Long-term tolerability of once-monthly injectable paliperidone palmitate in subjects with recently diagnosed schizophrenia

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Background: A post hoc analysis from a multiphase trial with open-label transition and maintenance phases, a double-blind relapse prevention phase, and an optional open-label extension examined the long-term tolerability with continuous once-monthly injectable paliperidone palmitate 39, 78, 117, or 156 mg (25, 50, 75, or 100 mg equivalents [mg eq] of paliperidone) in subjects with recently diagnosed (≤5 years; n = 216) versus chronic illness (>5 years; n = 429) schizophrenia.

Methods: Adverse events reported at a ≥2% margin between subgroups were identified. Relative risks (in the recently diagnosed compared with the chronically ill) and 95% confidence intervals (CI) were determined, and CI not including 1 were considered potentially significant.

Results: In both subgroups, the mean monthly dose was 109 mg (69.9 mg eq). Continuous mean exposures were 333.9 ± 271.9 and 308.7 ± 278.3 days in the recently diagnosed and chronic illness subgroups, respectively. Using the criteria outlined in the methods, nasopharyngitis was a potentially significant event reported in more chronically ill than recently diagnosed subjects at months 6, 9, 12, and endpoint (7.2% versus 2.8%; relative risk 0.384; 95% CI 0.163–0.907). Influenza (2.8% versus 0.7%; relative risk 3.9; 95% CI 1.003–15.730) and amenorrhea (3.2% versus 0.9%; relative risk 3.476; 95% CI 1.029–11.744) at endpoint were potentially significant events in more recently diagnosed than chronically ill subjects. Mean weight changes, sedation/somnolence, any extrapyramidal symptom-related or glucose-related events were generally similar between the groups. The mean prolactin level increased in both sexes in both subgroups (changes from baseline of +41.8 ng/mL and +26.5 ng/mL in recently diagnosed and chronic illness females and +12.3 ng/mL and +15.1 ng/mL in recently diagnosed and chronic illness males, respectively), and were higher in females with recently diagnosed illness than in females who were chronically ill (P = 0.0002 at endpoint). Prolactin-related events were reported by 7.9% of recently diagnosed subjects with schizophrenia and 3.5% of those who were chronically ill.

Conclusion: The long-term tolerability of paliperidone palmitate was generally similar in recently diagnosed schizophrenia subjects and those with more chronic illness, with the exception of some prolactin-related measures.

Keywords: paliperidone palmitate, long-acting antipsychotic, recently diagnosed, early illness, schizophrenia

Introduction
In subjects with schizophrenia, the first 5–10 years of illness has been identified as a critical period for effective intervention to prevent biological as well as psychosocial deterioration,1,2–7 optimize outcomes,1,8–12 reduce the risk for symptom relapse,13–17 and possibly mitigate disease progression.5,18 However, during these early years, there are...
some barriers to effective intervention and treatment. These patients are often nonadherent with medications due to lack of insight into both their illness and the need for treatment, as well as forgetfulness, lack of social support, and personal choice.19 Whereas recently diagnosed patients are often more responsive to antipsychotic medications than those with chronic schizophrenia,14,16,20,21 they may also be more sensitive to adverse drug effects.5,22–24 In particular, reports suggest that various extrapyramidal symptoms (EPS), weight gain, prolactin-related effects, and sedation are more frequent and problematic for persons early in their illness.8,29–36

Because early in the course of schizophrenia illness patients poorly adhere to daily therapy, it has been suggested that long-acting injectable antipsychotics may be a particularly appropriate treatment option for these patients. In addition to eliminating the daily need to remember to be adherent, these agents allow clinicians and caregivers to have immediate awareness of noncompliance. Guaranteed onboarding of antipsychotic medication allows clinicians to make more informed treatment decisions. Further, the long half-lives of these agents provide a wider window for missed doses (days or weeks rather than hours) before plasma levels drop below critical thresholds, where the risks for relapse, hospitalization, and suicide may be increased.21,37–41

On the other hand, tolerability is often a key factor in medication choice for recently diagnosed patients and long-acting agents may pose such concerns for both clinicians and patients. Unfortunately, few studies have assessed the safety of injectable medications in patients with a recent onset42–45 or first episode of psychosis,46 and their use in recently diagnosed patients is generally limited. Despite well known adherence issues with oral medications and an associated high risk for subsequent relapse,2,37 long-acting agents are often reserved for more treatment-refractory patients.

