Long-term Safety of Asenapine in Pediatric Patients Diagnosed With Bipolar I Disorder: A 50-Week Open-Label, Flexible-Dose Trial

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Abstract
Background Sublingually administered asenapine was approved in March 2015 by the United States Food and Drug Administration for patients aged 10–17 years with an acute manic or mixed episode associated with bipolar I disorder (BP-1). This is the first long-term safety and tolerability study of asenapine in this population.
Methods Following the 3-week randomized, double-blind, placebo-controlled trial of patients aged 10–17 years with an acute manic or mixed episode associated with BP-1, patients could enroll in this flexible-dose (2.5–10 mg twice daily) open-label extension (OLE) study for an additional 50 weeks, conducted from August 2011 to September 2014 in the United States and Russia. Treatment-emergent adverse events (TEAEs) were assessed and predefined TEAEs of interest reported in addition to metabolic and anthropometric parameters. The Young Mania Rating Scale (YMRS) and Clinical Global Impressions scale in bipolar illness (CGI-BP) were used to assess effectiveness.
Results A total of 321 patients (lead-in study treatment: placebo, n = 80; asenapine, n = 241) were included; 267 (83.2 %) reported one or more TEAE and 181 (56.4 %) discontinued early, 48 (15.0 %) due to TEAEs. Of the predefined TEAEs of interest, combined somnolence/sedation/hypersomnia occurred most frequently (42.4 %) followed by oral hypoesthesia/dysgeusia (7.5 %). In total, 109 (34.8 %) patients experienced clinically significant weight gain (≥7 % increase). No clinically meaningful changes were noted for laboratory parameters measured. Eighteen patients met the criteria for new-onset metabolic syndrome (MBS) post-baseline during the extension study, whereas 10 patients who met MBS criteria at baseline did not meet MBS criteria at endpoint. A total of 12 patients met MBS at baseline and endpoint. Mean change in YMRS total score from OLE baseline was −9.2 points at week 50, and change in CGI-BP severity overall score was similar among all treatment groups (those who initially received asenapine and those who initially received placebo). After 26 weeks of treatment in the OLE, 79.2 % of patients were classified as YMRS 50 % responders relative to acute trial baseline.
Conclusions Asenapine was generally well tolerated in pediatric patients with BP-1 during ≤50 weeks of open-label treatment; among predefined TEAEs of interest, the combination of somnolence/sedation/hypersomnia was the most common.

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Key Points

Long-term flexible dose administration of sublingual asenapine was generally well tolerated in pediatric patients with a manic or mixed episode associated with bipolar I disorder.

Fifteen percent of patients discontinued due to treatment-emergent adverse events; clinically significant weight gain was experienced by 34.8% of patients.

Among the predefined treatment-emergent adverse events of interest, the combination of somnolence, sedation, and hypersomnia was most frequent (42.4% of patients).

1 Introduction

Many patients with bipolar I disorder (BP-1) experience onset of mania prior to adulthood, and evidence suggests that BP-1 is relatively common in pediatric-aged patients; a cross-sectional survey of more than 10,000 American adolescents aged 13–18 years identified in household and school settings estimated that the lifetime prevalence of BP-1 was 1.7% [1]. These data are similar to results of an earlier meta-analysis of more than 16,000 young people aged 7–21 years, which estimated the lifetime prevalence of BP-1 to be 1.8%, with no difference observed between US and non-US populations [2]. Freeman and colleagues [3] reported that the quality of life associated with pediatric bipolar disorder is more burdensome than a variety of physical conditions and appears to have a greater impact than other pediatric psychiatric conditions.

Individuals who experience BP-1 earlier in life are expected to develop a more chronic, severe, and recurrent course of the disease when compared with individuals who developed the disorder in adulthood [4–7]. Bipolar disorder is a major cause of disability in young people [8].

Asenapine is a sublingually administered second-generation atypical antipsychotic. In the United States, asenapine was approved in adults for use as monotherapy (August 2009) or adjunctive therapy with lithium or valproate (September 2010) for manic or mixed episodes associated with BP-1. In March 2015, asenapine was approved by the US Food and Drug Administration (FDA) for use as monotherapy in pediatric patients aged 10–17 years with an acute manic or mixed episode associated with BP-1 [9]. Asenapine is not approved for the treatment of pediatric patients with BP-1 outside of the US, but is approved or registered in 65 countries as monotherapy for the treatment of adults with BP-1.

