Effect of Nonergot Dopamine Agonists on Symptoms of Restless Legs Syndrome

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ABSTRACT

PURPOSE We performed a meta-analysis of randomized placebo-controlled trials of nonergot dopamine agonists (NEDAs) for the treatment of restless legs syndrome.

METHODS A systematic literature search was conducted through July 2007. The primary outcome measures assessed were the percentage of responders to medication as determined by the Clinical Global Impression-Improvement (CGI-I) scale and the adjusted mean change in the International Restless Legs Syndrome Study Group Scale (IRLS) score from baseline compared with placebo. Meta-regression analysis was performed to evaluate the impact of study duration on the primary outcomes. Safety endpoints were also evaluated.

RESULTS A total of 14 trials (n = 3,197 subjects) were included in the meta-analysis. NEDA use resulted in greater response as measured by the CGI-I scale (relative risk [RR] 1.36; 95% CI, 1.24 to 1.49; \( P < .001 \)), and greater reductions in IRLS scores (weighted mean difference [WMD] –4.93; 95% CI, –6.42 to –3.43; \( P < .001 \)) from baseline vs placebo. Meta-regression analysis showed an inverse relationship between study duration and reduction in IRLS score. NEDAs were associated with a significant risk of adverse events (including nausea, dizziness, somnolence, and fatigue.)

CONCLUSIONS Use of NEDAs in patients with moderate-to-severe restless legs syndrome results in significant reductions in symptom severity, but a significant portion of patients will discontinue their use as a result of adverse events.

Ann Fam Med 2008;6:253-262. DOI: 10.1370/afm.845.

INTRODUCTION

Restless legs syndrome is a sensorimotor disorder affecting approximately 12% of the adult population.1,2 Although women and the elderly have a higher prevalence of restless legs syndrome, there is conflicting evidence of the impact ethnicity has on prevalence.2-5 Restless legs syndrome is characterized by an irresistible urge to move the legs, which may begin or worsen during periods of rest or inactivity and often affects sleep.6 Physical activity, such as walking or stretching, often relieve these urges.

The Medical Advisory Board of the Restless Legs Syndrome Foundation developed a treatment algorithm for restless legs syndrome in 2004.7 For patients with daily symptoms, treatment options included dopamine agonists, anticonvulsants (eg, gabapentin), and opioids.7

Dopamine agonists, particularly nonergot dopamine agonists (NEDAs), have become the mainstay of therapy for patients with daily symptoms of restless legs syndrome. NEDAs are generally preferred to ergot dopamine agonists (eg, pergolide, cabergoline), which have been associated with clinically important heart valve damage and resultant regurgitation.8-10

Numerous clinical trials have evaluated the efficacy and safety of NEDAs for restless legs syndrome with conflicting results. Most of these studies lacked adequate power to detect potential NEDA benefits and...
risks. We therefore performed a meta-analysis and meta-regression analysis to evaluate the effect of NEDAs on efficacy, withdrawal resulting from adverse effects, and overall risk of adverse effects in patients with restless legs syndrome.

METHODS

Study Selection

Included trials had to (1) be randomized trials of a NEDA, (2) be placebo controlled, and (3) report data on either the percentage of responders to medication (defined as “much improved” or “very much improved”) as determined by the Clinical Global Impression-Improvement (CGI-I) scale or the adjusted mean change on either the percentage of responders to medication NEDA, (2) be placebo controlled, and (3) report data

Statistical Analysis

Incidence of patient response, as measured by the CGI-I scale, was treated as a dichotomous variable. Weighted averages were reported as relative risk (RR) with associated 95% confidence intervals (CI). A DerSimonian and Laird random-effects model was used in calculating relative risk and 95% confidence intervals for this and all subsequent analyses. Risk difference (as well as number needed to treat) was calculated for response to the CGI-I scale, whereas the adjusted mean change in IRLS score from baseline was calculated as the difference between the adjusted mean IRLS score in the NEDA and placebo groups and reported as a weighted mean difference (WMD) and its 95% confidence interval. In addition, withdrawals caused by adverse events and incidence of prespecified adverse events were reported from the constituent studies. In cases where there was more than 1 published report on the same population or group of patients, the most recent article was selected, although previous articles were reviewed to supplement missing data where applicable. For studies with both an open-label and double-blind treatment portion, only the double-blind portion was used in this meta-analysis, because unblinded or open-labeled trials may exhibit exaggerated treatment effects.

