Clinical outcomes with olanzapine long-acting injection: impact of the 3-hour observation period on patient satisfaction and well-being

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Background: The objective of the present analysis is to determine the impact of the 3-hour observation period for olanzapine long-acting injection (LAI) on patient satisfaction and well-being by comparing data collected before and after its implementation.

Methods: This is a post hoc analysis of patients treated with olanzapine LAI in 1) a 6-month fixed-dose randomized controlled trial and/or 2) a 6-year open-label safety study. This analysis was limited to patients with schizophrenia who were treated with olanzapine LAI consistent with the approved indication and dosing recommendations of the European Union Summary of Product Characteristics (N=966). Of the 966 patients, the analysis further focused only on those patients who received both 1) at least one injection before the implementation of the 3-hour observation period and 2) at least one injection after implementation of the 3-hour observation period (N=487). Patient satisfaction was assessed with the three-item Patient Satisfaction with Medication Questionnaire-Modified. Responses were averaged across all postbaseline visits occurring before (ie, without) the implementation of the 3-hour observation period and across all postbaseline visits occurring after (ie, with) the implementation of the 3-hour observation period. In addition, the rate of postinjection delirium/sedation syndrome events was calculated.

Results: There was no meaningful change after implementation of the 3-hour observation period in satisfaction (before: mean [SD] = 4.0 [1.02] and after: mean [SD] = 4.1 [0.82]), preference for olanzapine LAI over oral medication (before: mean [SD] = 4.0 [0.90] and after: mean [SD] = 4.1 [0.77]), or ratings of satisfaction regarding side effects (before: mean [SD] = 1.9 [0.79] and after: mean [SD] = 1.8 [0.60]). For the total population (N=966), postinjection delirium/sedation syndrome occurred in 26 (0.07%) of 38,010 injections.

Conclusion: For patients with schizophrenia receiving treatment with olanzapine LAI, the 3-hour observation period had no impact on their satisfaction with the medication or on their subjective well-being.

Keywords: olanzapine long-acting injection, observation period, schizophrenia, Europe

Introduction

Olanzapine long-acting injection (LAI), a pamoate salt depot formulation of olanzapine, was formally introduced into clinical practice for the treatment of schizophrenia in 2008 following nearly a decade of prior research and development.1 In 2006, during this research and development program, an important safety signal was detected, namely postinjection delirium/sedation syndrome (PDSS). PDSS is believed to be associated with vessel injury during the injection process, resulting in accidental intravascular entry of a portion of the dose. Because olanzapine LAI is a salt-based formulation, it is more soluble in blood than in muscle tissue, resulting in symptoms consistent
with olanzapine intoxication, particularly excessive sedation (which could include coma) and/or delirium.\textsuperscript{2,4}

The identification of PDSS ultimately resulted in the European label for olanzapine LAI having a risk minimization plan associated with it, the key elements being a 3-hour observation period and for patients to be accompanied home at the end of this 3-hour time period.\textsuperscript{3} In 2013, the European Medicines Agency announced that the safety signal of olanzapine LAI was sufficient to warrant the removal of the requirement of patients to be accompanied home, but the mandatory stipulation of the 3-hour observation period still remains.\textsuperscript{3,5}

The objective of the present analysis was to determine the impact of the 3-hour observation period on patient satisfaction and well-being by comparing data collected before and after its implementation.

**Methods**

**Patients and design**

This is a post hoc analysis of data from two studies. Study F1D-MC-HGKA was a maintenance study in which patients (18–75 years of age) with schizophrenia were stabilized on oral olanzapine for 4–8 weeks and then randomized to 24 weeks of double-blind treatment with olanzapine LAI (150 mg every 2 weeks, 405 mg every 4 weeks, 300 mg every 2 weeks, or 45 mg every 4 weeks [total N=743]) or their stable oral olanzapine dose (N=322).\textsuperscript{4} Study F1D-MC-HGKB was a 6-year, single-arm, open-label Phase III study of olanzapine LAI in patients (18–76 years of age; N=931) with schizophrenia or schizoaffective disorder from three feeder studies (F1D-MC-HGJZ, F1D-MC-HGKA, and F1D-EW-LOBS).\textsuperscript{7}

In that study, patients received 210 mg olanzapine LAI at the first open-label visit. Patients received doses between 45 mg and 405 mg of olanzapine LAI at 2-week, 3-week, or 4-week intervals. Dosing after the first visit was determined on the basis of the investigators’ clinical judgment.