Because BP-1 is a chronic illness and a serious condition, long-term treatment is usually necessary. There are presently no long-term data available on the safety and tolerability of asenapine in pediatric patients with BP-1 of which we are aware. In general, there are limited empirical data that describe the long-term safety of any second-generation antipsychotics in pediatric patients. Completed studies have shown evidence that sedation, hyperprolactinemia, and metabolic abnormalities may develop during pharmacotherapy with this class of medications [10]. Therefore, in this study, we sought to characterize the safety of asenapine over 50 weeks in an open-label extension (OLE) trial.

2 Methods

Details of the antecedent 3-week, randomized, double-blind, placebo-controlled efficacy and safety trial of asenapine in pediatric patients with an acute manic or mixed episode associated with BP-1 (NCT01244815) have been previously published [11]. In brief, patients were randomized 1:1:1:1 to placebo, asenapine 2.5, 5, or 10 mg twice daily (bid) for 3 weeks (see electronic supplementary material [ESM], Online Resource 1). After completing the acute trial, patients could enroll in this 50-week OLE trial. The first day of the OLE trial overlapped with day 21/end-of-treatment period for the acute mania trial. The OLE was a flexible-dose, multicenter trial to evaluate the long-term safety and tolerability of asenapine monotherapy in pediatric patients aged 10–17 years with a manic or mixed episode associated with BP-1 (NCT01349907); the trial was conducted from August 16, 2011 to September 5, 2014, in the United States and Russia. Diagnosis of BP-1 was made according to the Diagnostic and Statistical Manual of Mental Disorders, fourth edition, text revision (DSM-IV-TR) with current manic (296.4x) or mixed (296.6x) episode with or without psychotic features required and confirmed by a structured clinical interview which consisted of a medical and psychiatric history, DSM-IV-TR checklist, and the Kiddie-Schedule for Affective Disorders and Schizophrenia—Present and Lifetime Version at screening before enrollment in the acute trial.

Independent Ethics Committees associated with each study site reviewed and approved the protocol and applicable amendments. The study was conducted in accordance with Good Clinical Practice standards and applicable country and/or local statutes and regulations regarding ethical committee review, informed consent, and the protection of human patients participating in biomedical research. Written informed consent was provided by the patient’s parent, legal representative, or caretaker and
written patient assent was also obtained before any trial procedures or assessments were performed. If at any time during the patient’s participation they turned 18 years of age, they were asked to provide written informed consent. This was an open-label trial and therefore did not require the use of blinding procedures although the blind was maintained with respect to the acute phase trial until that database was unlocked. An external Data Monitoring Committee met regularly to evaluate safety data in an unblinded fashion during the course of the trial. The OLE was initially designed as a 26-week study and amended to 50 weeks to meet European Medicines Agency requirements. The protocol was also amended to include subjects of younger age (10 and 11 years) and older age (18 years of age). Although patients aged 18 years were permitted to enroll in the OLE, predefined analyses were limited to ages ≤17 at OLE entry; the one patient who was aged 18 at entry was therefore excluded from analyses.

2.1 Inclusion Criteria

Patients aged 10–18 years who both completed the acute trial and who, in the investigator’s opinion, could potentially benefit from long-term asenapine treatment were eligible for enrollment. Key inclusion criteria from the acute trial included a primary diagnosis of BP-1 according to the DSM-IV-TR with a current manic (296.4x) or mixed (296.6x) episode with or without psychotic features (confirmed by a structured clinical interview that consisted of a medical and psychiatric history, DSM-IV-TR checklist, and the Kiddie-Schedule for Schizophrenia and Affective Disorders—Present and Lifetime Version at screening), a Young Mania Rating Scale (YMRS) score ≥20, and an overall severity of bipolar mania and depression ≥4 on the Clinical Global Impressions scale in bipolar illness (CGI-BP). For enrollment in the OLE, patients were required to demonstrate an acceptable level of adherence to antecedent trial medication, study visits, and other protocol requirements in the acute trial according to the investigators’ opinion, and to have a caregiver living with them able to provide continued support for adherence to the study protocol.

