A commentary on the efficacy of olanzapine for the treatment of schizophrenia: the past, present, and future

Abstract: Olanzapine is a second-generation atypical antipsychotic with proven efficacy for the treatment of schizophrenia. Approved in 1996, olanzapine is one of the most studied antipsychotics, resulting in a considerable amount of clinical data across diverse patient populations. Despite the fact that olanzapine is associated with a known risk of metabolic side effects, including weight gain, many clinicians continue to prescribe olanzapine for the treatment of schizophrenia with the expectation of additional therapeutic antipsychotic efficacy relative to other first-line atypical antipsychotics. The goal of this narrative is to revisit the role of oral olanzapine in the management of patients with schizophrenia, including those with recently diagnosed schizophrenia (“first-episode”), those with an established schizophrenia diagnosis who experience acute exacerbations, those receiving long-term antipsychotic treatment as a maintenance intervention, and those with suboptimal response to antipsychotic treatment, including treatment resistance. Collectively, data from published literature support the favorable efficacy of olanzapine compared with other first- and second-generation antipsychotics, including lower rates of treatment discontinuation and clinically meaningful improvements in the symptoms of schizophrenia. The development of antipsychotic medications with the favorable efficacy of olanzapine, but with reduced weight gain, could address a major unmet need in the treatment of schizophrenia.

Keywords: antipsychotic, efficacy, metabolic dysregulation, weight gain

Introduction

Olanzapine is an atypical antipsychotic first approved by the US Food and Drug Administration (FDA) in September 1996 for the treatment of schizophrenia.1,2 The availability of olanzapine and the atypical class in general provided psychiatrists with treatment options with reduced risk of motoric adverse effects.3 Several meta-analyses have been published establishing the efficacy and minimal extrapyramidal adverse effects of olanzapine compared to typical and some atypical antipsychotics in the treatment of schizophrenia.4–6 However, due to the risk of metabolic dysregulation and weight gain, most guidelines have positioned olanzapine as a second-choice therapy for the treatment of schizophrenia.7–11 This is similar to clozapine, which, despite its established efficacy for treatment-resistant symptoms, is positioned last because of safety concerns.11,12 Many experienced clinicians are of the opinion that olanzapine offers a distinctly favorable efficacy profile among the currently available antipsychotic medications,4–6,13 and therefore continue to prescribe the drug for patients with schizophrenia. This narrative review revisits the...
role of oral olanzapine in the management of adults with schizophrenia, examining evidence supporting the additional efficacy of olanzapine from the published literature since its initial approval.

**Initial discovery and preclinical data**

Olanzapine was first discovered while searching for a chemical analog of clozapine that would not require hematological monitoring. While investigating a series of thiophene isosteres on 1 of the phenyl rings in clozapine, a thienobenzodiazepine analog (olanzapine) was discovered.\(^1,14,15\) The receptor binding affinities for olanzapine were determined using in vitro assays\(^16\) and confirmed by in vivo functional assays.\(^17\) Olanzapine has high binding affinity for D1, D2, D4, 5HT2A, 5HT2C, 5HT3, α-1 adrenergic, histaminergic, and muscarinic receptors, which mirrors that of clozapine.\(^16\) In functional studies, olanzapine was a potent antagonist at 5HT and D2 receptors, suggesting it might be useful as an antipsychotic agent.\(^17\) In animal models, olanzapine produced antagonism of apomorphine-induced climbing and conditioned avoidance responding in rats, suggesting dopaminergic effects; antagonism of 5-hydroxytryptophan-induced head twitching and blockade of oxtremorine-induced tremor in mice indicate serotoninergic and anticholinergic activity, respectively.\(^18\) In addition, although not binding directly, research has indicated that olanzapine modulates N-methyl-d-aspartate (NMDA) receptors.\(^19,20\) These preclinical studies provided strong evidence that olanzapine could potentially be an effective antipsychotic drug with less drug-induced motoric activation associated with the existing antipsychotics at the time (see Safety section).\(^18,21\) The clinical pharmacokinetic profile of olanzapine supported once-daily oral administration, starting with a dose of 5–10 mg, with a target dose of 10 mg/day, and a maximum dose of 20 mg/day.\(^1\) Taken together, the preclinical and pharmacokinetic data helped to support the full clinical development and eventual approval of olanzapine for the treatment of schizophrenia.\(^2\)

