Efficacy and safety of lurasidone in acutely psychotic patients with schizophrenia: A 6-week, randomized, double-blind, placebo-controlled study

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Aim: The aim of this study was to evaluate the efficacy of lurasidone in acute schizophrenia in Japan and other countries.

Methods: Subjects (aged 18–74 years) diagnosed with schizophrenia were randomized to lurasidone 40 mg/day or placebo. The primary efficacy endpoint was change from baseline on the Positive and Negative Syndrome Scale (PANSS) total score at Week 6. Secondary efficacy assessments included the Clinical Global Impression-Severity Scale (CGI-S). Safety endpoints included adverse events, and laboratory and electrocardiogram parameters.

Results: A total of 483 subjects were randomized to lurasidone or placebo; 107 subjects were from Japan. Mean changes from baseline at Week 6 endpoint in PANSS total scores were −19.3 in the lurasidone group and −12.7 in the placebo group (treatment difference: P < 0.001, effect size = 0.41). Changes from baseline for Week 6 CGI-S scores were −1.0 for lurasidone and −0.7 for placebo (treatment difference: P < 0.001, effect size = 0.41). All-cause discontinuation during the 6-week, double-blind period was 19.4% for lurasidone and 25.4% for placebo, and discontinuation rates due to adverse event were 5.7% for lurasidone and 6.4% for placebo. The following common treatment-emergent adverse events occurred in more than 2% on lurasidone and at a rate at least twice that of the placebo group: akathisia (4.0%), dizziness (2.8%), somnolence (2.8%), abdominal discomfort (2.0%) and asthenia (2.0%). No significant changes in bodyweight or metabolic parameters were observed.

Conclusion: Lurasidone 40 mg once daily dosing demonstrated efficacy in a patient population with acute schizophrenia, including subjects from Japan, and was generally safe and well-tolerated.

Keywords: antipsychotic agents, clinical trial, efficacy, lurasidone, schizophrenia.

Globally, schizophrenia is estimated to affect at least 20 million people.1 Impairments in functioning are extensive with schizophrenia, ranking it among the 20 leading causes of disability.2,3 Reduced lifespan is also apparent among people with schizophrenia, primarily related to increased rates of suicide and cardiovascular disease.4,6

Lurasidone is a benzisothiazole, second-generation antipsychotic drug marketed in the USA, Canada, the European Union, Switzerland, Australia, and Brazil for the treatment of schizophrenia in recommended doses between 40 and 160 mg/day. This novel compound possesses potent antagonist affinity for dopamine D2 and serotonin 5-HT2A and 5-HT7 receptors, moderate antagonist affinity at α2A and α2C adrenergic receptors, and partial agonist affinity at 5-HT1A receptors.7 Unlike some other second-generation antipsychotics, lurasidone has either no or minimal affinity for the 5-HT2C receptors, histamine H1 receptors, and muscarinic M1 receptors that are related to weight gain, metabolic syndrome, or sedation.7

The efficacy, safety, and tolerability of lurasidone in the treatment of schizophrenia has been demonstrated in a number of studies conducted in the USA, Europe, Asia, and South America. A meta-analysis of eight placebo-controlled, short-term studies involving 2373 patients with schizophrenia found that lurasidone at doses up to 160 mg/day was superior to placebo with regard to change in total psychopathology, positive symptoms, negative symptoms, and general psychopathology.8 Lurasidone has not been found to be significantly

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associated with metabolic dysfunction as measured by changes in triglycerides, total cholesterol, high-density lipoprotein cholesterol, low-density lipoprotein cholesterol, fasting glucose, HbA1c, fast insulin, or prolactin level compared to placebo.\(^7\)