Validity Assessment

The Jadad scale was calculated by 2 investigators (W.L.B., C.I.C.) and used to assess the methodological quality of included trials. This rating scale uses the following quality assessment criteria—use of and methods for generating randomization, use of and methods for double-blinding, and description of patient withdrawals and dropouts—as these are inherent controls of bias. One point was given for each satisfied criterion. An aggregate score between 0 and 5 was calculated for each included trial (0 = weakest, 5 = strongest), with trials scoring less than 3 deemed to have lower methodological quality.

Data Abstraction

Two investigators (W.L.B., C.I.C.) independently abstracted all data using a standardized data abstraction tool, and disagreements were resolved by a third party (C.M.W.). Data related to each efficacy and safety endpoint were sought from the constituent studies. In cases where there was more than 1 published report on the same population or group of patients, the most recent article was selected, although previous articles were reviewed to supplement missing data where applicable. For studies with both an open-label and double-blind treatment portion, only the double-blind portion was used in this meta-analysis, because unblinded or open-labeled trials may exhibit exaggerated treatment effects.

Tolerability was assessed by evaluating study withdrawal rates that were due to adverse events and incidence of commonly reported adverse events (including headache, nausea, dizziness, somnolence, and fatigue).

Using the prespecified inclusion criteria, we conducted a systematic literature search of MEDLINE, EMBASE, CINAHL, and Web of Science, from the earliest possible date through July 2007, for all relevant articles published in English. We used the following Medical Subject Headings (MeSH) and keywords: “pramipexole,” “ropinirole,” “rotigotine,” “sumanirane,” and “dopamine agonist” in combination with “restless legs syndrome” and “RLS.” Results were limited to trials in humans. We manually searched references from reports of clinical trials or review articles to identify additional relevant studies. In addition, we reviewed the Food and Drug Administration (FDA) Web site (http://www.fda.gov; accessed August 7, 2007) and the European Agency for the Evaluation of Medicinal Products (EMEA) Web site (http://www.emea.eu.int; accessed August 7, 2007), as well as the Web sites for the manufacturers of the following agents for additional study data: pramipexole (Mirapex, Boehringer-Ingelheim, http://www.boehringer-ingelheim.com; accessed August 7, 2007), ropinirole (Requip, Glaxo-Smithkline, http://us.gsk.com; accessed August 7, 2007), rotigotine (Neupro, Schwarz Pharma, http://www.schwarzpharma.com; accessed August 7, 2007), and sumanirone (Pfizer, http://www.pfizer.com; accessed August 7, 2007). Two investigators (W.L.B., C.I.C.) reviewed all potentially relevant articles independently.
studies and ranges from 0% to 100%, with the higher percentage representing a higher likelihood of heterogeneity.\(^{14}\) Whereas categorization of values for \(I^2\) may not be appropriate in all situations, \(I^2\) values of 25%, 50%, and 75% have been regarded as representative of low, medium, and high statistical heterogeneity, respectively. We used visual inspection of funnel plots, Egger's weighted regression statistics, and the trim and fill method to assess for publication bias.\(^{15}\) The trim and fill method uses funnel plot symmetry to estimate the number of “missing” studies and the magnitudes of their effects. It reestimates the overall effect size after imputing potentially missing studies into the meta-analysis to determine whether the results of the original analysis were replicated. Statistics were performed using StatsDirect statistical software, version 2.4.6 (StatsDirect Ltd, Cheshire, England), and MIX statistical software (Bax L, Yu LM, Ikeda N, Tsuruta N, Moons KGM. MIX: Comprehensive Free Software for Meta-analysis of Causal Research Data - Version 1.54. 2006; at http://www.mix-for-meta-analysis.info). \(P < 0.05\) was considered statistically significant, except where otherwise indicated.

To evaluate the effect of heterogeneity between included studies on the meta-analysis’ conclusions, we performed the following subgroup and sensitivity analysis: (1) studies of less than 12 weeks’ follow-up and those with greater than or equal to 12 weeks’ follow-up were analyzed separately; (2) unpublished studies were excluded; (3) studies not using or reporting intention-to-treat principles were excluded; (4) studies with a Jadad score <3 were excluded; and (5) a fixed-effect model (Mantel-Haenszel fixed-effect model)\(^{16}\) was chosen for the primary analyses.