The approval of olanzapine in schizophrenia was initially based on data from short-term controlled trials (6 weeks).\(^22\) A longer-term maintenance trial (up to 8 months) of adults who met DSM-III-R/DSM-IV criteria for schizophrenia was later conducted.\(^1\) Since its approval over 2 decades ago, olanzapine has been one of the most investigated drugs in its class, leading to the generation of a considerable amount of clinical data on its efficacy and safety in patients with schizophrenia and other indications.\(^1,3,23\)

**Assessment of antipsychotic effects**

Schizophrenia is a chronic disease with a diverse psychopathology and multiple phases of illness.\(^11,24\) Consequently, numerous factors must be considered when assessing the benefits of a given treatment over short- and long-term periods.\(^11,25\) A number of scales and instruments are used in clinical trials of schizophrenia to assess treatment efficacy; these have been reviewed in detail elsewhere.\(^26–29\) Improvements in the psychopathologic symptoms of schizophrenia\(^26\) are commonly assessed using the 18-item Brief Psychiatric Rating Scale (BPRS),\(^30\) the Clinical Global Impression (CGI) scale,\(^29,31\) or the 30-item Positive and Negative Syndrome Scale (PANSS).\(^32\) Another well-recognized measure of treatment effectiveness that takes into consideration not only efficacy and tolerability but also patient experience is all-cause discontinuation.\(^33–35\)

The effect of antipsychotic treatment on hostility, given its close relationship to aggression and nonadherence to treatment,\(^36\) has also been extensively studied,\(^37\) commonly by examining treatment-associated changes to the hostility item of the PANSS\(^32\) or BPRS\(^30\) as a proxy measure of aggression.

Here, we summarize key evidence for the efficacy of olanzapine across various stages of treatment in patients with schizophrenia (Table 1).

**First-episode schizophrenia**

Patients presenting with a first psychotic episode require prompt administration of an effective antipsychotic medication to minimize the duration of untreated psychosis.\(^11\) The first episode of psychosis can be polymorphic and may evolve into a variety of conditions other than schizophrenia, with associated prognostic implications. Regardless of the eventual diagnosis, early intervention during a psychotic episode may prevent future deterioration and progression toward chronicity.\(^11\)

Several studies have provided evidence of olanzapine’s efficacy in first-episode schizophrenia patients.\(^38–41\) Notably, patients treated with olanzapine were less likely to discontinue treatment compared with other antipsychotics. In the large-scale European First-Episode Schizophrenia Trial (EUFEST), olanzapine had the lowest rate of all-cause discontinuation (Figure 1) at 1 year (33%) compared with haloperidol (72%), quetiapine (53%), ziprasidone (45%), and amisulpride (40%).\(^40\) In a separate trial, nearly 2 times as many patients remained on olanzapine (23%) compared with haloperidol (12%) at 2 years.\(^39\) In both studies, treatment with olanzapine
Table 1 Selected first-episode, acute, chronic/maintenance, and treatment-resistant schizophrenia trials assessing efficacy of olanzapine