None of the eight placebo-controlled trials involved individuals with schizophrenia from Japan.\(^8\) Two recent clinical trials have been conducted that included patients with schizophrenia from Japan. In the first trial, no significant endpoint differences on the primary outcome were found for either dose of lurasidone (40 mg/day or 80 mg/day) or risperidone versus placebo.\(^9\) However, the selectivity criteria of this trial were different from previous trials conducted in USA that had reported positive results for lurasidone.\(^10-12\) As with the US trials, the current study increased the severity criteria required for entry from a Positive and Negative Syndrome Scale (PANSS)\(^13\) total score ≥70 to a PANSS total score ≥80; and it required patients to be experiencing an acute exacerbation of mainly positive symptoms (e.g., exacerbation of delusion or hallucination; thought disorder). In the second trial, neither lurasidone 40 mg/day nor 80 mg/day significantly separated from placebo on the primary outcome in a modified intent-to-treat analysis. A full intent-to-treat (ITT) analysis conducted as a secondary analysis did find that lurasidone 40 mg and 80 mg were superior to placebo.\(^14\) Although the inclusion/exclusion criteria of this study were generally similar to those of the previous US trials, the study also included partially treatment-resistant patients and those with primarily negative symptoms; and these sample characteristics may have reduced the degree of study sensitivity.

The inconsistent results of the Japan studies raised the question of the generalizability of the lurasidone acute schizophrenia efficacy findings across various populations, including Japanese. Adding to this question were previous studies that have found ethnic/nationality differences in response to some antipsychotics.\(^15-17\) Further research on the generalizability of the effects of lurasidone in the treatment of schizophrenia was therefore warranted. The primary objective of the current study was to conduct a Phase 3 evaluation of the efficacy and safety of lurasidone in a geographically diverse sample of subjects, including some from Japan, with acute exacerbation of schizophrenia and the presence of positive symptoms.

**Methods**

**Study design and participants**

This was a multicenter, randomized, double-blind, placebo-controlled, parallel-group study designed to evaluate the efficacy and safety of lurasidone 40 mg/day administered over a 6-week period in patients with acute schizophrenia (clinical trial registration, EudraCT number: 2016–000060–42; study initiated May 2016 and completed November 2018). The study consisted of a screening/washout phase (up to 21 days, including a 3- to 7-day single-blind washout phase) followed by a double-blind treatment phase (6 weeks) and a follow-up visit (7 days following last dose of study drug). Hospitalization was required during the single-blind washout and from baseline to Week 2 (day 15 ± 2).

This study was conducted in accordance with the International Conference on Harmonization Guideline for Good Clinical Practice and the Declaration of Helsinki at 73 clinical sites in five countries (Japan, Ukraine, Russia, Romania, and Poland). The protocol was approved by the Ethics Committee at each participating center. Written informed consent was obtained from each patient following a detailed explanation of study procedures. This RCT manuscript was written according to the Consolidated Standards of Reporting Trials 2010 guideline.

Patients (aged 18–74 years) were diagnosed with schizophrenia according to a clinical interview using the Mini-International Neuropsychiatric Interview (MINI)\(^18\) 6.0.0 with Diagnostic and Statistical Manual of Mental Disorder, 4th Edition, Text Revision (DSM-IV-TR)\(^19\) criteria. To be included in the study, patients also had to meet the following key criteria: a PANSS\(^13\) total score ≥80; a PANSS item score ≥4 (moderate) on two or more of the following PANSS items: delusions, conceptual disorganization, hallucinations, suspiciousness, or unusual thought content at both screening and baseline; a score of 4 (moderately ill) or higher on the Clinical Global Impressions-Severity of Illness (CGI-S)\(^20\) at screening and baseline; an acute exacerbation of mainly positive symptoms (e.g., exacerbation of delusion or hallucination; thought disorder) for no longer than 2 months prior to the screening visit and marked deterioration of function from baseline (by history); and able to be hospitalized from Visit 2 (washout) through Visit 5 (Week 2) assessments. Key exclusion criteria were: continuous hospitalization for ≥3 months (90 days) immediately prior to screening; continuous hospitalization for ≥14 days for acute exacerbation of psychotic symptoms immediately prior to screening in patients who had been treated continuously with adequate doses of one or more antipsychotic agents for ≥4 weeks immediately prior to screening; a decrease of ≥20% in the PANSS total score between screening and baseline visits; PANSS total score below 80 at baseline; patient is considered to be at imminent risk of suicide or injury to self or others; history of treatment with clozapine for refractory psychosis or treatment-resistant schizophrenia; receiving a total dose of antipsychotic medication equivalent to ≥12.0 mg/day of haloperidol at the screening visit; received any depot antipsychotic drugs (sustained-release formulation) more recently than the minimum required washout prior to screening; and received fluoxetine within 1 month of screening.