2.2 Exclusion Criteria

Patients with a pervasive development disorder, schizophrenia, schizoaffective disorder, post-traumatic stress disorder, obsessive–compulsive disorder, or psychosis due to a medical condition or prohibited concomitant medication were excluded from the acute trial. Patients with a positive urine pregnancy test at OLE baseline or with the intention to become pregnant during the trial were excluded. Additional criteria that warranted exclusion from the OLE included being at imminent risk of self-harm or harm to others, in the investigator’s opinion, based on clinical interview and responses provided on the Columbia Suicide Severity Rating Scale (C-SSRS), involuntary inpatient commitment, poor adherence to the acute trial protocol, and known serological evidence of human immunodeficiency virus antibody. In addition, patients with an uncontrolled, unstable, clinically significant medical condition were excluded. Prohibited concomitant medications included all psychotropic drugs with the exception of stimulants (for attention-deficit hyperactivity disorder [ADHD]) and short-acting benzodiazepines (for transient agitation, irritability, restlessness, insomnia, and hostility). Anticholinergics were also allowed as a rescue medication for extrapyramidal symptoms (EPS).

2.3 Dosing

All patients, whether treated with asenapine or placebo in the acute trial, were administered open-label asenapine formulated as a fast-dissolving, black-cherry-flavored sublingual tablet at 2.5 mg bid on day 1. On the evening of day 4, the asenapine dose was increased to 5 mg bid. At the day 7 visit, the asenapine dose was increased to 10 mg bid, if tolerated, beginning with the evening dose. The dose was increased sooner, at the discretion of the investigator, if clinically warranted. From day 7 onward, asenapine dosing was flexible based on tolerability and/or symptomatology. Study visits also occurred on days 14, 28, 42, 70, 98, 126, 154, 182, 210, 238, 266, 294, 322, and 350 (see ESM, Online Resource 1).

2.4 Safety Outcomes

There were no prespecified key safety endpoints; exploratory safety events and tolerability parameters were assessed throughout the trial (i.e., vital signs were recorded at every study visit; laboratory tests were administered at visits 14, 18, 20, 25, and 28 [i.e., days 28, 126, 182, 266, 350]; and anthropometric parameters were assessed at visits 13, 14, 16, 18, and 20 through 28 [i.e., days 14, 28, 70, 126, and 182–350]). Adverse events were recorded based on spontaneous report or investigator observation; reported adverse events were monitored with the relevant clinical assessments and/or laboratory tests. An adverse event was considered treatment emergent (TEAE) if it was newly reported after open-label baseline or reported to have worsened in severity since open-label baseline. Predefined TEAEs of interest in the study were akathisia, dizziness, insomnia, somnolence/hypersomnia/sedation combined, oral hypoesthesia/dysgeusia, EPS, and incidence of clinically significant weight increase (defined as ≥7 % increase from baseline to endpoint according to the US FDA.
guidance and product prescribing information) [9]. The Extrapyramidal Symptom Rating Scale (ESRS) was used to measure EPS [12]. The ESRS consisted of scores on EPS-related items, leading to nine derived parameter scores. Ratings were made through a combination of clinical interview and motor examination. Further, events related to suicidality (e.g., suicidal ideation, suicidal behavior, and/or non-suicidal self-injurious behavior) were summarized based on responses to the C-SSRS at each visit [13].

Analysis of safety data also focused on mean changes from the OLE baseline in the following laboratory parameters: fasting glucose, fasting triglycerides, fasting cholesterol, prolactin, fasting insulin, and glycosylated hemoglobin (HbA1c). Laboratory values were obtained at baseline and on days 28, 126, 182, 266, and 350. TEAEs potentially related to new-onset diabetes mellitus were analyzed by grouping for summary purposes hyperglycemia, new-onset diabetes mellitus, increased weight, increased appetite, increased blood glucose, central obesity, hyperlipidemia, and loss of consciousness as defined in the Medical Dictionary for Regulatory Activities (MedDRA) by the standardized MedDRA query (SMQ) ‘hyperglycemia/new onset diabetes (broad)’. The criteria for metabolic syndrome (MBS) was defined by the International Diabetes Federation, which includes abdominal girth (measured throughout the trial, in centimeters) plus two other laboratory values (triglycerides, high-density lipoprotein cholesterol, blood pressure, or glucose), all of which had to be met at the same visit for a MBS diagnosis [14, 15]. MBS was further categorized based on the study period (i.e., baseline and/or treatment periods).