<table>
<thead>
<tr>
<th>Trial/Author</th>
<th>Design</th>
<th>Comparators</th>
<th>N (Patient Population)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sanger et al&lt;sup&gt;38&lt;/sup&gt;</td>
<td>6-week acute phase of a large, prospective, double-blind, multicenter, international</td>
<td>Olanzapine vs haloperidol</td>
<td>83 (first episode)</td>
</tr>
<tr>
<td>Green et al&lt;sup&gt;19&lt;/sup&gt;</td>
<td>2-year randomized, double-blind, multicenter, international</td>
<td>Olanzapine vs haloperidol</td>
<td>263 (first episode)</td>
</tr>
<tr>
<td>EUFEST (Kahn et al)&lt;sup&gt;40&lt;/sup&gt;</td>
<td>Open-label, randomized, multicenter, international</td>
<td>Haloperidol vs second-generation antipsychotics (amisulpride, olanzapine, quetiapine, ziprasidone)</td>
<td>498 (first episode)</td>
</tr>
<tr>
<td>CAFÉ (McEvoy)&lt;sup&gt;41&lt;/sup&gt;</td>
<td>52-week randomized, double-blind, multicenter</td>
<td>Olanzapine vs quetiapine vs risperidone</td>
<td>400 (first episode)</td>
</tr>
<tr>
<td>HGAP (US Clinical Trial)&lt;sup&gt;43&lt;/sup&gt;</td>
<td>6-week double-blind</td>
<td>Olanzapine vs placebo vs risperidone</td>
<td>152 (acute phase)</td>
</tr>
<tr>
<td>HGAD (North American Clinical Trial)&lt;sup&gt;44&lt;/sup&gt;</td>
<td>6-week double-blind</td>
<td>Olanzapine vs placebo and haloperidol</td>
<td>335 (acute phase)</td>
</tr>
<tr>
<td>E003 (Eastern Hemisphere Clinical Trial)&lt;sup&gt;45&lt;/sup&gt;</td>
<td>6-week double-blind</td>
<td>Olanzapine vs haloperidol</td>
<td>431 (acute phase)</td>
</tr>
<tr>
<td>HGAJ (International Clinical Trial)&lt;sup&gt;46&lt;/sup&gt;</td>
<td>6-week double-blind</td>
<td>Olanzapine vs haloperidol</td>
<td>1996 (acute phase)</td>
</tr>
<tr>
<td>Tran et al&lt;sup&gt;58&lt;/sup&gt;</td>
<td>46-week extension maintenance phase of 3 double-blind, acute-phase trials</td>
<td>Olanzapine vs haloperidol</td>
<td>807 (maintenance)</td>
</tr>
<tr>
<td>CATIE (Lieberman et al)&lt;sup&gt;56&lt;/sup&gt;</td>
<td>18-month randomized, double-blind</td>
<td>Perphenazine vs olanzapine, quetiapine, risperidone, and ziprasidone</td>
<td>1493 (chronic schizophrenia)</td>
</tr>
<tr>
<td>SOHO (Haro et al)&lt;sup&gt;59&lt;/sup&gt;</td>
<td>3-year observational, longitudinal</td>
<td>Differences between olanzapine, risperidone, quetiapine, amisulpride, clozapine, oral typicals, and depot typicals</td>
<td>7728 (maintenance)</td>
</tr>
<tr>
<td>Breier et al&lt;sup&gt;68&lt;/sup&gt;</td>
<td>6-week randomized, double-blind</td>
<td>Olanzapine and haloperidol</td>
<td>526 (treatment resistant)</td>
</tr>
<tr>
<td>Volavka et al&lt;sup&gt;59&lt;/sup&gt;</td>
<td>14-week randomized, double-blind</td>
<td>Clozapine, olanzapine, risperidone, and haloperidol</td>
<td>157 (suboptimal treatment response)</td>
</tr>
<tr>
<td>CATIE phase 2E&lt;sup&gt;70&lt;/sup&gt;</td>
<td>18 months (both phase 1 and 2) or 6 months in phase 2</td>
<td>Clozapine, olanzapine, quetiapine, and risperidone</td>
<td>99 (discontinued CATIE phase 1/1B)</td>
</tr>
<tr>
<td>Tollefsen et al&lt;sup&gt;71&lt;/sup&gt;</td>
<td>18-week randomized, double-blind with 3-year extension</td>
<td>Olanzapine and clozapine</td>
<td>180 (treatment resistant)</td>
</tr>
</tbody>
</table>

Abbreviations: BPRS, Brief Psychiatric Rating Scale; CGI, Clinical Global Impression; PANSS, Positive and Negative Syndrome Scale.
was associated with a more favorable efficacy profile, as evidenced by considerably lower rates of discontinuation due to lack of efficacy (14–19% vs 25–48%) during treatment with olanzapine compared with other antipsychotics. A post hoc analysis examining all-cause discontinuation rates from EUFEST found that treatment with olanzapine was associated with a number needed to treat (NNT) of 4 compared with haloperidol (indicating that 4 patients have to be treated with olanzapine instead of haloperidol to prevent 1 treatment discontinuation). For comparison, the NNT for quetiapine compared with haloperidol was 9.42

