p = 0.0009), and 42.2 ± 2.5 minutes (p < 0.0001) in the placebo, 10 mg, and 20 mg groups, respectively at one month and this efficacy was maintained for the three-month duration.
study duration. Significant improvement in mean NAASO was noted in the active treatment groups compared to placebo at one month with mean values of $1.6 \pm 0.1$, $1.3 \pm 0.1$ ($p < 0.0001$), and $1.1 \pm 0.1$ ($p < 0.0001$) in the placebo, 10 mg, and 20 mg groups, respectively and significance was demonstrated for all three months. Sleep quality was significantly improved for all three months in the active treatment groups as well.

To date no published trials have directly compared the efficacy of indiplon to a FDA-approved agent for sleep. However, one trial examined next day effects using a five-way crossover study in which adult patients received a single post-bedtime dose (four hours after lights-out) of indiplon IR 10 mg, indiplon IR 20 mg, zolpidem 10 mg, zopiclone 7.5 mg, and placebo. The primary outcome was to evaluate next day effects of the medications compared to placebo using the VAS-S, DSST, and the SCT. No significant differences in the VAS-S, DSST, or SCT were noted four or six hours post dose in the indiplon 10 mg or 20 mg groups. Baseline VAS-S values were $47.4 \pm 3$ in the zolpidem 10 mg group, $41.6 \pm 3$ in the zopiclone 7.5 mg group, and $48.1 \pm 3$ in the placebo group. Significant differences compared to placebo were noted in the VAS-S at four hours post-dose for zolpidem ($60.5 \pm 3$; $p = 0.042$) and zopiclone ($66.1 \pm 3$; $p < 0.0001$) with this persisting to six hours post-dose in the zopiclone group ($57.2 \pm 3$; $p = 0.002$).

**Elderly studies**

Indiplon IR was specifically studied in elderly patients in four trials. Walsh and colleagues conducted a two-week study involving patients who met criteria for primary insomnia for at least three months comparing indiplon IR 5 mg or 10 mg to placebo. End points were self-recorded in patient diaries. The mean LSO significantly improved at week 1 with a value of $34.6 \pm 1.8$ minutes ($p < 0.0001$) in the 5 mg group and of $30.4 \pm 1.6$ minutes ($p < 0.0001$) in the 10 mg group, compared to placebo with a value of $47.7 \pm 2.5$ minutes. This significant difference in mean LSO was maintained at week 2 for both the 5 mg ($p = 0.0160$) and 10 mg ($p = 0.0028$) doses. Mean TST at week 1 was $312.2 \pm 5$ minutes in the placebo group, $340.3 \pm 5$ minutes ($p < 0.0001$) in the 5 mg group, and $360 \pm 5$ minutes ($p < 0.0001$) in the 10 mg group. TST remained significantly better at week 2 compared to placebo for the 5 mg ($p = 0.0064$) and 10 mg ($p = 0.0015$) doses. Significant improvement was also noted at week 1 for NAASO with values of $2 \pm 0.1$, $1.6 \pm 0.1$ ($p = 0.0002$), $1.4 \pm 0.1$ ($p < 0.0001$) in the placebo, 5 mg, and 10 mg groups, respectively. At week 2, NAASO remained significant with values of $1.9 \pm 0.1$, $1.6 \pm 0.1$ ($p = 0.0014$), $1.5 \pm 0.1$ ($p < 0.0001$) in the placebo, 5 mg, and 10 mg groups, respectively. The WASO was significant in the indiplon 10 mg group at week 1 with $60.5 \pm 4$ minutes ($p < 0.0001$) and week 2 with $69.8 \pm 4$ minutes ($p = 0.0112$) compared to $83.3 \pm 4.1$ minutes at week 1 and $84.2 \pm 4$ minutes at week 2 in the placebo group. WASO was significant only in week 2 in the 5 mg group with values at week 1 of $73.1 \pm 4$ minutes ($p = 0.0725$) and week 2 with $70.8 \pm 4$ minutes ($p = 0.0180$). Sleep quality was also significantly improved in the 5 mg group at week 1 ($p < 0.0001$) and week 2 ($p = 0.0007$) as well as the 10 mg group at week 1 ($p < 0.0001$) and week 2 ($p = 0.005$), compared to placebo.