2.5 Effectiveness Outcomes

The following select exploratory parameters were assessed during the study: change in YMRS total score from baseline(s) (pre-treatment and open-label); time to failure to maintain effect from open-label baseline to any time point (i.e., a <50 % reduction from baseline in YMRS total score for patients who achieved a 50 % YMRS response during the acute trial and lost the effect during the OLE); total YMRS 50 % responders and remitters (i.e., those who achieved ≥50 % improvement in YMRS total score from each respective trial baseline to endpoint were considered responders, and remitters required a YMRS total score ≤12); and change in CGI-BP severity scale overall score from OLE baseline. The YMRS is an 11-item clinician-rated scale to assess the severity and symptoms of manic episodes and can range from 0 (all symptoms absent) to 60 (all symptoms extreme) [16–18]. A decrease in YMRS total score is indicative of improved symptoms. The YMRS was administered at baseline and on days 28, 70, 126, 182, 266, and 350. The CGI-BP severity scale is a single-item clinician-rated scale with a possible score of 1–7, where higher scores indicate greater severity [19]. The CGI-BP severity scale was administered on days 7, 14, 28, 42, 70, 98, 126, 154, 182, 210, 238, 266, 294, 322, and 350.

2.6 Sample Size

No formal sample size calculation was performed because this was a single-arm OLE trial and the primary objective was evaluation of safety and tolerability. The number of patients enrolled in this trial was dependent on the number of patients completing the double-blind acute trial and was expected to be ≥280 patients (approximately 70 % of patients enrolled in the acute trial).

2.7 Statistical Analysis

Analysis of safety data used the all-patients-as-treated (APaT) population, defined as all patients ≤17 years old who received one or more dose of open-label trial medication. For analysis of safety parameters from the OLE baseline (or last day of the antecedent acute trial treatment), categorical variables were summarized as numbers and percentages. For continuous variables (such as changes from baseline), mean, median, minimum, and standard deviation (SD) were provided as summary measures. Additionally, within-group point estimates were provided. For the select TEAEs of interest and TEAEs that occurred in four or more patients overall, 2-sided 95 % confidence intervals (CIs) were provided for the within-group percentage of patients with events using the 2-tailed exact binomial method proposed by Clopper–Pearson [20]. For continuous measures, 2-sided 95 % CIs were provided for within-group differences using an analysis of covariance model with terms for pooled site, identified group, and baseline value (if applicable). Predefined limits of change were summarized using frequency tables for each visit and endpoint (if applicable) and shifts from baseline to post-baseline and endpoint (if applicable) by identified group.

Changes in body weight and body mass index (BMI) were also adjusted for growth (age and sex) and presented as mean Z-scores [21]. The Z-scores are assumed to follow a standard normal distribution and indicate how many SDs an observed value is away from the expected growth (i.e., weight, height, BMI), based on an individual’s age and sex. Each Z-score corresponds to a percentile in the standardized growth curve. A Z-score of 0 corresponds to a 50th percentile (the median), whereas a negative Z-score corresponds to a percentile below the median and a positive Z-score corresponds to a percentile above the median. The distance of a Z-score from 0 represents the distance of a percentile from the median. For the countries included in this trial (USA and Russia), age- and sex-adjusted growth
was compared with expected growth seen in the general US population using the most recent growth charts at the time of database lock as provided by the National Center for Health Statistics (NCHS) [21]. A SAS program is available from the NCHS website that provides Z-scores and percentiles for weight, height, and BMI based on the US Centers for Disease Control and Prevention growth charts [21]. The Centers for Disease Control and Prevention reference database based on the US population may not be representative of sites outside the United States.