Paliperidone palmitate (Janssen Pharmaceuticals, Titusville, NJ), the palmitate ester of paliperidone, is a long-acting, once-monthly, injectable, atypical antipsychotic for the treatment of schizophrenia.47 Doses of paliperidone palmitate may be expressed as milligrams (mg) or as milligram equivalents (mg eq) of the pharmacologically active fraction, paliperidone (39, 78, 117, 156, and 234 mg of paliperidone palmitate corresponding to 25, 50, 75, 100, and 150 mg eq of active paliperidone). Its use for both acute and maintenance treatment of schizophrenia has been demonstrated in short-term48–51 and longer-term52–54 studies. To date, there are no published prospective studies of paliperidone palmitate in patients experiencing their first episode or early in their schizophrenia illness. A recent post hoc analysis of a large, international, 13-week, placebo-controlled trial reported on tolerability and efficacy of paliperidone palmitate in subjects with recently diagnosed schizophrenia receiving the recommended initiation doses (234 mg [150 mg eq] day 1 and 156 mg [100 mg eq] day 8).55 These data are informative but do not address longer-term tolerability concerns. A recently completed relapse prevention trial52,54 of paliperidone palmitate provides valuable long-term exposure data that can help to address this question. A post hoc safety and tolerability analysis of this trial focusing on the recently diagnosed subgroup as compared with the more chronically ill subgroup is reported herein.

Materials and methods

Design

This was a post hoc analysis of a five-phase international trial conducted from March 2005 to February 2008 (NCT001111189). Key subject inclusion criteria included being aged 18–65 years and a body mass index ≥ 15.0 kg/m², a diagnosis of schizophrenia for at least one year before screening, and a total Positive and Negative Syndrome Scale score below 120 at screening and at baseline, with no lower score eligibility limit. The study was conducted in accordance with the Declaration of Helsinki and consistent with Good Clinical Practice. Additional design details have been previously reported.52,54

Study phases and treatments

The five study phases (Figure 1) were: a screening/washout phase up to 7 days; an open-label, 9-week, transition phase for switching to paliperidone palmitate at 78 mg (50 mg eq) on days 1 and 8, followed by flexible dosing of 39, 78, or 156 mg (25, 50, or 100 mg eq) at week 5; an open-label 24-week maintenance phase with flexible paliperidone palmitate dosing (39, 78, or 156 mg monthly [25, 50, or 100 mg eq]) for the first 12 weeks followed by the established maintenance dose for the second 12 weeks; a randomized (1:1 ratio), double-blind, placebo-controlled, relapse prevention phase of variable duration (up to 63 weeks) for subjects stabilized on a fixed paliperidone palmitate dose during the maintenance phase; and an optional open-label extension phase up to 52 weeks with flexible paliperidone palmitate dosing (39, 78, 117, or 156 mg [25, 50, 75, or 100 mg eq]). Subjects who experienced a relapse, completed the relapse prevention phase, or received at least one injection of paliperidone palmitate when enrollment in the study was stopped were eligible to enter the open-label extension phase.
Post hoc population and tolerability assessments

The recently diagnosed subgroup was defined as those subjects within 5 years of their initial diagnosis of schizophrenia. The chronically ill subgroup included those subjects > 5 years out from their initial diagnosis of schizophrenia. This analysis was limited to subjects who received paliperidone palmitate continuously from entry into the open-label transition phase to study completion or discontinuation (Figure 1).