The studies were conducted in accordance with ethics principles derived from international ethics guidelines, including the Declaration of Helsinki and the Council for International Organizations of Medical Sciences (CIOMS) International Ethical Guidelines, the International Conference on Harmonization (ICH) Good Clinical Practices (GCP) Guideline, and applicable laws and regulations. In addition, the study protocols were approved by ethical review boards at each site (Table S1). After receiving a complete description of the study, all patients and/or their authorized legal representatives provided written informed consent before participation.

The present analysis was limited to patients who were treated with olanzapine LAI consistent with the approved indication and dosing recommendations of the European Union (EU) Summary of Product Characteristics (SmPC),\textsuperscript{5} that is, only those patients who met the diagnostic criteria for schizophrenia according to the *Diagnostic and Statistical Manual of Mental Disorders*, fourth edition, Text Revision,\textsuperscript{8} were included and patients with schizoaffective disorder were excluded. Patients who received oral olanzapine supplementation and who had a total daily dose of >20 mg/d oral olanzapine equivalent were also excluded. This patient population (N=966) was used in the calculation of the rate of PDSS events. For the analysis of patient satisfaction and well-being, the patient population was further refined to include only those patients who received both 1) at least one injection before the implementation of the 3-hour observation period and 2) at least one injection with the 3-hour observation period (N=487).

**Scales and measures**

All questionnaires and assessments (except the Clinical Global Impressions-Severity [CGI-S]) were completed every 6 months.

Patient satisfaction was assessed with the three-item Patient Satisfaction with Medication Questionnaire-Modified (PSMQ-Mod): satisfaction with current depot medication, preference between current depot medication and previous oral medications, and comparison of side effects between current depot medication and previous oral medications. The PSMQ-Mod uses a 5-point scale, with 5 representing highest satisfaction/preference for olanzapine LAI on the first two items and a score of 1 representing fewer side effects with olanzapine LAI on the third item.\textsuperscript{9}

Patient well-being was assessed with the Subjective Well-Being Under Neuroleptics Scale-Short Form (SWNS). The SWNS has five subscores (self-control, emotional regulation, mental functioning, physical functioning, and social integration), each consisting of four questions with a total of 20 statements (ten positive and ten negative) and a possible total score of 20–120, with higher scores indicating greater subjective well-being.\textsuperscript{10}

The 36-item Short Form Health Survey (SF-36) is a multipurpose, short-form patient-reported health survey used to evaluate overall physical and mental health. The SF-36 has eight subscales: bodily pain, general health, physical functioning, role-physical (contributing to the SF-12 physical score), mental health, role-emotional, social functioning, and vitality (contributing to the SF-12 mental score). Each...
scale has a total possible score of 0–100, with higher scores indicating less disability.11

The Quality of Life (QoL) scale was used to measure the
effect of schizophrenia symptoms on occupational,
social, and psychological functioning in patients with
schizophrenia. The total score of the scale is the sum of
21 items, with each item being rated on a 7-point scale. Four
categories contained in the scale are intrapsychic founda-
tions (items 13–17 and 20 and 21), interpersonal relations
(items 1–8), instrumental role (items 9–12), and common
objects and activities (items 18 and 19), with higher scores
indicating less impairment.12

The Positive and Negative Syndrome Scale (PANSS)
was used to evaluate the symptoms of schizophrenia. The
scale has a total possible score of 30–210, with higher scores
indicating more severe symptomatology.13

The CGI-S was used to assess the severity of schizoph-
phrenia. The single question is rated on a 7-point scale ranging
from 1 (normal, not at all ill) to 7 (among the most extremely
ill).14 The CGI-S was assessed at quarterly visits.