Efficacy analysis used the full analysis set population, defined as patients <17 years old who received one or more dose of open-label trial medication and had one or more open-label postbaseline in-treatment YMRS total score. Efficacy results (YMRS total score, CGI-BP severity overall score, and total YMRS 50 % responders) were evaluated using last observation carried forward (LOCF) and observed cases (OC) approaches. For LOCF analysis, the value of a particular assessment was the last available non-missing value prior to the particular assessment when the value was missing. In the OC approach, the assessment closest to the scheduled assessment day within the allowed window was used. If there were two assessments within the same absolute distance of the scheduled assessment, the one with the later date was chosen as OC. Kaplan–Meier analysis was used to estimate the time to failure to maintain response, and was limited to those patients at endpoint in the acute trial who were considered total YMRS 50 % responders. Days to failure to maintain response was defined as the number of days from first achieving responder status to the first subsequent time that the patient had less than a 50 % reduction from open-label baseline in YMRS total score. A patient was considered a YMRS responder at a given visit if they experienced a 50 % or more decrease from acute trial baseline in YMRS total score. A patient was considered a YMRS responder status to the first subsequent time that the patient had less than a 50 % reduction from open-label baseline in YMRS total score. A patient was considered a YMRS responder at a given visit if they experienced a 50 % or more decrease from acute trial baseline in YMRS total score. A patient was considered a YMRS responder status to the first subsequent time that the patient had less than a 50 % reduction from open-label baseline in YMRS total score. A patient was considered a YMRS responder status to the first subsequent time that the patient had less than a 50 % reduction from open-label baseline in YMRS total score. A patient was considered a YMRS responder status to the first subsequent time that the patient had less than a 50 % reduction from open-label baseline in YMRS total score.

3 Results

3.1 Patients

A total of 350 patients completed the acute trial and 322 (92.0 %) were treated with open-label asenapine; 321 were included in the analyses (Fig. 1). As predefined analyses were limited to patients aged <17 years at OLE entry, the one patient aged 18 years at entry was excluded from these analyses. Of the 321 included patients, 80 had been treated with placebo and 241 treated with asenapine in the antecedent acute phase trial. The mean age was 13.8 ± 2.0 years for the overall OLE population (Table 1). Other baseline demographic characteristics were similar between groups. The majority of patients in the total treatment group were White (69.8 %) and had a mean BMI of 24.6 ± 6.3. The most common comorbid Axis I disorder among the overall patient population was ADHD, which was diagnosed in 182 (56.7 %) patients, of whom 88 (48.4 %) were prescribed stimulants and 94 (51.6 %) were not. For the total treatment group, overall asenapine exposure ranged from 2 to 380 days, with a median duration of 182 days of treatment. Overall, 107 (33 %) patients were exposed to asenapine 2.5 mg, 5 mg, or 10 mg bid for at least 230 days. In the overall study population, 31 (9.7 %) patients received a modal dose of asenapine 2.5 mg bid, 105 (32.7 %) patients received 5 mg bid, and 170 (53.0 %) patients received 10 mg bid. There were 15 (4.7 %) patients that received treatment for <8 days for which no modal dose was determined.

Concomitant medication use was reported by 69.8 % of patients in the APaT population. The most commonly used medications were propionic acid derivatives (18.4 %; e.g., ibuprofen), sympathomimetics (16.5 %; e.g., methylphenidate), benzodiazepine derivatives (12.5 %; e.g., lorazepam), anilides (12.1 %; e.g., paracetamol), and mixed amphetamine salts (10.3 %).

3.2 Safety

No deaths occurred during the trial (Table 2). Overall, 267 (83.2 %) patients reported TEAEs in the total treatment group. Serious adverse events (SAEs) were reported in a total of 22 (6.9 %) patients. Most SAEs belonged to the psychiatric disorders class (i.e., suicidal ideation, aggression, bipolar disorder, depression, agitation). Overall, 13 (4.0 %) patients in the total treatment group discontinued treatment as a result of SAEs. The SAEs that led to discontinuation among these 13 patients included ADHD, aggression (n = 2), agitation, anxiety, bipolar disorder, depression, dystonia, intentional overdose, suicidal behavior, and suicidal ideation (n = 3). Additionally, 181 (56.4 %) patients discontinued from the total treatment group; the most common reasons were attributed to AEs (48, 15.0 %), nonadherence to the study protocol (46, 14.3 %), and consent withdrawal (38, 11.8 %). The most common individual TEAEs leading to discontinuation in the overall population were somnolence (1.9 %), sedation (1.2 %), and fatigue (1.2 %). Per the C-SSRS, 295 (91.9 %) of the 321 patients reported no positive responses (i.e., affirmative responses to questions on the C-SSRS) and one patient did not have C-SSRS data. Of the 25 (7.8 %) patients with positive responses on the C-SSRS, 23 (7.2 %) had some degree of suicidal ideation and/or behavior per the C-SSRS. The majority of the suicidal ideation was reported as passive or non-specific, or ideation without a plan or intent. A total of 8 of the 25 (32.0 %) patients with positive C-SSRS
responses reported self-injurious behavior with no suicidal intent on the C-SSRS. Further, 19 (5.9%) patients in the total treatment group had AEs or TEAEs of suicidality, including 10 patients for which these were considered serious. Of these 10 patients with serious TEAEs related to suicidality, one patient had a suicide attempt (melatonin overdose). Further, three patients discontinued due to suicidal ideation and two for suicidal behavior. During the trial, five accidental overdoses associated with asenapine were recorded as TEAEs; overdose was defined as any dose greater than the daily dose specified in the protocol.