Substantial improvements in several other efficacy-related outcome measures have also been observed. Greater improvements in total BPRS score have been reported with olanzapine compared with haloperidol, with patients over 2 times more likely to achieve clinical response (defined as ≥40% improvement from baseline in BPRS total score) with olanzapine (67% vs 29%, respectively; NNT=3).38 Significantly greater reductions in PANSS positive subscale scores have also been reported with olanzapine following 1 year of follow-up compared with quetiapine.41 In the EUFEST study, olanzapine provided numerical but not statistically significant differences in PANSS score reductions versus haloperidol; however, remission rates significantly favored olanzapine (57% vs 44%; NNT=8).39 Furthermore, in patients with first-episode schizophrenia, olanzapine was associated with a significantly greater reduction of hostility at 1 and 3 months.
Acute exacerbation in patients with schizophrenia

Patients suffer a variety of positive psychotic symptoms during acute exacerbations of schizophrenia, including delusions, hallucinations, and thought disorder, which may be present along with other associated behaviors (such as agitation and aggression). In addition, negative and affective symptoms may be the focus of intervention. During exacerbations, the goal of treatment is to reduce the severity of psychosis and associated symptoms as quickly as possible, and to formulate short- and long-term treatment plans to prevent relapse and slow or limit disease progression.

The initial efficacy of olanzapine in acute schizophrenia was based on data from 4 pivotal, 6-week, double-blind trials in which reductions in BPRS total scores were significantly or numerically greater with olanzapine compared to placebo and haloperidol. Subsequently, the efficacy of olanzapine in the treatment of acute exacerbations has been confirmed in several randomized, double-blind studies using the PANSS. Similar to findings in first-episode schizophrenia patients, olanzapine was also reported to be more effective than haloperidol in reducing aggressive behavior in a study that specifically enrolled physically assaultive patients with schizophrenia.

The efficacy of olanzapine in the treatment of acute exacerbations is supported by a meta-analysis involving direct and indirect comparisons of 212 randomized controlled studies of 15 first- and second-generation antipsychotics. Olanzapine was among 4 antipsychotics (which also included clozapine, amisulpride, and risperidone) determined to be significantly more effective than other choices in the acute treatment of schizophrenia, based on improvements in PANSS or BPRS scores, and among 5 antipsychotics (also including amisulpride, clozapine, paliperidone, and risperidone) associated with significantly lower rates of all-cause discontinuation. This latter finding is hypothesized to be related, at least in part, to the emergence of improved insight during olanzapine treatment and subsequent downstream effects.

Beyond acute treatment: long-term outcomes

Up to 50% of patients with schizophrenia do not adhere to treatment in the short term, and approximately 75% experience recurrent relapse and continued disability. As such, long-term treatment of schizophrenia is focused on sustaining symptom remission, maintaining (or improving) the patient’s level of functioning and quality of life, and supporting patient adherence to recommended treatment over time. The long-term effectiveness of olanzapine has been established across a number of long-term outcome domains, including relapse prevention (defined as preventing or delaying recurrence of acute exacerbations of the symptoms of schizophrenia, assessed by rates of hospitalization), robustness of response (defined as the relative absence of active schizophrenia symptoms during long-term treatment, assessed by rates of remission), and durability of therapy (defined as the relative duration of time the patient remains on any given antipsychotic, commonly assessed by all-cause discontinuation and subjective response).

One of the initial studies assessing the long-term efficacy of olanzapine was an extension study for patients who responded well to treatment in olanzapine’s pivotal acute-phase trials. Time to relapse, defined as hospitalization for psychopathology, was significantly lower for olanzapine than haloperidol, with a 1-year risk of relapse of 20% compared with 28% for haloperidol. Data from 2 large studies comparing the efficacy of first- and second-generation antipsychotics, the Clinical Antipsychotic Trials of Intervention Effectiveness (CATIE) study and the Schizophrenia Outpatient Health Outcomes (SOHO) study, consistently found that olanzapine had lower rates of hospitalization for exacerbation of schizophrenia. Further, in the CATIE study, the rate of remission, based on the Andreasen Criteria, was highest for olanzapine (12% vs 6–8% with other antipsychotics). Olanzapine also had the highest rate of remission, based on CGI scores, in the SOHO study.