We further evaluated the impact of time on the primary outcomes using random-effects meta-regression estimation via iterative maximum likelihood (REML). Meta-regression analysis is an indirect way to examine the possibility of effect modification by duration of treatment. The logarithmically transformed relative risks in the CGI-I scale and change in IRLS score were examined. Meta-regression analysis was performed with SPSS, version 11.0 (SPSS, Chicago, Illinois).

RESULTS

Study Characteristics

The initial search yielded 1,194 potential literature citations (Figure 1). Of those, 1,150 were excluded through review of abstracts, leaving 44 articles for full publication review. We found 14 studies (\(n = 3,197\) subjects) that conformed to our inclusion criteria (Table 1).\(^{17,30}\) We excluded 30 trials from analysis, including those that had an open-label design,\(^{31-33}\) a cross-over design,\(^{34-38}\) or were withdrawal studies (meaning that the effect of continuing treatment vs withdrawing treatment after a double-blind treatment period was assessed).\(^{39,40}\)

In the included trials, patients received ropinirole in 7 trials (\(n = 1,698\) subjects),\(^{17,19,21,24-26}\) pramipexole in 4 trials (\(n = 825\) subjects),\(^{22,24,27}\) rotigotine in 2 trials (\(n = 404\) subjects),\(^{18,26}\) and sumanirole in 1 trial (\(n = 270\) subjects).\(^{25}\) Thirteen trials provided CGI-I scale responder rates,\(^{17-29}\) and 10 provided mean change in IRLS score from baseline.\(^{17,19,24,27,28,30}\) Enrollment ranged from 41 to 381 patients. The mean patient age ranged from 51 to 76 years, and approximately two-thirds were female. Duration of follow-up ranged from 1 to 12 weeks. All trials enrolled patients with similar disease severity (baseline IRLS scores ranged between 22 and 28).

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**Figure 1. Flow diagram of trial identification, inclusion, and exclusion.**

- 1,194 Citations identified and screened
- 1,069 Citations excluded
  - 112 Not human studies
  - 957 Not clinical trials
- 125 Abstracts retrieved for detailed evaluation
- 81 Abstracts excluded as not relevant
  - 81 No useable endpoint reported
- 44 Full-Text articles retrieved for detailed evaluation
- 30 Articles excluded
  - 25 Not randomized, controlled trial
  - 5 No useable endpoint reported
- 14 Trials included\(^{18-31}\)
  - 13 Reported response to CGI-I\(^{19,30}\)
  - 10 Reported change in IRLS from baseline\(^{18,20,25,28,29,31}\)

IRLS = International Restless Legs Syndrome Study Group Scale score; CGI-I = Clinical Global Impression-Improvement scale.
Quantitative Data Synthesis

Upon meta-analysis, NEDA use resulted in a statistically significantly greater response to therapy, as measured by the CGI-I scale compared with placebo (Figure 2A). This corresponded to a risk difference of 0.18 (95% CI, 0.13-0.23; \( P < .001 \)) and a number needed to treat of 6 (95% CI, 5-8). A statistically significant 5-point reduction in the adjusted mean change in the IRLS score from baseline with the NEDAs compared with placebo was also seen (Figure 2B). Statistical heterogeneity was suggested for the adjusted mean change in the IRLS score (Q statistic, \( P < .001 \), \( I^2 = 69\% \)), although it was nonsignificant for the CGI-I scale (Q statistic, \( P = 0.12 \), \( I^2 = 33\% \)). Review of the forest plots for each (Figure 2) shows that included studies were in general agreement on the positive effects of NEDAs, but not the magnitude of benefit.

On visual inspection of the funnel plots, little asymmetry was noted for either the CGI-I scale or the IRLS score, suggesting a low risk of publication bias (not shown). The addition of theoretical studies using the trim and fill method did not significantly alter