**Randomization and masking**

Randomization was implemented (1:1 ratio of drug to placebo) using an Interactive Voice/Web Response System (IXRS) performed at baseline. A unique subject number was assigned by the IXRS when a subject entered the screening period. The unique subject number allocated a patient to a particular treatment group and identified the subject for data collection purposes. Patients, investigators, and all research staff remained blinded to the identity of the treatment from the time of randomization until database lock and unblinding. Study drugs were all identical in packaging, labeling, schedule of administration, and appearance.

**Drug administration and concomitant medications**

Drug study consisted of tablets containing either lurasidone 40 mg/day or placebo and was administered orally, once daily in the evening, with food or within 30 min after eating.

When a patient was an inpatient, the hospital pharmacist ensured daily compliance with the dosing regimen. When/if a patient was an outpatient, compliance was monitored closely at each visit. Patients were instructed to bring all unused study drugs with them to each visit. Compliance was assessed by counting tablets and dividing the actual number of doses taken (per tablet count) by the number of doses the subject should have taken within a visit period and multiplying by 100. All patients were reminded of the importance of strict compliance with taking the study drug as directed, with food, for the effectiveness of treatment and for the successful outcome of the study.

Patients were required to discontinue prohibited medications, including antipsychotics, antidepressants, mood stabilizers, and other psychotropics. Potent inducers or inhibitors of the CYP3A4 enzyme system were prohibited during all phases of this study. Biperiden, trihexyphenidyl, diphenhydramine, or promethazine were also permitted if benzotropine was not available for the management of treatment-emergent movement disorders, or if a subject had an inadequate response or intolerability to benzotropine treatment. Treatment with propranolol (≤120 mg/day) was permitted as needed for akathisia. Concomitant use of lorazepam, zolpidem, temazepam, brotizolam, triazolam, lormetazepam, zopiclone, or eszopiclone was permitted within protocol-specified dose limits and timing constraints (not administered within 8 h prior to PANSS rating or other assessments).
**Efficacy assessments**

The PANSS and CGI-S were administered weekly throughout the study. The primary endpoint was change from baseline in PANSS total score at Week 6. Secondary endpoints included change from baseline in: PANSS total score at Weeks 1, 2, 3, 4, and 5, CGI-S score at each post-baseline visit, PANSS subscale scores at each post-baseline visit, and PANSS 5-factor Lindenmayer model scores, including negative symptoms, excitement, cognitive disorders, positive symptoms, and anxiety/depression, at each post-baseline visit. Additional secondary outcomes included the proportion of subjects who achieved a clinical response, defined as 20% or greater improvement from baseline in PANSS total score at Week 6 using last observation carried forward (LOCF); change from baseline in Calgary Depression Scale for Schizophrenia (CDSS) total scores at Weeks 3 and 6; and time to all-cause discontinuation from baseline. The EuroQOL-5 Dimensions-3 Levels (EQ-5D-3L) scale, evaluated by subjects using a paper scale at baseline and Week 6, was included as an exploratory endpoint.

**Safety assessments**

Safety endpoints included assessment of treatment-emergent adverse events (AE), laboratory tests, vital signs, waist circumference, bodyweight, body mass index, QTc interval determined from electrocardiography measurements, and use of concomitant antiparkinsonian drugs. The clinician-rated Drug-Induced Extrapyramidal Symptom Scale (DIEPSS) was used to assess extrapyramidal symptoms induced by antipsychotics. The emergence of suicidality was examined using the Columbia-Suicide Severity Rating Scale (C-SSRS). AE, vital signs, bodyweight, DIEPSS, and C-SSRS were measured weekly throughout the 6-week study and additionally at a Week-7 follow-up visit. Electrocardiography and laboratory tests were conducted at baseline and Week 6 (or end of study).