In the long-term management of schizophrenia, it is important to understand the significance patients and providers place on the benefits and risks of different antipsychotics when making treatment decisions. Surveys have found that improvements in the positive symptoms of schizophrenia are ultimately more important to both healthcare providers and patients than avoiding adverse side effects of antipsychotics such as weight gain and extrapyramidal symptoms, although more severe adverse events may shift this opinion in the other direction. Likewise, in an analysis of data from 32 schizophrenia trials, discontinuation of medication was largely driven by lack of efficacy rather than intolerability. Therefore, a key finding supportive of the distinctly favorable efficacy of olanzapine is its durability of treatment. As shown in Figure 1, time to all-
cause discontinuation is consistently longer for olanzapine compared with other antipsychotics.\textsuperscript{35,40,56,59} In CATIE, only 15% of patients treated with olanzapine discontinued treatment due to lack of efficacy, as compared with 24–28% of patients treated with other antipsychotics.\textsuperscript{36} Further, a naturalistic study comparing olanzapine, risperidone, and haloperidol found better subjective response and compliance in patients treated with olanzapine compared with the other antipsychotics.\textsuperscript{64} Collectively, these data suggest that patients taking olanzapine remain on treatment longer, despite the drug’s known side effects, than do patients taking other antipsychotics, potentially owing to olanzapine’s superior efficacy.

Furthermore, post hoc analysis of data from the CATIE study identified a significant difference in favor of olanzapine for hostility in schizophrenia,\textsuperscript{36} consistent with the post hoc analysis of hostility in first-episode patients from the EUFEST study.\textsuperscript{37}

### Suboptimal treatment response

Treatment resistance, commonly defined as little or no symptomatic response to at least 2 previous antipsychotics, occurs in approximately 10–30% of patients with schizophrenia.\textsuperscript{11} This definition is used as the criteria for recommending treatment with clozapine.\textsuperscript{65} It is also not uncommon for patients to experience partial response to a prescribed antipsychotic with a clinical status that lies somewhere between full responder and treatment resistance.\textsuperscript{65,66} The management of these patients is a major challenge and constitutes a serious public health concern given the associated social and economic burden.\textsuperscript{67}

The use of olanzapine in patients with treatment-resistant schizophrenia or suboptimal response has been evaluated in several studies.\textsuperscript{68–74} In a study of patients with suboptimal treatment response randomized to olanzapine, clozapine, risperidone, or haloperidol, improvements in global psychopathology were significantly greater at week 14 with olanzapine compared with haloperidol.\textsuperscript{59} In phase 2E of the CATIE study, which included patients who discontinued from phase 1 primarily due to lack of efficacy, no significant differences were found between clozapine and olanzapine with regard to all-cause discontinuation or improvements in PANSS total score at 3 months, with clozapine being superior to treatment with either quetiapine or risperidone.\textsuperscript{20} Other studies in patients with treatment-resistant schizophrenia or suboptimal treatment response have also reported similar reductions with olanzapine and clozapine in PANSS total score, CGI severity score, and BPRS total score.\textsuperscript{69,71–73}

In many of the aforementioned studies, olanzapine dosages above the maximum recommended dose of 20 mg/day were used in treatment-resistant patients.\textsuperscript{13} This is not without controversy, as doses of 40 mg/day did not confer an advantage over lower doses in a large (N=599) randomized clinical trial of patients with suboptimal response to their current treatment (but who were not defined as treatment resistant).\textsuperscript{75}