Of the predefined TEAEs of interest (Table 2), the combination of somnolence, sedation, and hypersomnia occurred most frequently (136 or 42.4% of the total treatment group; 60.0% of placebo/asenapine patients and 36.5% of asenapine/asenapine patients) followed by the combination of oral hypoesthesia and dysgeusia (7.5% overall), EPS (6.2% overall), and dizziness (4.7% overall). With the exceptions of insomnia and ≥7% weight gain from baseline to endpoint, which were reported with similar frequency between the OLE groups, as expected the placebo/asenapine group had a greater frequency of all other TEAEs of interest compared with the asenapine/asenapine group. Most TEAEs were reported to be mild or moderate in severity. The most commonly reported TEAEs that were reported in ≥5% of the total treatment group were somnolence, increased weight, sedation, headache, nausea, vomiting, fatigue, increased appetite, upper abdominal pain, and oral hypoesthesia (see ESM, Online Resource 2).

Mean increases in all growth parameters (weight, height, BMI, and abdominal girth) were observed; however, relative increases in weight exceeded those in height (children and adolescents are expected to gain weight and grow in height as part of normal development) as suggested by the observed increases in BMI. In the total treatment group, 109 (34.8%) patients experienced clinically significant (≥7%) weight gain (Table 2). TEAEs of weight gain were reported for 58 (18.1%) patients. After correcting for age and sex, the mean weight and BMI scores at baseline and endpoint demonstrated an increase (Table 3). Weight category shifts were observed from the OLE baseline; more patients had upward shifts than downward shifts (see ESM, Online Resource 3). The mean change from baseline in weight percentile increased over time during the acute trial, and continued to increase initially during the OLE and appeared to stabilize temporally with continued treatment (see ESM, Online Resource 4). Results indicate that positive mean changes in BMI percentiles did not increase after week 18 (mean BMI percentile change of 3.9 at day 266) (see ESM, Online Resource 5). This is consistent with the corresponding weight percentile, where the maximum mean change in weight percentile occurred at week 18 (4.9 at day 266).

No clinically meaningful changes were noted in the laboratory parameters measured (Table 4). There were no patients with reported TEAEs with the specific preferred terms of hyperglycemia or new-onset diabetes. However, using the broad MedDRA definition, additional terms were included in the definition of new-onset diabetes mellitus (increased weight, increased appetite, increased blood glucose, central obesity, hyperlipidemia, and loss of consciousness). In the total treatment group, 73 (22.7%) patients reported TEAEs within the broad scope of those pertaining to new-onset diabetes mellitus. Additionally,
there were two TEAEs of increased blood glucose and six cases of increased blood insulin; all of which were of mild or moderate intensity. In total, 18 patients met new-onset MBS criteria post-baseline during the extension study, whereas 10 patients who met MBS criteria at baseline did not meet MBS criteria at endpoint. A total of 12 patients met MBS at baseline and endpoint.

### 3.3 Efficacy

Change in YMRS total score from the acute trial baseline in this uncontrolled trial indicates that the improvement in mania, as measured by the mean change in YMRS total score from baseline, was maintained over the course of the extension trial for patients who were treated with asenapine in the acute trial (Fig. 2a). Mean change in YMRS total score from OLE baseline at week 26 was −6.9 points in the total treatment group and was of a greater magnitude in the placebo/asenapine group (−13.0 points) versus the asenapine/asenapine group (−4.9 points). Week 50 results for total YMRS score were similar with a mean change of −9.2 points observed in the total treatment group, −15.2 in the placebo/asenapine group, and −6.5 in the asenapine/asenapine group. Similar trends over time were observed with the LOCF approach.

Likewise, the change in CGI-BP severity overall score was similar among all treatment groups, indicating effectiveness was maintained over the course of the trial (Fig. 2b). The LOCF approach yielded consistent results.