Table 1. Clinical Trial Characteristics

| Reference   | Design | N  | Baseline IRLS Mean + SD | Follow-up (weeks) | Drug/Dose         | Adjusted Mean Difference IRLS Mean + SE | Treatment CGI-I Responder Rate (%) | Jadad Score |
|-------------|--------|----|-------------------------|-------------------|-------------------|----------------------------------------|-----------------------------------|-------------|
| Allen (2004) | P, R, DB, PC | 55 | 25.0 ± 5.0              | 12                | Ropinirole 0.25-4.0 mg/dl | -1.2 ± 2.1 | Ropinirole = 17/32 (53) Placebo = 17/33 (52) | 3 |
| Stiasny-Kolster (2004) | P, R, DB, PC | 63 | 24.4 ± 5.8              | 1                | Ropinirole 0.25-4.0 mg/dl | -3.0 ± 1.1 | Ropinirole = 78/146 (53) Placebo = 6/14 (43) | 4 |
| Walters (2004) | P, R, DB, PC | 284 | 23.6 ± 5.9              | 12               | Ropinirole 0.25-4.0 mg/dl | -2.5 ± 1.1 | Ropinirole = 78/131 (60) Placebo = 56/138 (41) | 5 |
| Bogan (2006)   | P, R, DB, PC | 381 | 22.0 ± 5.0              | 12               | Ropinirole 0.25-4.0 mg/dl | -3.7 ± 0.9 | Ropinirole = 53/135 (39) Placebo = 137/187 (73) | 5 |
| Inoue (2006)   | P, R, DB, PC | 41 | 22.7 ± 4.1              | 6                | Pramipexole 0.125-0.75 mg/dl | -11.5 ± 3.0 | Pramipexole = 16/20 (80) Placebo = 11/21 (52) | 3 |
| Partinen (2006) | P, R, DB, PC | 107 | 22.7 ± 4.1              | 3                | Pramipexole 0.125-0.75 mg/dl | -9.2 ± 1.7 | Pramipexole = 65/66 (78) Placebo = 9/21 (43) | 3 |
| Winkelmann (2006) | P, R, DB, PC | 339 | 23.4 ± 5.1              | 12               | Pramipexole 0.25-0.75 mg/dl | -4.3 ± 1.1 | Pramipexole = 183/254 (72) Placebo = 44/85 (52) | 4 |
| Garcia-Borreguero (2007) | P, R, DB, PC | 270 | 25.4                    | 8                | Sumanirole 0.5-4.0 mg/dl | N/A | Sumanirole = 104/212 (49) Placebo = 26/51 (51) | 3 |
| Oertel (2007)  | P, R, DB, PC | 341 | 28.0 ± 6.3              | 6                | Rotigotine 0.5-4 mg/24 h | N/A | Rotigotine = 212/280 (76) Placebo = 29/53 (55) | 5 |
| RRL100013P     | P, R, DB, PC | 338 | 24.7 ± 5.2              | 6                | Pramipexole 0.125-0.75 mg/dl | -6.6 ± 1.1 | Pramipexole = 141/224 (63) Placebo = 37/114 (32) | 3 |
| ROP101892P     | P, R, DB, PC | 359 | 26.0 ± 4.4              | 12               | Ropinirole 0.5-6.0 mg/dl | -4.1 ± 1.0 | Ropinirole = 124/175 (71) Placebo = 92/184 (50) | 4 |
| SKF101468P     | P, R, DB, PC | 54  | 26.6 ± 5.3              | 7                | Ropinirole 0.25-4.0 mg/dl | -9.9 ± 2.9 | Ropinirole = 76/146 (52) Placebo = 76/146 (52) | 3 |

CGI-I = Clinical Global Impression – Improvement scale; DB = double-blind; IRLS = International Restless Legs Syndrome Study Group scale; N/A = not available; P = prospective; PC = placebo control; R = randomized.
Figure 2. Nonergot dopamine agonist's impact on response to clinical global impression-improvement scale (A) and International Restless Legs Syndrome Study Group Scale score (B).

A Relative Risk Meta-Analysis Plot (Random Effects)

| Study              | Relative Risk (95% CI) |
|--------------------|------------------------|
| Allen, 2004        | 1.03 (0.64 to 1.65)    |
| Stiasny-Kolster, 2004 | 1.62 (0.97 to 3.31)  |
| Trenkwalder, 2004  | 1.32 (1.03 to 1.70)    |
| Walters, 2004      | 1.52 (1.18 to 1.96)    |
| Bogan, 2006        | 1.30 (1.12 to 1.52)    |
| Inoue, 2006        | 1.53 (0.98 to 2.55)    |
| Partinen, 2006     | 1.76 (1.16 to 3.12)    |
| Winkelmann, 2006   | 1.39 (1.14 to 1.76)    |
| Garcia-Borreguero, 2007 | 0.96 (0.73 to 1.34)  |
| Oertel (R), 2007   | 1.38 (1.11 to 1.84)    |
| Oertel (P), 2007   | 1.94 (1.48 to 2.60)    |
| RRL100013          | 1.42 (1.20 to 1.69)    |
| SFK101468          | 1.16 (0.95 to 1.43)    |
| Combined (random)  | 1.36 (1.24 to 1.49)    |