Tolerability was evaluated by treatment-emergent adverse events reported from the open-label transition phase baseline through months 1, 3, 6, 9, and 12 and to the open-label extension endpoint. These included reports of adverse events (either reported by the subject voluntarily or collected by means of interviewing subjects in a nondirected manner), clinical laboratory tests, vital sign measurements, extrapyramidal symptom rating scales (Simpson Angus Scale, Barnes Akathisia Rating Scale, and Abnormal Involuntary Movement Scale) and findings on physical examination. Changes from baseline in physical assessments of weight and body mass index, and measurement of serum prolactin and glucose levels were recorded.

Analysis sets and statistical analysis

The safety population consisted of 216 recently diagnosed subjects and 429 chronically ill subjects who received paliperidone palmitate continuously from study entry through study completion or discontinuation. Demographic and baseline characteristics between groups were compared by t-test for continuous variables and Chi-square or Fisher’s exact test for categorical variables. Between-subgroup differences in continuous measures for post-baseline scores were assessed using analysis of covariance (model with fixed effects) for recently diagnosed versus chronically ill subgroups and country, and the baseline (transition phase) value as a covariate. Change from baseline within groups was assessed by paired t-test. Mean monthly doses of paliperidone palmitate (for days on drug only) and exposure in days were summarized.

Frequencies and percentages of adverse events were summarized from the time of first injection to months 1, 3, 6, 9, 12 weeks.

**Figure 1** Study design and subject disposition in recently diagnosed and chronically ill subgroups treated continuously with paliperidone palmitate.

**Note:** +74 subjects who were in the maintenance phase at the time of its termination and were considered to have completed this phase were eligible to enroll in the open-label extension without entering the double-blind phase. Of the 58 that enrolled, 26 were recently diagnosed and 32 were in the chronic illness subgroup.
and 12 and endpoint. Adverse events reported during these time periods that occurred at a margin of $\geq 2\%$ between the subgroups are reported and displayed by percentage rate in each treatment subgroup. The relative risk (RR) and 95% confidence interval [CI] in the recently diagnosed group compared with the chronically ill subgroup were determined. Unadjusted CI were utilized as a flagging device to identify potentially significant adverse events when the unadjusted CI did include 1. Any adjustment for multiplicity was considered to be counterproductive for identifying potential safety risks.

Any events related to extrapyramidal symptoms, glucose, or prolactin were also summarized. Logistic regression models examined the association between probability of these events with demographic and baseline characteristics that differed between subgroups. These included diagnosis status (recently diagnosed versus chronically ill), age, age at time of diagnosis of schizophrenia, race (black versus Caucasian or other versus Caucasian), weight, body mass index, and smoking status (yes or no).

Changes in weight and glucose and prolactin levels from transition phase baseline through open-label extension were summarized. In addition to analysis of covariance at endpoint, repeated-measures analysis of covariance assessed between-group differences using observed scores to compare mean response profiles over time. The $P$ values for the group effect and group-by-visit interactions were provided. The group effect measured the deviation from the hypothesis of equality of mean changes between groups, averaged over the treatment duration. The group-by-visit interaction tested the hypothesis of parallel response profiles over time. Multiple linear regression models examined relationships between changes in weight, and glucose and prolactin levels, with demographic and baseline characteristics that were found to be different at baseline.

## Results

### Subject baseline demographics, disposition, and dosing

Subjects in the recently diagnosed subgroup were younger than the chronically ill subgroup at study entry, with an older age at the time of diagnosis of schizophrenia, and with a lower weight and body mass index (Table 1). There was a higher percentage of Caucasians and a lower percentage of smokers in the recently diagnosed subgroup than in the chronically ill subgroup. The mean years of illness was $2.9 \pm 1.5$ (range 1.0–5.0) in the recently diagnosed subgroup and $16.2 \pm 8.1$ (range 6.0–47.0) in the chronically ill subgroup. The subgroups were similar with respect to sex (male 60.2% and 58.0%, respectively), mean total Positive and Negative Syndrome Scale score (70.7 and 73.3, $P = 0.0811$, respectively), and Clinical Global Impressions Scale scores (not ill to moderate in 87.0% and 81.8%, respectively). Completion rates were generally similar between the subgroups during each study phase (Figure 1). Among subjects receiving paliperidone palmitate who entered the double-blind relapse prevention phase, the time to relapse was not significantly different between the recently diagnosed and chronically ill subgroups ($P = 0.0999$, log-rank test).