Of the 141 patients who were total YMRS 50% responders at the end of the acute trial and who were included in the OLE, 46 patients (32.6%) failed to
maintain effect in the OLE. At day 182 (week 26), the Kaplan–Meier estimate of the cumulative percentage of failures to maintain effect was 31.1 % in the total treatment group.

After 26 weeks of treatment in the OLE, a total of 118 of 149 (79.2 %) patients were total YMRS 50 % responders using the acute trial baseline; YMRS remission (YMRS total score ≤12) was reached by 102 of 149 (68.5 %) patients at the end of treatment in the OLE. Of note, 26-week data were used owing to the limited number of patients with data values after week 26.

4 Discussion

Flexibly dosed asenapine (2.5 mg, 5 mg, and 10 mg bid) was generally well tolerated in pediatric patients with an acute manic or mixed episode associated with BP-I treated

| Table 2 | Summary of select TEAEs\(^a\) in pediatric patients with bipolar I disorder |
|---------|----------------------------------------------------------|
|         | Placebo/asenapine | Asenapine/asenapine | Total |
|         | \(N = 80\) | \(n (\%)\) | \(N = 241\) | \(n (\%)\) | \(N = 321\) | \(n (\%)\) |
| Any TEAE\(^b\) | 74 (92.5) | 193 (80.1) | 267 (83.2) |
| Death | 0 | 0 | 0 |
| SAE | 5 (6.3) | 17 (7.1) | 22 (6.9) |
| Discontinuation caused by TEAE | 11 (13.8) | 37 (15.4) | 48 (15.0) |
| \(\geq 1\) predefined TEAE of interest during the study\(^b\) | | | |
| Somnolence, sedation, and hypersomnia combined | 48 (60.0) | 88 (36.5) | 136 (42.4) |
| Dizziness | 9 (11.3) | 6 (2.5) | 15 (4.7) |
| Insomnia | 2 (2.5) | 6 (2.5) | 8 (2.5) |
| Oral hypoesthesia and dysgeusia | 15 (18.8) | 9 (3.7) | 24 (7.5) |
| EPS\(^c\) | 6 (7.5) | 14 (5.8) | 20 (6.2) |
| Akathisia | 3 (3.8) | 7 (2.9) | 10 (3.1) |
| \(\geq 7\) % weight increase from baseline to endpoint, \(n/n1\) (\%) | 28/75 (37.3) | 81/238 (34.0) | 109/313 (34.8) |

\(\text{EPS}\) extrapyramidal symptom, \(n1\) number of patients available for assessment, \(\text{OLE}\) open-label extension, \(\text{SAE}\) serious adverse event, \(\text{TEAE}\) treatment-emergent adverse event

\(a\) TEAEs were counted from the open-label baseline; TEAEs were newly reported in the OLE or worsened in severity during the OLE compared with the acute phase

\(b\) Every patient was counted a single time for each applicable row and column

\(c\) Includes the preferred terms akathisia, dyskinesia, dystonia, and Parkinson-like events

| Table 3 | Z-scores\(^a\) for weight and BMI asenapine-treated pediatric patients with bipolar I disorder |
|---------|----------------------------------------------------------|
|         | Placebo/asenapine | Asenapine/asenapine | Total |
|         | \(N = 80\) | \(\text{Z-score for age and sex, mean (SD)}\) | \(N = 241\) | \(\text{Z-score for age and sex, mean (SD)}\) | \(N = 321\) | \(\text{Z-score for age and sex, mean (SD)}\) |
| Weight, kg | | | | | |
| Baseline | 1.11 (1.16) | 0.83 (1.19) | 0.90 (1.18) |
| Endpoint | 1.21 (1.14) | 0.91 (1.18) | 0.98 (1.17) |
| Change\(^b\) | 0.10 (0.32) | 0.08 (0.36) | 0.08 (0.35) |
| BMI, kg/m\(^2\) | | | | | |
| Baseline | 1.15 (0.98) | 0.88 (1.15) | 0.94 (1.12) |
| Endpoint | 1.25 (0.98) | 0.93 (1.19) | 1.00 (1.15) |
| Change\(^b\) | 0.10 (0.31) | 0.05 (0.49) | 0.06 (0.45) |