B Summary Meta-Analysis Plot (Random Effects)

| Study              | Weighted Mean Difference (95% CI) |
|--------------------|----------------------------------|
| Allen, 2004        | -1.20 (-5.32 to 2.92)            |
| Trenkwalder, 2004  | -3.01 (-5.17 to -0.85)           |
| Walters, 2004      | -2.50 (-4.66 to -0.34)           |
| Bogan, 2006        | -3.70 (-5.46 to -1.94)           |
| Inoue, 2006        | -11.50 (-17.38 to -5.62)         |
| Partinen, 2006     | -9.20 (-12.53 to -5.87)          |
| Winkelmann, 2006   | -4.30 (-6.46 to -2.14)           |
| Oertel (P), 2007   | -6.60 (-8.76 to -4.44)           |
| RRL100013          | -4.11 (-6.07 to -2.15)           |
| SFK101468          | -9.91 (-15.52 to -4.30)          |
| Combined            | -4.93 (-6.42 to -3.43)           |

CI = confidence interval.

Note: The squares represent individual studies and the size of the square represents the weight given to each study in the meta-analysis. Error bars represent 95% CIs. The diamond represents the combined results. The solid vertical line extending upwards from 0 (CGI) or 1 (IRLS) is the null value.
the study results (subgroup analysis, Figures 3 and 4). Egger’s weighted regression analysis confirmed the low-likelihood of publication bias for both the CGI-I and IRLS measures ($P = .59$ and .10, respectively).

Upon subgroup analysis, pramipexole, ropinirole, and rotigotine use resulted in a greater response than placebo, as measured by CGI-I scale ($P < .01$ for each), whereas sumanireole did not have a significant effect on CGI-I scale scores ($P = .80$) (Figure 3). Both pramipexole and ropinirole showed a statistically significant reduction in the adjusted mean change in the IRLS score from baseline as compared with placebo ($P < .001$ for each), but no study using either rotigotine or sumanireole was eligible for inclusion in the IRLS analysis (Figure 4). In sensitivity analysis the conclusions of the meta-analysis remained robust to methodological changes.

Upon meta-analysis of tolerability endpoints, patients receiving NEDAs were more likely to withdraw because of adverse events than those taking placebo ($RR = 1.35; 95\% CI, 1.00-1.81; number needed to harm = 77, P = .048$) (Table 2). When analyzed separately, patients receiving pramipexole, rotigotine, and sumanireole had no significant increase in withdrawal rates, whereas those receiving ropinirole had significantly higher rates of withdrawals caused by adverse events ($P = .02$). NEDAs as a class were found to significantly increase patients’ risk of nausea, dizziness, somnolence, and fatigue ($P < .05$ for all). A trend toward a higher risk of headaches was noted for all NEDAs ($P = .09$). When each drug was analyzed individually, pramipexole significantly increased nausea risk; ropinirole significantly increased nausea, dizziness, somnolence, and fatigue risk; and sumanireole significantly increased headache risk.

Random-effects meta-regression analysis showed a significant relationship between the study duration and the adjusted mean change in the IRLS score from baseline ($P < .001$) (Figure 5). Studies of a shorter duration showed more robust improvements in symptoms than those with longer durations. Approximately a 0.66-point lesser score reduction was seen for each additional study week. A similar relationship was not seen

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**Figure 3. Subgroup and sensitivity analyses of nonergot dopamine agonists evaluating response to Clinical Global Impression-Improvement scale.**

| Subgroup Analyses | RR (95\% CI) |
|-------------------|--------------|
| All NEDAs         | 1.36 (1.24 to 1.49) |
| Pramipexole       | 1.60 (1.34 to 1.92) |
| Ropinirole        | 1.32 (1.21 to 1.43) |
| Rotigotine        | 1.41 (1.12 to 1.79) |
| Sumanireole       | 0.96 (0.71 to 1.30) |

| Sensitivity Analyses | RR (95\% CI) |
|----------------------|--------------|
| >12 weeks’ duration only | 1.32 (1.22 to 1.43) |
| <12 weeks’ duration only | 1.47 (1.16 to 1.86) |
| Published trials only | 1.38 (1.24 to 1.54) |
| ITT trials only       | 1.35 (1.23 to 1.49) |
| Trim and fill analysis | 1.31 (1.18 to 1.45) |
| Fixed-effects model   | 1.37 (1.28 to 1.47) |

CI = confidence interval; DA = dopamine agonists; ITT = intention to treat; NEDA = nonergot dopamine agonist; RR = relative risk.