The longest indiplon IR trial in the elderly was a six-month open-label extension trial. Patients with a three-month or more history of primary insomnia self-evaluated the benefit of indiplon on their insomnia. Patients received either indiplon IR 5 mg or 10 mg as needed at bedtime. Only 81 of 121 patients completed six months of the study, with 72% of patients on the 5 mg dose and 92% of patients on the 10 mg dose reporting indiplon IR improved their insomnia.
The efficacy of indiplon IR was also examined in a four-period crossover study in elderly patients with a diagnosis of primary insomnia comparing indiplon IR 5 mg, 10 mg, or 20 mg to placebo.\(^{33}\) PSG measured mean LPS was significantly improved with values of 13.8, 10.4, 9.8, and 25.2 minutes in the indiplon IR 5 mg, 10 mg, 20 mg, and placebo groups, respectively (p < 0.001). Significant improvement in PSG measured mean TST was only demonstrated in the indiplon IR 10 mg group with a value of 372.1 minutes (p = 0.027) and in the 20 mg group with a value of 385.6 (p < 0.001) minutes. TST values for the 5 mg and placebo groups were 363.7 and 354.4 minutes, respectively. Subjectively measured mean LSO was significantly improved with values of 28.8 minutes in the 5 mg group, 24.7 minutes in the 10 mg group, 20.2 minutes in the 20 mg group, and 41.8 minutes in the placebo group (p < 0.004). No difference in next day residual effects as measured by DSST, SCT, and VAS-S was found in the indiplon groups compared to placebo.

Farber and colleagues\(^{26}\) studied indiplon IR in an indirect manner with an approved comparative agent in elderly patients to evaluate next day residual effects. In a one-night, four-way crossover, single post-bedtime dose (four hours after lights-out) trial, indiplon IR 5 mg and 10 mg were compared to placebo and zopiclone 3.75 mg as an active control. The primary outcome measured next day residual effects using the VAS-S, DSST, and the SCT. No significant difference in VAS-S or SCT was found in the medication arms versus placebo. Also, no significant difference was noted in the indiplon IR 5 mg group versus the placebo group in the DSST at four, six, or eight hours post-dose. However, the indiplon IR 10 mg group did demonstrate a significant difference at four hours post-dose compared to placebo (p = 0.022) in DSST, but no difference was noted at six and eight hours post-dose. Finally, zopiclone demonstrated a significant reduction in DSST at both four hours (p = 0.002) and eight hours (p = 0.003) post-dose compared to placebo.

**Indiplon MR**

Indiplon has also been studied in a MR form. The MR dosage form provides drug initially and then gradually releases medication providing therapeutic drug concentrations for an extended period during the night.\(^{34}\) It should be noted that study data related to indiplon MR is limited and three of the studies are only available in abstract form.\(^{35-37}\)

**Adult studies**

A two-week study compared indiplon MR 30 mg to placebo in adults with a diagnosis of primary insomnia for greater than three months.\(^{35}\) LSO was 30 minutes in the indiplon MR group compared to 37 minutes in the placebo group (p = 0.003) at week 1 and 27 minutes compared to 33 minutes at week 2 (p = 0.0131), respectively. Subjective TST was significantly improved at week 1 with 375 minutes in the indiplon MR group compared to 328 minutes in the placebo group (p < 0.0001) and this continued at week 2 with 367 minutes in the indiplon MR group versus 336 minutes in the placebo group (p = 0.0013). Subjective WASO was significantly improved at 54 minutes compared to 79 minutes at week one (p < 0.0001) and 51 minutes compared to 73 minutes at week 2 (p = 0.003) in the indiplon MR and placebo groups, respectively. Sleep quality and NAASO were also significantly improved at weeks 1 and 2 (p < 0.0001) compared to placebo (p-value not reported).

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**Table 4** Efficacy measures used in indiplon clinical trials\(^{38-40}\)

<table>
<thead>
<tr>
<th>Efficacy term</th>
<th>Measured end point</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Latency to sleep onset (LSO)*</td>
<td>Sleep onset</td>
<td>Time it takes to go from lying in bed to the beginning of sleep</td>
</tr>
<tr>
<td>Latency to persistent sleep (LPS)</td>
<td>Sleep onset</td>
<td>An objective PSG measurement of the time it takes to fall asleep</td>
</tr>
<tr>
<td>Total sleep time (TST)*</td>
<td>Sleep duration</td>
<td>Actual amount of sleep time which includes all stages of sleep</td>
</tr>
<tr>
<td>Wake after sleep onset (WASO)*</td>
<td>Sleep maintenance</td>
<td>Amount of time spent awake after initial sleep onset until the end of the sleep cycle</td>
</tr>
<tr>
<td>Number of Awakenings after sleep onset (NAASO)*</td>
<td>Sleep maintenance</td>
<td>Number of times a subject awakens from sleep during the night</td>
</tr>
</tbody>
</table>