\(\text{BMI}\) body mass index, \(\text{SD}\) standard deviation

\(a\) Z-score indicates how many SDs a value is away from the expected value for a person of the same sex and age. A Z-score of 0 corresponds to the 50th percentile (i.e., the median), a negative Z-score corresponds to a percentile below the median, and a positive Z-score corresponds to a percentile above the median. Percentiles are corrected for age and sex

\(b\) Change from baseline to endpoint was calculated using the open-label baseline

\(\Delta\) Adis
for up to 50 weeks. Somnolence (26.8 %), weight gain (18.1 %), sedation (15.9 %), and headache (12.8 %) were the most commonly reported individual TEAEs and somnolence, sedation, and fatigue were the most common TEAEs leading to discontinuation in the OLE (1.9, 1.2, and 1.2 %, respectively). These TEAEs, combined with dizziness and oral hypoesthesia, occurred more frequently in asenapine-naive patients and may, therefore, be TEAEs to consider monitoring with initiation of asenapine in this population. Additionally, SAEs that led to discontinuation included ADHD, aggression, agitation, anxiety, dystonia, intentional overdose, suicidal behavior, and suicidal ideation. These SAEs are as expected for a long-term, open-label study in this patient population, making it difficult to interpret if, or how, they are related to the study medication. That said, during the acute trial the incidence of these SAEs was higher in those randomized to receive asenapine and, in addition, healthy lifestyle instruction should accompany treatment.

An average improvement in the YMRS total score and CGI-BP severity score was seen over time in this uncontrolled trial. In addition, patients who transitioned to asenapine in the OLE after receiving placebo during the acute phase essentially achieved the same level of response as the asenapine/asenapine group by the end of the extension study. Of note, almost one third of patients who met the criterion for response (50 % improvement in YMRS) at the end of the acute trial failed to sustain this level of improvement at all visits during the OLE. This may be a reflection of the strict maintenance criterion since patients were assessed as having failed to maintain response even if the drop below the 50 % improvement threshold was temporary and response was regained at subsequent visits. Additionally, since fluctuation of symptoms in bipolar disorder may be more common in pediatric patients than in adults [5], patients who failed to maintain response at every assessment may have experienced symptom fluctuation that is characteristic of the pediatric course of the disorder. However, after 26 weeks of open-label asenapine treatment, 79 % of patients were YMRS 50 % responders and 68 % of patients achieved YMRS remission (YMRS total score ≤12), suggesting that a large proportion of patients achieved clinically meaningful symptom improvement.

### 4.1 Limitations

Although this study had a relatively large sample size, it was an open-label study and efficacy and safety results should be interpreted with caution. In addition, over half of the patients in this study discontinued prematurely which may limit generalizability of these results; however, the completion rate (44.6 %) in this study was similar to that seen in other long-term studies of antipsychotics in adolescent populations [23–25] and may be related to the length of the trial and the ages of the patients. Furthermore,
results may not be generalizable to the entire child and adolescent population with BP-1 due to the inclusion and exclusion criteria used in this study. While not a limitation of the study per se, accurate diagnosis of BP-1 in pediatric patients may be challenging; symptoms of bipolar illness can be seen in other populations without bipolarity [26]. In the DSM, fifth edition, increased energy has been added into the ‘A criterion’ for the diagnosis of the manic features specifier which is more inclusive than the DSM-IV-TR diagnosis of manic episode and requires at least three opposite pole symptoms [27]. It is therefore unclear how this may affect results of clinical trials that previously enrolled patients on the basis of DSM-IV-TR criteria.

5 Conclusions

Long-term therapy with asenapine was generally well tolerated in pediatric patients aged 10–17 years with BP-1, although 15.0% of patients discontinued due to TEAEs
and 34.8% experienced clinically significant (≥7%) weight gain with long-term asenapine therapy. Further studies are warranted in children and adolescents with BP-I to improve our understanding of weight gain and other metabolic abnormalities associated with long-term ase-napine treatment, and their relation to the normal changes that occur in this patient population during growth. The combination of somnolence, sedation, and hypersomnia was the most commonly reported TEAE of interest. Somnolence and sedation were also among the most common TEAEs leading to discontinuation and may be treatment-limiting in a minority of patients.

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Compliance with Ethical Standards

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Research involving human participants All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Informed consent Informed consent was obtained from all individual participants included in the study.

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