**Statistical analyses**

The ITT population was defined as all randomized patients who received at least one dose of study drug and had both baseline and at least one post-baseline assessment on the PANSS total score. The primary efficacy endpoint of change from baseline in the PANSS total score at Week 6 was analyzed using a mixed model for repeated measures (MMRM), with fixed factors of pooled study center, visit, treatment, and treatment-by-visit interaction, and baseline PANSS total score as a covariate. An unstructured covariance matrix was used for the within-subject correlation and the Kenward-Roger’s approximation was used to calculate the denominator degree of freedom. The continuous secondary efficacy variables were analyzed in a similar way as done for the primary efficacy variable. Effect sizes for continuous measures on the primary and secondary variables were calculated as the absolute least squares mean difference divided by the model estimate of standard deviation. The counts, percentages, and odds ratio (OR) of PANSS responders at Week 6 were analyzed using a logistic regression model with terms for baseline PANSS total score, pooled study center, and treatment. Numbers needed to treat (NNT) were calculated. Time to all-cause discontinuation was examined using a Kaplan-Meier product-limit survival curve. Analyses of
efficacy within the subgroup of Japanese patients focused on reporting of effect sizes comparing drug and placebo in change from baseline to Week 6 because the study was not powered for subgroup analyses. Safety was evaluated in the safety population, defined as all patients randomized who received at least one dose of study drug. All safety variables were summarized using descriptive statistics. For patients with missing data, PANSS responders are analyzed using LOCF method. All statistical inference, unless otherwise stated, was performed with two-sided tests at the significance level of 0.05. All data analyses were conducted using SAS Version 9.4.

**Determination of sample size**

In a previous clinical trial designed to evaluate the efficacy and safety of lurasidone (40 mg/day and 80 mg/day) treatment for 6 weeks compared to placebo in Asian (Japanese, Korean, Taiwanese, and Malaysian) subjects with acute schizophrenia, the effect size in change from baseline of the PANSS total score for the lurasidone group over placebo at Week 6 in the ITT population was 0.30 in the full sample and 0.37 in a subgroup of patients who met the inclusion/exclusion criteria of the present study. The effect size in the present study was estimated at 0.32, resulting in a sample size of 207 per treatment group to yield a 90% power with a two-sided 5% significance level. Allowing for attrition, an inflation factor of 1.14 was calculated for an MMRM method as primary efficacy analysis using the method by Lu et al., producing the target number of randomized subjects of 472 (236 per treatment group). Retention rates and correlation matrix used for the power calculation were estimated from the previous trial.

**Results**

**Patient disposition and baseline characteristics**

A total of 593 patients were screened for the study to achieve 483 randomized (Fig. 1). There were 247 randomized to the lurasidone group, all of whom received at least one dose of medication, and 236 randomized to placebo, one of whom did not receive the study drug and was excluded from the ITT and safety population. Two subjects in the lurasidone group and three in the placebo group had no
Table 1. Baseline characteristics (ITT population)