### Safety

The safety profile of olanzapine has been reviewed extensively elsewhere.\textsuperscript{3,76–79} In brief, second-generation antipsychotics are generally better tolerated than are the first-generation antipsychotics with regard to antipsychotic-induced movement disorders.\textsuperscript{33} However, many of these drugs are associated with clinically significant metabolic disturbances, including increased risk of diabetes, dyslipidemia, and weight gain.\textsuperscript{80} Olanzapine has one of the highest risks for metabolic syndrome among the second-generation atypical antipsychotics, especially during long-term treatment and with the administration of high doses, in accordance with a report from professional associations of psychiatrists, endocrinologists, and diabetologists.\textsuperscript{81} In particular, hypertriglyceridemia can represent a sensitive marker of insulin resistance, a condition that induces obesity, especially central obesity, a primary risk factor for metabolic syndrome.\textsuperscript{82} In a recent animal model study, olanzapine exacerbated atherosclerosis by deregulating hepatic lipid metabolism and worsening hyperlipidemia and aortic inflammation.\textsuperscript{83} Antagonistic effects of olanzapine at 5-HT\textsubscript{2c}, histaminergic H1, and muscarinic M3 receptors have been implicated in olanzapine-induced weight gain. Although a 2010 meta-analysis\textsuperscript{84} reported significantly less weight gain when adjunctive metformin, fenfluramine, sibutramine, topiramate, and reboxetine versus placebo were added on to olanzapine therapy, and more recent studies support the efficacy of betahistine,\textsuperscript{85–88} to date, data are insufficient to recommend the routine use of any of these medications to mitigate olanzapine-induced weight gain.\textsuperscript{84,85}

Olanzapine’s adverse effects on glucose and lipid metabolism, weight, and body composition are of serious concern and require close monitoring.\textsuperscript{77} In an analysis of 13 short-term, placebo-controlled olanzapine monotherapy studies (median exposure: 6 weeks), the average weight gain with olanzapine was 2.6 kg (5.7 lb) compared to a
weight loss of 0.3 kg (0.6 lb) with placebo. The percentage of patients who gained at least 7% of their baseline weight was 22% with olanzapine versus 3% with placebo. In long-term studies (at least 48 weeks), 64%, 32%, and 12% of patients gained 7%, 15%, or 25% or more of their baseline body weight, respectively. Shifts from normal/borderline to abnormal plasma glucose or lipid variables have also been observed and are a concern. Several guidelines and publications have included strategies to help mitigate these side effects, such as with concomitant off-label use of metformin, as well as greater attention to promoting positive lifestyle modifications with respect to diet and exercise. However, there are currently no FDA-approved treatments indicated to mitigate the metabolic dysregulation, including weight gain, associated with olanzapine or any other currently available atypical antipsychotics.

Despite these well-known adverse metabolic effects, olanzapine does have some safety advantages relative to other antipsychotics. Olanzapine is associated with a lower incidence of extrapyramidal symptoms than typical and some atypical antipsychotics, an important cause of noncompliance and treatment discontinuation. Further, agranulocytosis, a potentially fatal adverse event that limits clozapine’s use, was not reported in any of the more than 2000 patients receiving olanzapine in the 4 pivotal efficacy and safety trials. Olanzapine also has generally little impact on prolactin compared with some typical and atypical antipsychotics. A significantly lower proportion of patients had elevations in prolactin during treatment with olanzapine compared with haloperidol (18–35% vs 82%) or with risperidone (51% vs 94%). Although prolactin effects with olanzapine are likely dose related, increases in prolactin associated with olanzapine were generally mild, transient, and of lesser magnitude than haloperidol and risperidone. Several studies have subsequently confirmed a favorable impact of olanzapine on prolactin levels. As a result, patients taking olanzapine may be at reduced risk for complications of hyperprolactinemia, including gynecomastia, galactorrhea, menstrual irregularities, and sexual dysfunction, compared with other antipsychotics. Lastly, olanzapine treatment was associated with a low incidence of seizures, 22/2500 (0.9%), during premarketing studies. Based on this finding, the Epilepsy Foundation lists olanzapine as 1 of 3 antipsychotics recommended on the basis of both epilptogenesis and side effect profile.