**Note:** *Objective measured by PSG or subjectively measured by patient diary or post-sleep questionnaires.

**Abbreviations:** PSG, polysomnogram; REM, rapid eye movement.

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38. Farber and colleagues
39. Adult studies
40. Drug Design, Development and Therapy downloaded from https://www.dovepress.com/ by 54.70.40.11 on 12-Feb-2019
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Indiplon MR 40 mg was studied in healthy male volunteers compared to placebo in a venipuncture model of insomnia.\textsuperscript{18,26,32,33} Blood samples were taken throughout the night to impair sleep initiation and sleep maintenance. Subjectively measured improvements in LSO were noted in the indiplon MR group at 30.2 minutes versus 72.2 minutes in the placebo group (p < 0.001). Significant improvement in subjective TST was found in the indiplon MR group with 315.6 minutes versus 246.6 minutes in the placebo group (p = 0.02). The NAASO was improved, but not significantly in the indiplon MR group compared to placebo and this lack of significance was believed to be due to the multiple blood draws during the night. Overall patient rated sleep quality was significantly improved compared to placebo (p = 0.04). No change in next day residual effects was noted compared to baseline using the DSST, SCT, and VAS-S assessments.

Elderly studies

Indiplon MR 15 mg was studied in elderly patients with primary insomnia of at least three months duration in a two-week trial.\textsuperscript{34} All data was self-recorded by patients in daily sleep diaries. Significant improvement in mean subjective TST occurred in the indiplon MR group at week 1 with a value of 376 ± 4.2 minutes (p < 0.0001) compared to the placebo group with a value of 327.5 ± 4.4 minutes. LSO was significantly improved in the indiplon group at weeks 1 and 2 with mean values of 22.0 ± 1.1 and 21.2 ± 1.2 minutes (p < 0.0001) compared to the placebo group at 34.9 ± 1.8 and 31 ± 1.8 minutes, respectively. Other significant findings at weeks 1 and 2 in the indiplon MR group included mean WASO with 55.7 ± 3.1 minutes at week 1 (p < 0.001) and 56.4 ± 3.6 minutes at week 2 (p < 0.0001), compared to the placebo group at 84.9 ± 3.2 minutes at week 1 and 80.5 ± 3.7 minutes at week 2. Mean NAASO significantly improved in the indiplon MR group to 1.4 ± 0.1 at week 1 (p < 0.0001) and 1.3 ± 0.1 at week 2 (p < 0.0001) compared to 2 ± 0.1 at week 1 and 1.9 ± 0.1 at week 2 in the placebo group. Sleep quality was also significantly improved in week 1 at 64% (p < 0.0001) compared to 32% and in week 2 at 57% (p < 0.002) compared to 35% in the indiplon MR and placebo groups, respectively. The authors did note that the study design may not completely represent elderly patients with chronic insomnia since patients with unstable medical or psychiatric illnesses were excluded and due to the strict time-in-bed requirements of the study.

Indiplon MR was examined in elderly patients comparing doses of 10 mg, 20 mg, 30 mg, and 35 mg to placebo with each patient receiving each dose of medication on two consecutive nights.\textsuperscript{37} Significant improvement in mean LPS values was found with values of 26 minutes in the placebo group, 17.6 minutes in the 10 mg group, 11.2 minutes in the 20 mg group, 11 minutes in the 30 mg group, and 9.9 minutes in the 35 mg group (p < 0.01). Mean sleep efficiency was significantly improved in the 20 mg, 30 mg, and 35 mg dosages only (p < 0.0001). WASO in the placebo group was 103.2 minutes and 102.1 minutes in the 10 mg group (NS). Significant improvements in WASO occurred only in the 20 mg (87.3 minutes), 30 mg (81.9 minutes), and 35 mg (83 minutes) dosage groups, compared to placebo (103.2 minutes) (p < 0.01).