|                                      | Placebo N = 233 | Lurasidone N = 245 | Overall N = 478 |
|--------------------------------------|-----------------|--------------------|-----------------|
| Sex, (Male), n (%)                   | 120 (51.5)      | 119 (48.6)         | 239 (50.0)      |
| Age (years), mean ± SD               | 39.3 ± 11.4     | 41.0 ± 11.0        | 40.2 ± 11.2     |
| Race, n (%)                          |                 |                    |                 |
| White                                | 183 (78.5)      | 183 (74.7)         | 366 (76.6)      |
| Asian                                | 50 (21.5)       | 59 (24.1)          | 109 (22.8)      |
| Other                                | 0               | 3 (1.2)            | 3 (0.6)         |
| Country (Japan), n (%)               | 49 (21.0)       | 58 (23.7)          | 107 (22.4)      |
| Duration of illness (years), mean ± SD| 10.4 ± 8.3     | 10.6 ± 8.0         | 10.5 ± 8.2      |
| Duration of current acute exacerbation of symptoms (days), mean ± SD | 20.0 ± 12.9    | 20.1 ± 13.1        | 20.1 ± 13.0     |
| Number of prior hospitalizations (4 or more), n (%) | 135 (57.9) | 141 (57.6) | 276 (57.7) |
| Diagnosis of paranoid type, n (%)    | 212 (91.0)      | 229 (93.5)         | 441 (92.3)      |
| Body mass index (kg/m²), mean ± SD  | 25.2 ± 4.9      | 25.3 ± 4.5         | 25.2 ± 4.7      |
| PANSS total score, mean ± SD        | 101.7 ± 11.5    | 102.8 ± 11.0       | 102.3 ± 11.3    |
| PANSS positive subscale, mean ± SD  | 25.6 ± 3.9      | 25.9 ± 4.0         | 25.7 ± 3.9      |
| PANSS negative subscale, mean ± SD  | 24.3 ± 4.4      | 24.6 ± 4.0         | 24.5 ± 4.2      |
| PANSS general subscale, mean ± SD   | 52.3 ± 6.4      | 51.8 ± 6.6         | 52.0 ± 6.5§     |
| CGI-S score, mean ± SD              | 4.9 ± 0.6       | 5.0 ± 0.6          | 5.0 ± 0.6       |
| Prior antipsychotic use, n (%)      |                 |                    |                 |
| Typical                             | 61 (26.2)       | 68 (27.8)          | 129 (27.0)      |
| Atypical                            | 106 (45.5)      | 116 (47.3)         | 222 (46.4)      |
| Typical + atypical                  | 56 (24.0)       | 48 (19.6)          | 104 (21.8)      |
| Not used                            | 10 (4.3)        | 13 (5.3)           | 23 (4.8)        |

Note: the decision to analyze the prior antipsychotic use was made after unblinding of the data.

1Duration of illness indicates years from the initial episode of schizophrenia to the informed consent.

2Duration of current acute exacerbation of symptoms is days from the onset of current episode to the informed consent.

3The mean ± SD was not pre-specified by statistical analysis plan.

CGI-S, Clinical Global Impressions-Severity of Illness; PANSS, Positive and Negative Syndrome Scale; SD, standard deviation.

post-baseline PANSS assessments, leaving final ITT sample sizes of 245 and 233, respectively.

There were no subjects who were considered noncompliant. The most common reasons for discontinuing the intervention were withdrawal of consent, lack of efficacy, and adverse events. Demographic and clinical characteristics of the randomized sample were similar between the lurasidone and placebo groups (Table 1; Supplementary Materials Table S1). Sample sizes by country were Japan 107, Ukraine 193, Russia 146, Romania 23, and Poland 9.

Concomitant medications in safety population were used by 42.1% (n = 104) of those in the lurasidone group and 45.1% (n = 106) of those in the placebo group. These were primarily anxiolytics (n = 66, 26.7%, for lurasidone; n = 74, 31.5% for placebo), most commonly lorazepam, and hypnotics/sedatives (n = 50, 20.2% for lurasidone; n = 53, 22.6% for placebo), most commonly brotizolam. Antiparkinsonian medications were used by 4.9% (n = 12) of those in the lurasidone group and 1.3% (n = 3) of those in the placebo group.

Efficacy

Using the MMRM model, the mean change from baseline to Week 6 in PANSS total score (primary endpoint) was significantly greater for the lurasidone group compared to the placebo group with a difference of −6.6 (standard error [SE] = 1.58, 95% confidence interval [CI], −9.7, −3.5; P < 0.001; effect size = 0.41; LS mean of −19.3, −12.7, respectively). Treatment differences in change from mean baseline PANSS total score were observed at Week 2 and at subsequent visits (all had nominal P-values < 0.001, not adjusted for multiple comparisons; Fig. 2a). Compared with placebo, treatment with lurasidone resulted in significantly greater proportions of PANSS responders at the LOCF endpoint (odds ratio [OR] = 2.36, 95% CI, 1.57, 3.55; NNT = 6). The response for lurasidone-treated patients was 60.0% compared to 42.5% for placebo patients (Supplementary Materials Table S2).