Summary

Two decades ago, the safety risks associated with olanzapine were not as well understood as they are today. Now, olanzapine is no longer the first choice of treatment for patients with schizophrenia. Even so, olanzapine remains widely used, lending indirect support to the hypothesis that olanzapine’s effectiveness differs from other antipsychotic treatment options. This commentary aims to highlight the significant body of data supporting the hypothesis that olanzapine may be more effective than other non-clozapine antipsychotics currently considered the first choice of treatment for schizophrenia. Although our focus was to highlight olanzapine’s unique efficacy qualities, we acknowledge that olanzapine’s efficacy is not a universal finding across all studies; however, for the purposes of the current commentary, it was not our intention to conduct an exhaustive meta-analysis. The relative efficacy seems to depend, in part, on the clinical situation in which it is used. Collectively, these data suggest that if efficacy alone were the only consideration, olanzapine may be a preferred first-line treatment option for patients with chronic schizophrenia requiring effective long-term treatment (ie, for prevention of relapse, robustness of response, and treatment persistence aspects) and in those with suboptimal responses during treatment with other antipsychotics. Despite the known risk for metabolic side effects, including weight gain, olanzapine continues to be prescribed due to the overall favorable drug profile. As with any treatment, selection of the most appropriate antipsychotic agent for patients with schizophrenia must be individualized and thoughtfully considered, taking into account both the benefits and risks associated with its use. Therefore, the development of interventions that offer the efficacy of olanzapine but reduce the weight gain and potential long-term complications associated with its use could address a major unmet need in the treatment of schizophrenia. These interventions could include agents with new mechanisms of action or single-pill combination products that contain both olanzapine and agent(s) that could potentially mitigate olanzapine’s propensity for weight gain to serve as potential alternatives in clinical situations in which olanzapine would have been considered.

Abbreviations

BPRS, Brief Psychiatric Rating Scale; CATIE, Clinical Antipsychotic Trials of Intervention Effectiveness; CGI, Clinical Global Impression; EUFEST, European First-
Episode Schizophrenia Trial; FDA, US Food and Drug Administration; NNT, number needed to treat; PANSS, Positive and Negative Syndrome Scale; SOHO, Schizophrenia Outpatient Health Outcomes.

Acknowledgments
This study was sponsored by Alkermes, Inc. Medical writing and editorial support was provided by Peloton Advantage, LLC, an OPEN Health company, Parsippany, NJ, and funded by Alkermes, Inc. Alkermes, Inc. is developing a combination product of olanzapine and samidorphan for the treatment of schizophrenia, and has funded the development of this manuscript.

Disclosures
LC has, in the past 5 years, engaged in collaborative research with, or received consulting or speaking fees, from: Acadia, Alexza, Alkermes, Allergan, AstraZeneca, Avanir, Bohringer Ingelheim, Bristol-Myers Squibb, Eli Lilly, Forum, Genentech, Intra-Cellular Therapeutics, Janssen, Jazz, Lundbeck, Merck, Medivation, Mylan, Neurocrine, Novartis, Noven, Otsuka, Pfizer, Indivior/Reckitt Benckiser, Reviva, Shire, Sunovion, Takeda, Teva, Valeant, Vanda. In the past 12 months, consultant: Acadia, Alkermes, Allergan, Indivior, Intra-Cellular Therapeutics, Janssen, Lundbeck, Merck, Neurocrine, Noven, Otsotica, Otsuka, Pfizer, Shire, Sunovion, Takeda, Teva, Vanda. In the past 12 months speaker: Acadia, Alkermes, Allergan, Janssen, Lundbeck, Merck, Neurocrine, Otsotica, Pfizer, Shire, Sunovion, Takeda, Teva. Stocks (small number of shares of common stock): Bristol-Myers Squibb, Eli Lilly, J & J, Merck, Pfizer purchased >10 years ago. Royalties: Wiley (Editor-in-Chief, International Journal of Clinical Practice), UpToDate (reviewer), Springer Healthcare (book). JPM serves on advisory boards for Neurocrine and Alkermes, MST, DM, and PJW are full-time employees of Alkermes, Inc., and may own stock in the company.

References
6. Davis JM, Chen N, Glick ID. A meta-analysis of the efficacy of second-generation antipsychotics. Arch Gen Psychiatry. 2003;60(6):553–564. doi:10.1001/archpsyc.60.6.553