The rate of PDSS events was also summarized.

Statistical analyses
All patients who received both at least one injection before
the implementation of the 3-hour observation period and at
least one injection with the 3-hour observation period were
included in this post hoc analysis. Demographics and baseline
characteristics were summarized by their frequencies. For all
health outcome measures and efficacy measures, responses
were averaged within patients across all postbaseline visits
occurring before (ie, without) the implementation of the
3-hour observation period and across all postbaseline visits
occurring after (ie, with) the implementation of the 3-hour
observation period. No adjustments were made to variance
estimates to account for the different number of measure-
ments taken between patients. Due to exploratory nature of
the analyses, only descriptive statistics are provided.

Clinical trial registration
The trials from which data are reported here were registered on
ClinicalTrials.gov as NCT00088465 and NCT00088491.

Results
A total of 966 patients were treated with olanzapine LAI
consistent with the approved indication and dosing recom-
endations of the EU Summary of Product Characteristics
(SmPC). This patient population was used for the calculation
of the rate of PDSS events. Of these 966 patients, the present
analysis further focused on those patients (n=487) who
received at least one injection before the implementation of
the 3-hour observation period and at least one injection with
the 3-hour observation period. The mean (SD) age of these
patients was 38.2 (11.75) years, with most patients being
male (66.3%) and Caucasian (67.1%) (Table 1).

Among patients who were treated both before and after
implementation of the 3-hour observation period, there was
no meaningful change in their satisfaction with olanzapine
LAI (before: mean [SD] =.4.0 [1.02] and after: mean [SD] =.4.1
[0.82]), preference for olanzapine LAI over oral medication
(before: mean [SD] =.0.0 [0.79] and after: mean [SD] =.4.1
[0.77]), or ratings of satisfaction regarding side effects
(before: mean [SD] =.1.9 [0.79] and after: mean [SD] =.8.00
[0.60]) (Figure 1).

Total scores on the SWNS were similar before and after
implementation of the 3-hour observation period (before:
mean [SD] =78.14 [14.28] and after: mean [SD] =88.6
[13.14]). Similar results were reported for subscale scores
of self-control, emotional regulation, mental functioning,
physical functioning, and social integration (Figure 2).

No meaningful changes were reported on any of the sub-
scapes of the SF-36, including the SF-12 physical subscale
(before: mean [SD] =.50.8 [6.68] and after: mean [SD] =.50.8
[6.69]) and SF-12 mental subscale (before: mean [SD] =.45.2
[9.49] and after: mean [SD] =.46.6 [8.35]), scales after imple-
mentation of 3-hour observation period compared to before
implementation (Figure 3).

On the QoL scale, the total score was numerically higher
after implementation of 3-hour observation period compared
to before implementation of 3-hour observation period, but
the difference in scores was not considered clinically mean-
ingful (total score, before: mean [SD] =.70.5 [19.7] and after:
mean [SD] =.77.3 [21.07]). Similar results were reported in
the subscale scores of the QoL (Figure 4).

Table I Demographics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Patients (N=487)</th>
</tr>
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<tr>
<td>Age, mean (SD)</td>
<td>38.2 (11.75)</td>
</tr>
<tr>
<td>Sex, n (%)</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>164 (33.7)</td>
</tr>
<tr>
<td>Male</td>
<td>323 (66.3)</td>
</tr>
<tr>
<td>Race, n (%)</td>
<td></td>
</tr>
<tr>
<td>African</td>
<td>30 (6.2)</td>
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<tr>
<td>Caucasian</td>
<td>327 (67.1)</td>
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<tr>
<td>East Asian</td>
<td>22 (4.5)</td>
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<tr>
<td>Hispanic</td>
<td>92 (18.9)</td>
</tr>
<tr>
<td>Native American</td>
<td>1 (0.2)</td>
</tr>
<tr>
<td>West Asian (Indian subcontinent)</td>
<td>15 (3.1)</td>
</tr>
</tbody>
</table>

Notes: N, number of patients; n, number of affected patients.
The PANSS total score, as well as the positive symptoms, negative symptoms, and general psychopathology subscale scores, was similar before and after implementation of the 3-hour observation period (Figure 5). Scores on the CGI-S were also similar before and after implementation of the 3-hour observation period (before: mean [SD] =2.7 [0.91] and after: mean [SD] =2.5 [0.86]).