Note: The dashed vertical line represents the combined treatment effect for the original analysis.
with response to treatment, according to the CGI-I score ($P = .27$) (Figure 5).

**DISCUSSION**

In this meta-analysis of 14 randomized, controlled trials, patients receiving NEDAs for treatment of restless leg syndrome showed significant improvement in their symptoms and disease severity as evidenced by improvements in CGI-I scale and IRLS scores from baseline compared with placebo. These beneficial effects must be weighed against a statistically significant increase in withdrawals resulting from adverse events, as well as an increased incidence of individual adverse events.

When analyzed separately, all NEDAs, with the exception of sumanirole, significantly improved response to treatment and reduced the IRLS score from baseline. Studies of sumanirole did not seem to find a beneficial effect on the CGI-I scale and did not report IRLS scores in a manner conducive to meta-analysis; however, only 1 study of sumanirole met the criteria for inclusion into our meta-analysis. In contrast, the effects on CGI-I scale and IRLS scores qualitatively seem more robust with pramipexole than the other NEDAs, but

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**Table 2. Adverse Events**

| Group            | W/D due to ADEs RR (95% CI) | Headache RR (95% CI) | Nausea RR (95% CI) | Dizziness RR (95% CI) | Somnolence RR (95% CI) | Fatigue RR (95% CI) |
|------------------|-----------------------------|----------------------|-------------------|-----------------------|------------------------|---------------------|
| Dopamine agonists| 1.35 (1.00-1.81)*            | 1.20 (0.98-1.47)     | 3.25 (2.36-4.48)* | 1.47 (1.02-2.13)*     | 1.94 (1.45-2.61)*      | 1.37 (1.01-1.86)*   |
| Pramipexole      | 1.15 (0.49-2.69)            | 0.99 (0.64-1.54)     | 2.68 (1.51-4.76)* | 1.03 (0.51-2.07)      | 1.55 (0.75-3.20)       | 1.09 (0.64-1.85)    |
| Ropinirole       | 1.49 (1.06-2.10)*           | 1.21 (0.95-1.54)     | 3.95 (2.76-5.66)* | 1.72 (1.13-2.62)*     | 2.03 (1.47-2.80)*      | 1.72 (1.18-2.68)*   |
| Rotigotine       | 0.46 (0.08-2.58)            | 1.34 (0.54-3.33)     | 0.82 (0.16-4.24)  | 0.58 (0.19-1.73)      | N/A                    | 0.78 (0.32-1.92)    |
| Sumanirole       | 1.11 (0.06-19.45)           | 2.68 (1.01-7.11)*    | 4.77 (0.65-34.69) | 4.53 (0.62-33.04)     | N/A                    | N/A                 |

ADEs = adverse drug events; CI = confidence interval; N/A = not available; RR = relative risk; W/D = withdrawal.

* $P < .05$.
we could not determine the direct comparative efficacy between NEDAs in our meta-analysis. The qualitatively greater improvements with pramipexole could potentially be explained by the relatively short duration of pramipexole studies (3 to 6 weeks), because it is during this time the drug’s effects are most prominent.

Meta-regression analysis showed that the beneficial effects of NEDAs, in terms of improvements in IRLS scores from baseline, are most prominent during the first few weeks of therapy ($P < .001$). These effects appear to diminish somewhat in trials evaluating a 12-week treatment period. Results must be interpreted cautiously, however, given that confounders other than study duration may have an impact on effect size. Thus, any relationships that are identified may not be causal. Notably, previously conducted longer-term follow-up extension studies (up to 2 years) have shown that NEDAs maintain their beneficial effects, although augmentation is seen in 33% to 50% of patients.\textsuperscript{39,41,42}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure5}
\caption{Random-effects meta-regression evaluating effect of NEDAs on mean change in IRLS (A) and CGI-I (B) over time.}
\end{figure}

\begin{itemize}
\item NEDAs = nonergot dopamine agonists; IRLS = International Restless Legs Syndrome Study Group Scale; CGI-I = Clinical Global Impression-Improvement; RR = relative risk.
\item Note: The circles represent individual studies, and the area of the circle is proportional to the weight of each study. The dark line represents the regression equation as represented by the following equation: (A) weighted mean difference $= -11.3545 + 0.6574 \times$ study duration; (B) $\ln \text{RR} = 0.5304 - 0.0219 \times$ study duration.
\end{itemize}
relatively short-term nature (up to 12 weeks), the
benefits found in our meta-analysis are representative of
the overall benefit seen in long-term clinical use.