Safety and tolerability

Adult and elderly adverse events

Indiplon IR

The clinical studies reporting specific adverse events related to indiplon are summarized in Table 5.\textsuperscript{18,26,27,29,31,32,41} Other studies did not report details on adverse events but reported no serious adverse events.\textsuperscript{18,28,30,33} Based on available study data, no specific serious adverse events attributable to indiplon have been identified. Currently it is difficult to determine if there is a dose dependent increase in the incidence of specific adverse events. A 12-month randomized, double-blind, safety and tolerability study using as needed indiplon IR 10 mg or 20 mg in 536 adults with insomnia found indiplon to be well tolerated without any dose dependent side effects.\textsuperscript{41} However, a three-month study by Scharf and colleagues\textsuperscript{11} reported discontinuation rates secondary to adverse events as higher in the indiplon 20 mg group (19.7%) compared to the indiplon 10 mg group (6.4%) and placebo (5.2%). Adverse events in this study were reported as transient, with median durations of 3–19 days for the adverse events occurring at a >5% incidence (see Table 5).

Indiplon’s safety in the elderly was specifically evaluated in four studies,\textsuperscript{18,26,32,33} two of which are only available in abstract form.\textsuperscript{18,33} The largest of these studies evaluated two weeks of nightly use of indiplon IR 5 mg and 10 mg vs placebo in patients aged 65–80 years. A dose-related increase in study discontinuation due to adverse events was reported (8.4% on 10 mg, 5% on 5 mg, and 0.8% on placebo).\textsuperscript{32}

Rebound insomnia

The potential for rebound insomnia following indiplon discontinuation was examined in two adult studies and one elderly study.\textsuperscript{28,31} The first adult study found after three months of consecutive nightly use, rebound insomnia occurred
on the first night in 22.1% of subjects on indiplon 20 mg, 18.9% on indiplon 10 mg, and 8.8% on placebo; however, occurrence was much less on the second night, with rates of 11.4%, 11.1%, and 1.1% of patients, respectively (no p value reported). In the second adult study, rebound insomnia, assessed subjectively and by PSG, was not found with indiplon after five weeks of use. In a two-week elderly study, rebound insomnia was noted the first night after stopping active treatment in 2% of the placebo group, 12.9% of the 5 mg group (p = 0.004), and 8.6% (p < 0.05) of the 10 mg group. These values decreased to 0%, 1.1%, and 1.1% in the placebo, 5 mg, and 10 mg groups, respectively on night 2 after discontinuing therapy (NS).

Withdrawal effects
The potential for indiplon to cause withdrawal effects was assessed in two adult studies and one elderly study using the Benzodiazepine Withdrawal Symptoms Questionnaire (BWSQ). The BWSQ uses a three-point severity scale with 20 questions designed to evaluate withdrawal symptoms from medications that work at GABA receptors. After three months of indiplon use, no discontinuation effects were seen in adults at the end of treatment and one week after stopping therapy. There was also no difference in discontinuation effects compared among the 10 mg and 20 mg groups in a five-week adult trial. Indiplon discontinuation following a two-week elderly study did not cause any withdrawal symptoms either.

Abuse potential
The abuse potential of indiplon was studied using large doses of indiplon (30 mg, 50 mg, and 80 mg) versus placebo and triazolam (0.25 mg, 0.5 mg, and 0.75 mg) in 21 adult patients with a history of recreational substance abuse. Investigators