The mean monthly dose was similar in each subgroup (approximately 109 mg [69.9 mg eq]), with a mean duration of exposure of 333.9 $\pm 271.9$ days in the recently diagnosed subgroup and 308.7 $\pm 278.3$ days in the chronically ill subgroup. During the study, 42.1% of those in the recently diagnosed subgroup received benzodiazepines and 10.2% received medications for extrapyramidal symptoms, respectively. In the chronically ill subgroup, rates were 46.4% for benzodiazepines and 17.0% for medications used to treat extrapyramidal symptoms.

### Reported adverse events

During the month following the first injection, 31.5% (68 of 216) of recently diagnosed and 42.7% (183 of 429) of subjects in the chronically ill subgroup reported an adverse

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Transition baseline demographics and characteristics that differed between subgroups</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years (mean $\pm$ SD)</td>
<td>Recently diagnosed (n = 216)</td>
</tr>
<tr>
<td>$&lt;25$</td>
<td>$31.0 \pm 9.3$</td>
</tr>
<tr>
<td>Age distribution, years, n (%)</td>
<td></td>
</tr>
<tr>
<td>$26–50$</td>
<td>$68 (31.5%)$</td>
</tr>
<tr>
<td>$&gt;50$</td>
<td>$6 (2.8%)$</td>
</tr>
<tr>
<td>Race, n (%)</td>
<td></td>
</tr>
<tr>
<td>Caucasian</td>
<td>$156 (72.2%)$</td>
</tr>
<tr>
<td>Black</td>
<td>$13 (6.0%)$</td>
</tr>
<tr>
<td>Asian</td>
<td>$43 (19.9%)$</td>
</tr>
<tr>
<td>Other</td>
<td>$4 (1.9%)$</td>
</tr>
<tr>
<td>Weight, kg (mean $\pm$ SD)</td>
<td>$75.3 \pm 15.94$</td>
</tr>
<tr>
<td>BMI, kg/m$^2$ (mean $\pm$ SD)</td>
<td>$25.9 \pm 4.92$</td>
</tr>
<tr>
<td>Current smoker, n (%)</td>
<td>$92 (42.6%)$</td>
</tr>
<tr>
<td>Age at schizophrenia diagnosis, years (mean $\pm$ SD)</td>
<td>$28.0 \pm 9.1$</td>
</tr>
</tbody>
</table>

Abbreviations: BMI, body mass index; SD, standard deviation.
### Table 2
Subjects with at least one adverse event in the recently diagnosed and chronically ill subgroups (from time since first injection to specified time point)

<table>
<thead>
<tr>
<th>Intervals from first injection</th>
<th>Recently diagnosed (n = 216)</th>
<th>Chronically ill (n = 429)</th>
<th>RR^a (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>To month 1</td>
<td>68 (31.5%)</td>
<td>183 (42.7%)</td>
<td>0.738 (0.589–0.924)</td>
</tr>
<tr>
<td>To month 3</td>
<td>112 (51.9%)</td>
<td>264 (61.5%)</td>
<td>0.843 (0.726–0.978)</td>
</tr>
<tr>
<td>To month 6</td>
<td>129 (59.7%)</td>
<td>296 (69.0%)</td>
<td>0.866 (0.763–0.982)</td>
</tr>
<tr>
<td>To month 9</td>
<td>141 (65.3%)</td>
<td>312 (72.7%)</td>
<td>0.898 (0.801–1.005)</td>
</tr>
<tr>
<td>To month 12</td>
<td>148 (68.5%)</td>
<td>320 (74.6%)</td>
<td>0.919 (0.826–1.021)</td>
</tr>
<tr>
<td>To open-label extension endpoint</td>
<td>153 (70.8%)</td>
<td>330 (76.9%)</td>
<td>0.921 (0.833–1.018)</td>
</tr>
</tbody>
</table>

Notes: Recently diagnosed subgroup compared with the chronically ill subgroup. Data are cumulative; each interval includes events from the previous interval.