Similar results were evident on the primary endpoint for the subgroup of patients in the Japan population. The reduction in mean PANSS total score from baseline to Week 6 was −13.7 for the lurasidone group and −6.8 for the placebo group for the treatment difference (SE) of −6.8 (3.52) (P = 0.054, effect size = 0.42). A treatment-by-country interaction included in the full ITT analysis was not significant (P = 0.836).

Treatment with lurasidone resulted in significantly greater reductions from mean baseline CGI-S scores beginning at Week 2 (Fig. 2b). The effect size for the ITT population at Week 6 was 0.41 (P < 0.001 from MMRM analysis). Within the subgroup of patients from Japan, the reduction in mean CGI-S scores from baseline to Week 6 was −0.7 for the lurasidone group and −0.6 for the placebo group (effect size = 0.08).

Compared to placebo-treated patients, lurasidone-treated patients showed significantly greater reductions from baseline to Week 6 on the PANSS Positive Symptom scale, PANSS Negative Symptom scale, PANSS General Psychopathology scale, and all of the PANSS Lindenmayer model scores (negative symptoms, excitement, cognitive disorders, positive symptoms, and anxiety/depression; Table 2). Effect
The incidence of any AE occurred in generally similar proportions between treatment groups (Table 3). Two patients in the lurasidone group had serious AE (schizophrenia; n = 2). Four patients in the placebo group had serious AE (schizophrenia; n = 2; hand fracture, n = 1; suicide attempt, n = 1). Fourteen patients in the lurasidone group and 15 in the placebo group had AE that led to study drug discontinuation. There were no deaths during the study.

Common AE that occurred in more than 2% on lurasidone and at a rate at least twice that of the placebo group were akathisia (4.0%), dizziness (2.8%), somnolence (2.8%), abdominal discomfort (2.0%), and asthenia (2.0%; Table 3 and Supplementary Tables S4 and S5). There were 11 (4.5%) lurasidone-treated patients who had a hyperglycemia/new-onset diabetes AE compared to seven (3.0%) placebo-treated patients. No patients had any dyslipidemia AE. Weight gain AE were reported by four (1.6%) patients in the lurasidone group and one (0.4%) patient in the placebo group. There were 11 (4.4%) patients in the lurasidone group and eight (3.4%) in the placebo group who had any hypersensitivity AE.

There were no clinically meaningful between-group differences in the mean changes from baseline to Week 6 in blood chemistry values or serum prolactin (Table 4). No statistically significant between-group differences were evident for change from baseline to Week 6 in weight gain, RR interval, PR interval, or QT interval (Table 4). No clinically meaningful changes in vital signs were observed from baseline to Week 6 in either treatment group.

### Table 2. Mean (SE) change from baseline to Week 6 on outcomes

| Outcome                        | Placebo N = 233 | Lurasidone N = 245 | Effect size | P-value | Effect size |
|--------------------------------|-----------------|---------------------|-------------|---------|-------------|
| PANSS total score              | −12.7 (1.15)    | −19.3 (1.10)        | 0.41        | <0.001  | 0.42        |
| CGI-S score                    | −0.66 (0.07)    | −1.03 (0.06)        | 0.41        | <0.001  | 0.08        |
| PANSS subscales                |                 |                     |             |         |             |
| Positive subscale              | −3.9 (0.37)     | −6.1 (0.35)         | 0.44        | <0.001  | 0.35        |
| Negative subscale              | −2.5 (0.27)     | −3.3 (0.26)         | 0.22        | 0.030   | 0.40        |
| General Psychopathology subscale | −6.8 (0.61) | −10.0 (0.58)        | 0.39        | <0.001  | 0.33        |
| PANSS Lindenmayer 5-factor scales |             |                     |             |         |             |
| Negative symptoms              | −2.6 (0.27)     | −3.4 (0.26)         | 0.23        | 0.019   | 0.33        |
| Excitement                     | −1.5 (0.22)     | −2.5 (0.21)         | 0.32        | 0.002   | 0.21        |
| Cognitive disorders            | −1.6 (0.18)     | −2.4 (0.17)         | 0.31        | 0.002   | 0.48        |
| Positive symptoms              | −2.4 (0.23)     | −3.7 (0.22)         | 0.40        | <0.001  | 0.21        |
| Anxiety/depression             | −2.9 (0.23)     | −4.0 (0.22)         | 0.34        | <0.001  | 0.29        |
| CDSS score                     | −1.17 (0.18)    | −1.43 (0.17)        | 0.11        | 0.256   | 0.12        |
| EQ-5D-3L index value           | 0.03 (0.01)     | 0.05 (0.01)         | 0.11        | 0.228   | 0.02        |

†The effect size was not pre-specified by statistical analysis plan.