### Table 5: Clinical studies with reported ADR's with indiplon IR greater than placebo

<table>
<thead>
<tr>
<th>Author (year)</th>
<th>ADRs Incidence (%)</th>
<th>Specific ADR</th>
<th>Placebo</th>
<th>Indiplon 5 mg</th>
<th>Indiplon 10 mg</th>
<th>Indiplon 20 mg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Farber (2008)</td>
<td>Placebo 2.8</td>
<td>Headache</td>
<td>0</td>
<td>2.8</td>
<td>2.8</td>
<td>N/A</td>
</tr>
<tr>
<td>Elderly N = 36</td>
<td>Indiplon 5 mg 2.8</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Indiplon 10 mg 2.8</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Farber (2008)</td>
<td>Placebo 8.6</td>
<td>Tiredness</td>
<td>0</td>
<td>N/A</td>
<td>0</td>
<td>2.9</td>
</tr>
<tr>
<td>Adults N = 35</td>
<td>Indiplon 10 mg 5.9</td>
<td>Headache</td>
<td>5.7</td>
<td>5.9</td>
<td>5.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Indiplon 20 mg 11.8</td>
<td>Retarded motor activity</td>
<td>0</td>
<td>0</td>
<td>2.9</td>
<td></td>
</tr>
<tr>
<td>Roth (2007)</td>
<td>Placebo = 26</td>
<td>Headache</td>
<td>2.3</td>
<td>N/A</td>
<td>2.4</td>
<td>6.7</td>
</tr>
<tr>
<td>Adults N = 264</td>
<td>Indiplon 10 mg 39</td>
<td>Somnolence</td>
<td>3.5</td>
<td>6.0</td>
<td>7.8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Indiplon 20 mg 34</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rosenberg (2007)</td>
<td>Placebo = 10</td>
<td>Headache</td>
<td>1</td>
<td>N/A</td>
<td>2</td>
<td>3.1</td>
</tr>
<tr>
<td>Adults N = 593</td>
<td>Indiplon 10 mg 10.1</td>
<td>Nausea</td>
<td>0.5</td>
<td>0.5</td>
<td>2.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Indiplon 20 mg 15.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Scharf (2007)</td>
<td>Not reported</td>
<td>URI</td>
<td>5.2</td>
<td>N/A</td>
<td>5.9</td>
<td>6</td>
</tr>
<tr>
<td>Adults N = 702</td>
<td>Amnesia</td>
<td></td>
<td>0</td>
<td>1.3</td>
<td>6.4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Dizziness</td>
<td></td>
<td>3</td>
<td>4.7</td>
<td>6.9</td>
<td></td>
</tr>
<tr>
<td>Walsh (2007)</td>
<td>Placebo = 42</td>
<td>Headache</td>
<td>2.5</td>
<td>7.5</td>
<td>5.9</td>
<td>N/A</td>
</tr>
<tr>
<td>Elderly N = 358</td>
<td>Indiplon 5 mg = 31.1</td>
<td>Somnolence</td>
<td>0</td>
<td>1.7</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Indiplon 10 mg = 38.3</td>
<td>Depression</td>
<td>0</td>
<td>5</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Decreased appetite</td>
<td>0</td>
<td>5.9</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Dizziness</td>
<td>1.7</td>
<td>4.2</td>
<td>6.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Nausea</td>
<td>0</td>
<td>5</td>
<td>1.7</td>
<td></td>
</tr>
<tr>
<td>Black (2006)</td>
<td>Not reported</td>
<td>Headache</td>
<td>11.8</td>
<td>8.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adults N = 536</td>
<td>Back pain</td>
<td></td>
<td>7.9</td>
<td>N/A</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Somnolence</td>
<td></td>
<td>7.9</td>
<td>N/A</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>URI</td>
<td></td>
<td>N/A</td>
<td>6.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nasopharyngitis</td>
<td></td>
<td>N/A</td>
<td>5.9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: Abstract-only data.
Abbreviations: ADR, adverse drug reaction; URI, upper respiratory infection; N/A, not applicable.
concluded that the abuse potential of indiplon at these high doses was not different when compared to triazolam; however, psychomotor and cognitive impairment might be less with indiplon versus triazolam.

**Daytime residual effects**
Studies evaluating daily and as needed use for up to 12 months in adults, and up to six months in the elderly, have found indiplon IR to be relatively safe,\(^\text{18,41}\) without complaints of next day residual effects.\(^\text{27,28,31,33}\) Details of these effects were previously described in the clinical trials section of this article.

**Indiplon MR safety**
The longest and largest study to date for the MR formulation investigated the use of indiplon MR 15 mg for two weeks in elderly patients.\(^\text{44}\) Forty-four percent of patients in the indiplon MR group had an adverse reaction compared to 28% in the placebo group, with <1% deemed severe. The following adverse events occurred at an incidence two-fold greater than placebo: dizziness (8%), headache (6%), and somnolence (4%). No falls were reported. There was evidence of rebound insomnia on the first night after discontinuation, but incidence returned to baseline by the second night. No evidence of withdrawal symptoms was found using the BWSQ. Safety data from the three other trials looking at the MR formulation are only available in abstract form but report no serious adverse reactions.\(^\text{35–37}\)

**Dosing and administration**
If granted FDA approval, it is anticipated that indiplon IR will be available in 5 mg and 10 mg strengths. It should be noted that efficacy studies in adults have used indiplon IR doses of 10 mg and 20 mg while efficacy studies in the elderly have used indiplon IR doses of 5 mg, 10 mg, and 20 mg.\(^\text{18,32,33}\) Studies have assessed daily nighttime dosing, as needed dosing, and as needed post-bedtime dosing.