Abbreviations: CI, confidence interval; RR, relative risk.

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**Figure 2** Percentage, relative risk (recently diagnosed versus chronic illness), and 95% confidence intervals of adverse events reported by a margin of ≥2% in recently diagnosed or chronically ill subgroups.

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0 2 3 4 5 6 7 8 9 10 12 14 16 18

- **Incidence (%)**
- **Relative risk (95% confidence interval) recently diagnosed compared to chronic illness**

### Diagram Details
- **Month 1**
  - Insomnia
  - Schizophrenia
  - Agitation
- **Month 2**
  - Insomnia
  - Schizophrenia
  - Psychotic disorder
  - Nasopharyngitis
  - Weight increased
- **Month 3**
  - Insomnia
  - Schizophrenia
  - Psychotic disorder
  - Nasopharyngitis
  - Blood cholesterol increased
- **Month 4**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Month 5**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Month 6**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Month 7**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Month 8**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Month 9**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Month 10**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Month 11**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Month 12**
  - Insomnia
  - Schizophrenia
  - Nasopharyngitis
  - Blood glucose increased
- **Open-label extension endpoint**
  - Insomnia
  - Schizophrenia
  - Agitation
  - Psychotic disorder
  - Nasopharyngitis
  - Urinary tract infection
  - Delirium

- **Favors recently diagnosed subgroup**
- **Favors chronic illness subgroup**
event (Table 2). In general, incidence rates, RR, and 95% CI suggested that adverse events were less likely in the recently diagnosed subgroup than in the chronically ill subgroup at each time interval.

In the month following the first injection, no adverse events were reported at a margin of ≥2% in more of the recently diagnosed subjects than in those with chronic illness (Figure 2). Insomnia, worsening of schizophrenia, and agitation were reported at a margin of ≥2% in more of the chronically ill subjects than in recently diagnosed subjects. These differences in incidence were not considered potentially significant based upon the 95% CI. Nasopharyngitis was reported by more subjects with chronic illness than by those in the recently diagnosed subgroup of subjects at months 6 (4.9% versus 1.4%; RR 0.28; 95% CI 0.086–0.941 respectively), 9 (5.6% versus 1.9%; RR 0.33; 95% CI 0.116–0.942), and 12 (6.1% versus 2.3%; RR 0.38; 95% CI 0.149–0.981), and endpoint (7.2% versus 2.8%; RR 0.38; 95% CI 0.163–0.907); the 95% CI did not include 1 and were considered potentially significant. Influenza (2.8% versus 0.7%; RR 3.97; 95% CI 1.003–15.730) and amenorrhea (3.2% versus 0.9%; RR 3.48; 95% CI 1.029–11.744) were reported by more subjects in the recently diagnosed subgroup at endpoint. Given that the 95% CI did not include 1, these events were also considered potentially significantly different between subgroups.

In the recently diagnosed subgroup, sedation was reported by none, one, or two subjects at each time point (0.0%–0.93%), with somnolence reported by one or two subjects at each time point (0.46%–0.93%), and five subjects at endpoint (2.31%). In the chronically ill subgroup, sedation was reported by two subjects at each time point (0.47%), with somnolence reported by 1–5 subjects at each time point (0.23%–1.17%).

**Events related to extrapyramidal symptoms**

Extrapyramidal symptom-related adverse event rates included parkinsonism, hyperkinesia, dystonia, tremor, and dyskinesia. Rates of any extrapyramidal symptom-related events were numerically lower in the recently diagnosed subgroup at each period assessed (through months 1, 3, 6, 9, and 12 and open-label extension). Rates ranged from 2.3% to 9.3% in the recently diagnosed subgroup and from 5.8% to 12.6% in the chronically ill subgroup (Figure 3). Rates of individual extrapyramidal symptom-related events and their specific preferred terms are summarized in Table 3. Among the specific preferred terms, nonspecific extrapyramidal disorder was reported by more recently diagnosed subjects (4.6%) than by chronically ill subjects (2.3%), and akathisia was reported by more chronically ill subjects (3.3%) than recently diagnosed subjects (1.9%).