CDSS, Calgary Depression Scale for Schizophrenia; CGI-S, Clinical Global Impressions-Severity of Illness; EQ-5D-3L, EuroQOL-5 Dimensions-3 Levels; PANSS, Positive and Negative Syndrome Scale; SE, standard error.

### Table 3. Summary of adverse events

| Category                          | Placebo N = 235 | Lurasidone N = 247 |
|-----------------------------------|-----------------|---------------------|
| Overall                           |                 |                     |
| Any AE                            | 120 (51.1)      | 116 (47.0)          |
| Serious AE                        | 4 (1.7)         | 2 (0.8)             |
| AE leading to discontinuation     | 15 (6.4)        | 14 (5.7)            |
| AE of special interest            |                 |                     |
| Any extrapyramidal AE             | 12 (5.1)        | 20 (8.1)            |
| Hyperglycemia/new-onset diabetes  | 7 (3.0)         | 11 (4.5)            |
| Weight gain                       | 1 (0.4)         | 4 (1.6)             |
| ≥7% increase in weight            | 2 (0.9)         | 8 (3.3)             |
| Hypersensitivity                  | 8 (3.4)         | 11 (4.5)            |
| Common adverse events†            |                 |                     |
| Akathisia                        | 4 (1.7)         | 10 (4.0)            |
| Dizziness                         | 3 (1.3)         | 7 (2.8)             |
| Somnolence                        | 0               | 7 (2.8)             |
| Abdominal discomfort              | 0               | 5 (2.0)             |
| Asthenia                          | 2 (0.9)         | 5 (2.0)             |

†Treatment-emergent adverse events reported in ≥2% on lurasidone and ≥ 2x placebo.

AE were classified using MedDRA Version 19.1. Number (%) of patients is shown. AE, treatment-emergent adverse event.

At Week 6, adjusted LS means (SE) for DIEPSS total scores (excluded overall severity item) were similar between the lurasidone group (−0.16 [0.058]) and the placebo group (−0.14 [0.061]) (P = 0.817). The proportions of subjects who had at least one
occurrence of suicidal ideation as measured by the C-SSRS were similar between the lurasidone group (17 patients, 6.9%) and placebo group (16 patients, 6.8%). One patient in the lurasidone group and two patients in the placebo group attempted suicide.

Discussion

In this 6-week placebo-controlled study, the efficacy and safety of lurasidone was demonstrated in acutely psychotic patients with schizophrenia recruited in Japan and other countries. Statistically significant differences between lurasidone and placebo on the PANSS total score were evident as early as Week 2 and were apparent at all subsequent visits, with a moderate effect size (0.41) at Week 6. The difference in improvements at endpoint was clinically meaningful as evidenced by a sizeable difference in response rates (60.0% for lurasidone; 42.5% for placebo) and single-digit number needed to treat for response rate (NNT = 6). The effects of lurasidone were broad-based: the CGI-S and all subscale scores for the PANSS (Positive, Negative, and General Psychopathology) and PANSS 5-factor Lindenmayer model (negative symptoms, excitement, cognitive disorders, positive symptoms, and anxiety/depression) scores showed significant superiority for lurasidone compared to placebo on change from baseline to Week 6.