**Contraindications, warnings, and precautions**
Although labeling is not currently available, indiplon may not be an option for all patients. As with any medication, indiplon should be contraindicated in anyone with previous hypersensitivity to this medication. In 2007 the FDA recommended a label change for sleep-inducing agents. This label change provided warnings regarding anaphylaxis, angioedema, and other “complex sleep-related behaviors which may include sleep-driving, making phone calls and preparing and eating food (while asleep)”\(^\text{44}\). If approved, indiplon will likely receive the same warning. Because of its hepatic metabolism, use should be avoided in those with impaired liver function. Alcohol use >5 drinks/day or >14 drinks/week was excluded in most trials; however, one study showed there was little or no interaction when indiplon IR 10 mg was co-administered with ethanol in 10 healthy patients.\(^\text{25}\) Because indiplon acts centrally, concomitant alcohol use with indiplon should probably be avoided. Furthermore, most trials excluded concomitant use of centrally acting medications (anxiolytics, anticonvulsants, narcotics, antihistamines). More studies are needed to determine if there would be any difference in the effectiveness or safety of indiplon when used with other centrally acting medications.

Benzodiazepine-like withdrawal symptoms were not reported after discontinuation of indiplon during clinical trials,\(^\text{28,31,32}\) however use should probably be avoided in those with previous dependence on other centrally acting medications, such as benzodiazepines or other hypnotics. One study in a small number of patients with a history of substance abuse found that the potential for abuse of large doses of indiplon was not different when compared with triazolam.\(^\text{45}\)

**Special populations**
Indiplon IR and MR have been studied in patients aged 18–80 years with lower doses in the elderly population. Neither the IR nor MR formulations have been studied in those <18 years of age or in pregnant women, therefore avoiding use of indiplon in these populations is warranted until more safety and efficacy data is available. Two of the three requirements identified in the 2007 FDA Approvable Letter included an objective/subjective clinical trial in the elderly and a pre-clinical study for use during the third trimester of pregnancy.\(^\text{10}\)

**Discussion**
Indiplon may have several advantages to offer if approved for the treatment of primary insomnia. Indiplon IR appears safe and efficacious in the treatment of insomnia in both adult and elderly populations as demonstrated by improvements in LPS, LSO, TST, WASO, NAASO, and sleep quality. A potentially unique item to indiplon is post-bedtime dosing or middle of the night dosing.\(^\text{27}\) However, zaleplon has been used off-label in a post-bedtime or middle of the night dosing method similar to indiplon.\(^\text{49}\) Indiplon appears to be well tolerated with no specific serious adverse events identified during clinical trials. However, the FDA has
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Indiplon in the management of insomnia

requested a study comparing adverse events of indiplon versus an FDA approved product for the treatment of primary insomnia.10

A potential issue related to indiplon in the treatment of primary insomnia is that the majority of adult studies used the 10 mg and 20 mg dosages and the elderly studies used 5 mg and 10 mg dosages. FDA approval was sought only for the 5 mg and 10 mg dosages. It is unclear if this will affect the potential efficacy in the adult population by seeking approval for only one of the dosages studied in adults. Based on available data, it does not appear the 5 mg dose was studied in adults. Also, it should be noted that at various time points in clinical studies, indiplon was unable to demonstrate statistical improvement in values such as TST, WASO, NAASO, and sleep quality depending on the dose of medication and the duration of the trial. Even though indiplon was studied specifically in the elderly, the FDA did request an additional safety and efficacy trial. Indiplon studies also excluded patients with current psychiatric illnesses and patients with acute medical illnesses which are two very common areas of use for sedative hypnotics. At the time of this writing, it is not known what specific indications indiplon will have or if it will have a specific advantage or indication over other products already approved for the treatment of primary insomnia. The post-bedtime or middle of the night dosing advantage of indiplon may be offset by the off label use of zaleplon in such a dosing method. Also, it is not known if indiplon will be indicated for sleep maintenance based on its short half-life. The MR formulation was investigated for that indication, however, indiplon MR received a Not Approvable Letter in 2006 and it does not appear that its approval was pursued following that letter. From the current Approvable Letter it appears that sleep initiation or as needed post-bedtime dosing would be the primary indications for indiplon. Finally, the cost of indiplon is unknown and it is difficult to predict how insurance companies would view indiplon compared to other nonbenzodiazepine sedative hypnotics, some of which are available generically.

Conclusion

Indiplon IR appears safe and effective in the treatment of insomnia in both adult and elderly populations. Indiplon appears to be well tolerated with few side effects and no specific serious adverse events reported in clinical trials. Studies evaluating daily and as needed indiplon IR dosing did not reveal complaints of next day residual or withdrawal effects. The abuse potential of indiplon appears minimal, and it has a low, transient incidence of rebound insomnia. If FDA approved, indiplon may provide a unique dosing option for the treatment of insomnia related to post-bedtime or middle of the night dosing.

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Disclosure

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