Although not associated with heart valve prob-
lems, as is treatment with ergot dopamine agonists,
the NEDA use is somewhat limited by its high rate of
adverse events. We found significant increases in the
rate of withdrawals caused by adverse events, as well as
increased incidence of nausea, dizziness, somnolence,
and fatigue, with the NEDAs as a class compared with
placebo. When evaluated separately, ropinirole signi-
ficantly increased the number of withdrawals that were
due to adverse effects and significantly increased the risk
of nausea, dizziness, somnolence, and fatigue compared
with placebo. In contrast, pramipexole did not increase
the risk of withdrawals because of adverse effects and
only increased nausea risk compared with placebo.
Although rotigotine did not increase the risk of with-
drawals because of adverse events or of any individual
adverse event, and although sumanireole only increased
the risk of headache, there was low power to identify
adverse effects, as there were only 2 studies for roti-
gotine and 1 study for sumanireole included in the meta-
alysis. A subsequent reevaluation of tolerability will be
needed when these drugs are more rigorously studied.

It is possible that the NEDA agents might differ in
terms of efficacy and safety as a result of pharmacologic
and pharmacokinetic differences. Whereas ropinirole
and sumanireole show activity primarily toward the
D2 receptor and serotonin receptors, pramipexole
predominantly acts on the D1 receptor.41 Rotigotine
acts on D3, D2, and D1 receptors, as well as serotonin
and α2-adrenergic receptors. Sumanireole has a rela-
tively short half-life of 3 to 5 hours, ropinirole and
rotigotine have a modest half-life of approximately 6
hours, and pramipexole has a long half-life that ranges
from 8 to 12 hours.43 The direct link between recep-
tor specificity, half-life, and efficacy and safety has not
been determined, however, and more work is needed.

There are some limitations to this meta-analysis.
First, we included trials that were not published and
available only online, making critical evaluation of their
methods and results challenging. Even so, our main
study results remained robust despite the removal of
these trials in sensitivity analysis. Second, as with any
meta-analysis, the potential for publication bias is a con-
cern. Publication bias is defined as “the tendency on the
parts of investigators, reviewers, and editors to submit or
accept manuscripts for publication based on the direc-
tion or strength of the study findings.”44 Although visual
inspection of our meta-analysis’ funnel plot could not
rule out publication bias, the results of the trim and fill
analysis showed it is unlikely that publication bias signi-
ficantly affected our study results. This conclusion is fur-
ther supported by the nonsignificant Egger’s weighted
regression statistic. Finally, limitations of symptom score
and disease severity scales should be mentioned. A cli-
ically important difference between treatment options
on the IRLS is a point of ongoing debate, which makes
interpretation of the differences in the current meta-
analysis (approximately 3 to 7 points on a 40-point IRLS
score, or a 7.5%-17.5% improvement) difficult to inter-
pret. It should be noted, however, that the benefits of
therapy on the IRLS score were above that derived from
placebo, which has been shown to be robust in several
studies.19,20 Further research correlating differences in
the IRLS score to quality-of-life scores or clinical events
is necessary to be able to say what magnitude of differ-
ence in these scales are clinically significant.

In conclusion, the use of NEDAs (pramipexole,
ropinirole, rotigotine) in patients with moderate-to-
severe restless legs syndrome results in significant
reduction in symptom severity but increases the risk
of withdrawal of therapy as a result of adverse effects.
There may be qualitative differences in both efficacy
and safety between agents, although no definitive
comparisons can be made. Meta-regression analysis
suggests that these agents are most beneficial in the
early stages of treatment. Future studies should be con-
ducted to compare directly the efficacy of individual
NEDAs against each other in patients with restless legs
syndrome, as well as include a longer period follow-up
period to assess the long-term effects of these agents.

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Submitted September 12, 2007; submitted, revised, October 30, 2007;
accepted November 27, 2007.

Key words: Restless legs syndrome; dopamine agonists; nonergot;
meta-analysis

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