Logistic regression models showed that none of the baseline characteristics that differed between the subgroups was associated with risk of extrapyramidal symptom-related events (ie, age, $P = 0.7057$; weight, $P = 0.8921$; body mass index, $P = 0.9823$; age at diagnosis of schizophrenia, $P = 0.7727$; current smoking status, $P = 0.3507$; and race [black, $P = 0.0950$; other, $P = 0.1602$]).

**Weight changes**

Weight changes (least squares mean ± standard error) at endpoint were 2.6 ± 0.9 kg in the recently diagnosed subgroup and 3.4 ± 0.7 kg in the chronically ill subgroup ($P = 0.42$; least squares mean difference 0.8 ± 0.99, 95% CI: −1.15–2.75). Observed scores at each time point were evaluated using repeated-measures analysis of covariance. Average changes from baseline to endpoint between groups were similar (between-group comparison, $P = 0.4342$); this finding did not differ throughout the trial (group-by-visit interaction, $P = 0.9520$). Linear regression models assessing the impact of baseline differences did not reveal any potential

<table>
<thead>
<tr>
<th>Table 3 Extrapyramidal symptom-related adverse events from first injection through open-label extension phase, in recently diagnosed and chronically ill subgroups</th>
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</thead>
<tbody>
<tr>
<td><strong>Extrapyramidal adverse events and preferred terms</strong></td>
</tr>
<tr>
<td><strong>Subjects with an event, n (%)</strong></td>
</tr>
<tr>
<td>Parkinsonism</td>
</tr>
<tr>
<td>Extrapyramidal disorder</td>
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<tr>
<td>Hypertonia</td>
</tr>
<tr>
<td>Musculoskeletal stiffness</td>
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<tr>
<td>Drooling</td>
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<tr>
<td>Muscle tightness</td>
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<tr>
<td>Hyperkinesia</td>
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<tr>
<td>Akathisia</td>
</tr>
<tr>
<td>Restlessness</td>
</tr>
<tr>
<td>Dystonia</td>
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<tr>
<td>Muscle spasms</td>
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<tr>
<td>Dystonia</td>
</tr>
<tr>
<td>Oculogyric crisis</td>
</tr>
<tr>
<td>Tremor</td>
</tr>
<tr>
<td>Dyskinesia</td>
</tr>
<tr>
<td>Dyskinesia</td>
</tr>
<tr>
<td>Muscle twitching</td>
</tr>
<tr>
<td>Tardive dyskinesia</td>
</tr>
</tbody>
</table>

**Notes:** *A subject experiencing more than one adverse event within a system organ class/preferred term is counted once within that system organ class/preferred term for incidence.*
Glucose-related measures

Rates of glucose-related adverse events were 5.1% (22 of 429) in the chronically ill subjects and 2.8% (6 of 216) in recently diagnosed subjects, with “blood glucose increased” being the most frequent (4.0% versus 1.9%, Table 4). In logistic regression models, age (P = 0.0001; odds ratio [OR] 1.087; 95% CI 1.047–1.129) had a significant association with incidence of glucose-related adverse events (one-year increase in age is associated with an 8.7% increase in odds of experiencing “blood glucose increased”), whereas body mass index (P = 0.0516; OR 1.057; 95% CI 1.000–1.118) and age at diagnosis of schizophrenia (P = 0.0578; OR 1.040; 95% CI 0.999–1.082) showed trends towards significance. Other baseline characteristics had no impact (ie, weight, P = 0.4310; current smoking status, P = 0.6082; and race [black, P = 0.1737; other, P = 0.0810]).