Of the 38,010 injections given to patients with schizophrenia who were treated with olanzapine LAI consistent with the approved indication and dosing recommendations of the EU SmPC (N=966), 26 (0.07%) were followed by a PDSS event. The percentage of injections resulting in PDSS events was similar before (10/13, 660=0.07%) and after (16/24, 350=0.07%) implementation of the 3-hour observation period.

**Discussion**

On measures of patient satisfaction, well-being, health, QoL, and symptoms of schizophrenia, we found no evidence of an effect of the 3-hour observation period. On each measure, scores were only slightly different before and after implementation of the 3-hour observation period. Also, as expected, the percentage of injections resulting in a PDSS event was essentially the same before and after implementation of
the 3-hour observation period. The percentage of PDSS events that occurred following injections with olanzapine LAI in this patient sample is similar to that previously reported (<0.1%).

It should also be noted that, overall, these patients had relatively low symptom levels, as patients were generally clinically stable when they began treatment with olanzapine LAI in this clinical study. This is evident by their responses on the PANSS and CGI-S and supported by their responses on the SWNS, SF-36, and QoL scale.

Overall, on the PSMQ-Mod, the mean responses both before and after implementation indicated that patients experienced less side effects with olanzapine LAI and preferred olanzapine LAI compared to their oral medication and that they were satisfied with olanzapine LAI. As expected, these results agree with a previous report.
(from a largely overlapping patient sample), indicating that 73.7% of patients being treated with olanzapine LAI were somewhat or very satisfied with their medication, 69.5% preferred or much preferred olanzapine LAI versus previous oral medications, and 74.2% thought that olanzapine LAI was associated with “less” or “much less side effects”.

Though it might not be expected that measures of efficacy and well-being would be affected by the 3-hour observation period, it seems reasonable to hypothesize that the initiation of a 3-hour observation period would result in a decrease in satisfaction among olanzapine LAI patients because of the potential resulting inconvenience and interruption of daily routine. However, if this effect is influencing scores at all, it is counterbalanced by other factors. For example, one might expect scores to improve somewhat during the course of an open-label study as some patients continue to improve with treatment and others do not do as well and drop out. However, there is a less obvious possibility. Anecdotal evidence suggests that patients may find that additional time with their mental health care team may be a positive experience for them. Naber\textsuperscript{16} suggests that positive benefits (eg, patient acceptance and better therapeutic relationship) may result from regular contact with their health care providers. Indeed, the 3-hour observation period could well be further utilized by offering psychotherapy or psychoeducation.

**Limitations**

There are a few limitations of this analysis that should be recognized. First, with no a priori sample size calculation, it is unclear whether the sample of subjects used was large enough to detect a difference in our measures, if one existed. Second, our conclusions are based on interpretation of a body of descriptive statistics rather than predefined inferential tests. Third, this analysis was limited to patients with schizophrenia who were treated with olanzapine LAI consistent with the approved indication and dosing recommendations of the EU SmPC. Results that included responses of patients with schizoaffective disorder, or patients who were given supplementary oral doses of olanzapine beyond those recommended by the EU SmPC, could have been different.

**Conclusion**

For patients with schizophrenia receiving treatment with olanzapine LAI consistent with the approved indication and dosing recommendations of the EU SmPC, the 3-hour observation period had no impact on their satisfaction with the medication and on their subjective well-being, overall health, QoL, or symptoms of schizophrenia. It is hoped that these results will help EU clinicians further define and refine how best they would like to use olanzapine LAI in treating their patients with schizophrenia, especially in keeping within the current requirements of the EU SmPC.

**Acknowledgments**

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Disclosure
All authors are the employees and minor shareholders of Eli Lilly and Company. The authors report no other conflicts of interest in this work.

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