Mean glucose levels at open-label endpoint were not significantly different between the groups (least squares mean difference −0.1, 95% CI −0.61–0.32, P = 0.5436). Regression models showed that age (P = 0.0126, regression coefficient 0.025 ± 0.010) and age at diagnosis of schizophrenia (P = 0.0021, regression coefficient 0.042 ± 0.014) were associated with changes at endpoint.

Prolactin-related measures

Prolactin levels increased in both sexes in both subgroups. Mean prolactin values were consistently higher in recently diagnosed female subjects compared with chronically ill female subjects (Table 5). Potentially prolactin-related adverse events were reported by 7.9% of recently diagnosed subjects compared with 3.5% of chronically ill subjects through endpoint (Table 6). The most commonly reported events were amenorrhea (7 [3.2%] and 4 [0.9%], respectively) and galactorrhea (2 [0.9%] and 6 [1.4%], respectively).

Discussion

The primary objective of this post hoc analysis was to examine the long-term tolerability of once-monthly injectable paliperidone palmitate in subjects with a more recent diagnosis of schizophrenia compared with those with a longer duration of illness. Among the range of events anticipated to occur more commonly in the recently diagnosed subgroup, only events related to prolactin elevation emerged as more likely to be increased. Weight increases, sedation/somnolence, overall extrapyramidal symptoms, and glucose-related events did not occur at a greater rate in recently diagnosed subjects. Strengths of this analysis include the length and
Recently diagnosed (n = 216) | Chronic illness (n = 429)
---|---
Subjects reporting a potentially prolactin-related adverse event, n (%) | 17 (7.9) | 15 (3.5)

**Specific potentially prolactin-related event, n (%):**

- Amenorrhea: 7 (3.2) | 4 (0.9)
- Galactorrhea: 2 (0.9) | 6 (1.4)
- Menstruation irregular: 2 (0.9) | 3 (0.7)
- Erectile dysfunction: 2 (0.9) | 2 (0.5)
- Sexual dysfunction: 2 (0.9) | 1 (0.2)
- Oligomenorrhea: 1 (0.5) | 1 (0.2)
- Breast pain: 1 (0.5) | 0 (0)
- Breast tenderness: 0 (0) | 1 (0.2)
- Gynecomasia: 1 (0.5) | 0 (0)
- Hyperprolactinemia: 1 (0.5) | 0 (0)
- Blood prolactin increased: 1 (0.5) | 0 (0)
- Libido decreased: 0 (0) | 1 (0.2)

**Notes:** A subject experiencing more than one adverse event within a system organ class/preferred term is counted once within that system organ class/preferred term for incidence.

Size of the study. This multiphase study database provides the longest exposure data currently available with the paliperidone palmitate once-monthly injection in patients early in the course of their illness. Nevertheless, these findings are limited in that this work represents the post hoc analysis of a single study.

Relevant to the tolerability findings, the recently diagnosed and chronically ill subgroups had similar doses, durations of treatment exposure (>300 days in each subgroup), and discontinuation rates. These treatment similarities deserve comment because one might have expected lower doses and/or shorter exposures to be used in subjects with early illness if they tolerated the drug less well than more chronically ill subjects. However, it is relevant to note that the paliperidone palmitate doses allowed in this study (39, 78, 117, and 156 mg [25, 50, 75, and 100 mg eq]) did not include the highest available 234 mg (150 mg eq) dose. Further, initiation doses were lower than the currently recommended initiation dosage (234 mg day 1 and 156 mg day 8 [150 mg eq day 1 and 100 mg eq day 8]). Therefore, while similarities were noted between the two subgroups, the overall rates of adverse events reported during the first month may have been lower than what would have been seen with the currently recommended initiation regimen. Pertinent to this, somewhat higher overall adverse event rates were observed in a recently published post hoc analysis of a double-blind, placebo-controlled trial in recently diagnosed subjects receiving the recommended day 1 and day 8 doses. During the week following the initial 234 mg (150 mg eq) paliperidone palmitate or placebo injection, 37.6% (41 of 109) and 29.7% (11 of 37), respectively, of subjects reported an adverse event. During the month following the day 8 injection of paliperidone palmitate 156 mg (100 mg eq) or placebo, adverse event rates were 41.0% (16 of 39) and 37.8% (14 of 37), respectively. During the first week, a broad range of events reported in one or two paliperidone palmitate subjects contributed to the higher rate; the events reported more often by patients on active treatment than by those on placebo were injection site pain, agitation, and headache. In the month following the second injection, anxiety was the most common event, and was reported more often by patients on active treatment than by those on placebo. These events did not emerge as more likely to occur in the recently diagnosed versus more chronic subjects in the present analysis (Figure 2).

Rates of any extrapyramidal symptom-related events, which are of particular concern in patients with early illness,
were generally similar or numerically lower in this analysis of recently diagnosed compared with chronically ill subjects. Rates tended to stabilize after month 3, with incremental increases becoming smaller with continued treatment. Findings related to specific types of extrapyramidal symptoms also require consideration. As reported, there was a somewhat higher rate of nonspecifically coded extrapyramidal symptoms in the recently diagnosed subgroup. In the previously mentioned post hoc analysis of recently diagnosed subjects from a double-blind, placebo-controlled trial, movement-related event rates were 10.3% (4 of 39) with paliperidone palmitate (234 mg day 1 and 156 mg day 8 [150 mg eq day 1 and 100 mg eq day 8] and monthly thereafter) and 8.1% (3 of 37) with placebo over 13 weeks (RR 1.3; 95% CI 0.30–5.27; *P > 0.05*). Among the specific types of events, the most common in the recently diagnosed population was parkinsonism (7.7% paliperidone palmitate and 0% placebo). In the current long-term evaluation, parkinsonism was again the most common movement disorder-related event, with the same rate in recently diagnosed and more chronic subjects (5.6%).

Prolactin findings were also anticipated by prior work. Levels increased in both subgroups and in both sexes, with higher levels in females with early illness compared with those having chronic illness, and a higher percentage of recently diagnosed versus chronically ill subjects reported prolactin-related adverse events (7.9% versus 3.5%, respectively).

The limitations to these findings include the fact that the original study was not designed to assess long-term tolerability of the drug in patients with recently diagnosed schizophrenia. The 5-year cut point used to define early illness relied on historical information and patient self-reporting, which may have had variable reliability, and many patients may have been ill for some time prior to receiving a formal diagnosis. Nonetheless, it is generally accepted that the first 5–10 years of illness is a critical period for effective intervention. Using this 5-year cut point likely captured a population that was enriched with those at an earlier stage of schizophrenia. Also, although there was no comparison with a placebo group in this analysis, these findings are still relevant to the question at hand regarding the long-term tolerability of paliperidone palmitate in subjects early in the course of their illness compared with those with more chronic illness. An unexpected finding of this safety analysis is the similar or even somewhat lower rate of total adverse events or any extrapyramidal-related events reported by recently diagnosed compared with subjects with chronic illness. This is at variance with other reports in the literature. Also, with few exceptions, baseline phenotypes did not have a significant effect on the results.

In conclusion, these long-term findings complement prior tolerability analyses of the initiation dosing of injectable paliperidone palmitate in subjects with recently diagnosed schizophrenia, and may help guide clinicians in the management of these patients.

**Disclosure**

LA, JKS, D-JF, and CAB are employees of Janssen Scientific Affairs LLC, Titusville, NJ. IT is an employee of Janssen Research and Development LLC, Titusville, NJ. This research was funded by Janssen Scientific Affairs LLC. Writing, editorial, and technical support services were provided by Susan Ruffalo of MedWrite Inc, Newport Coast, CA. Some of these data were presented at the 164th annual meeting of the American Psychiatric Association, held on May 14–18, 2011, Honolulu, HI, and at the 13th International Congress on Schizophrenia Research, April 2–6, 2011, Colorado Springs, CO.

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