The drug–placebo difference on the PANSS total score for the current study was similar to that in a previous trial for lurasidone in the treatment of schizophrenia. In that trial that recruited patients from the USA, Colombia, Lithuania, India, and the Philippines, Meltzer et al.\textsuperscript{11} found a slightly larger difference between lurasidone 40 mg/day and placebo in change from baseline to Week 6 on the PANSS total score (9.7 points difference). The baseline PANSS total score for the current study (mean = 102.3 ± 11.25), however, was approximately half a standard deviation higher than that in the Meltzer et al.\textsuperscript{11} trial (baseline mean PANSS total score of 97), indicating a somewhat more severe sample enrolled in the current study compared to the previous trial. In contrast, the Higuchi et al.\textsuperscript{9} trial that did not demonstrate a significant difference for lurasidone 40 mg, 80 mg or risperidone versus placebo had a much lower mean baseline PANSS total score (91.9 ± 11.8) that may have hindered finding a drug-placebo difference. The low mean baseline severity of that trial was a function of a PANSS total score inclusion criterion set at ≥70, which was lower than the minimum PANSS severity that was required in the previous lurasidone trials.\textsuperscript{10–12} In addition, unlike the previous trials, the Higuchi et al.\textsuperscript{9} trial did not have a CGI-S inclusion criterion and did not include patients with worsening psychotic symptoms at screening. Treatment-resistant schizophrenia was also common in the sample of patients in the trial.\textsuperscript{9} Somewhat different inclusion/exclusion criteria were used in the second Higuchi et al.\textsuperscript{14} trial. In that trial, lurasidone 40 mg and 80 mg did not significantly separate from placebo on the primary outcome in a modified intent-to-treat analysis, but a full intent-to-treat analysis conducted as a secondary analysis did find that lurasidone 40 mg and 80 mg were superior to placebo. The PANSS total score indicated that severity of the sample at baseline was high (mean = 102.8 ± 15.5). However, the lack of drug-placebo differences in this study may\textsuperscript{14} have been due to the enrollment of patients who had taken a total dose of antipsychotic medication equivalent to 12 mg or more haloperidol, and the high prevalence of both predominantly negative symptoms and treatment-resistant schizophrenia at screening. The inclusion and exclusion criteria of the current study were different from these trials (see Supplementary Tables S6a,b). The criteria of the current study included PANSS total score ≥ 80, CGI-S score ≥ 4, inclusion of patients with acute schizophrenia, predominantly positive symptoms prior to screening, and exclusion of haloperidol-equivalent dose ≥12 mg (to exclude patients with treatment-resistant schizophrenia). These differences in inclusion criteria may explain the positive findings of the current study compared to the lack of drug-placebo differences in the primary analyses of two previous trials conducted in Asian countries.\textsuperscript{9,14}

Because of the inconsistent results across the lurasidone trials including Japanese for acute schizophrenia, a network meta-analysis was performed that included two previous trials and the current study.\textsuperscript{2} In this analysis, lurasidone 40 mg/day and 80 mg/day led to

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### Table 4. Baseline and mean change to Week 6 (LOCF) in weight, BMI, laboratory parameters, and ECG parameters (safety population)

| Parameter mean (SD) | Placebo N = 235 | Lurasidone N = 247 |
|---------------------|----------------|-------------------|
|                     | Baseline | Change | n | Baseline | Change | n |
| Bodyweight (kg)     | 71.90 (15.84) | −0.18 (1.96) | 233 | 72.21 (14.78) | 0.10 (2.07) | 246 |
| BMI (kg/m²)         | 25.24 (4.85) | −0.06 (0.70) | 233 | 25.31 (4.48) | 0.03 (0.74) | 246 |
| Waist circumference (cm)\textsuperscript{7} | 86.80 (13.00) | −0.26 (2.58) | 226 | 86.35 (12.18) | −0.03 (3.31) | 240 |
| Triglycerides\textsuperscript{4} (mg/dL) | 127.07 (71.13) | −0.62 (61.59) | 204 | 143.94 (136.58) | −14.33 (125.37) | 211 |
| Total cholesterol\textsuperscript{7} (mg/dL) | 183.86 (40.23) | −4.13 (32.99) | 205 | 188.91 (48.27) | 0.68 (56.75) | 211 |
| HDL cholesterol\textsuperscript{7} (mg/dL) | 107.10 (33.78) | −4.97 (26.39) | 202 | 110.76 (41.88) | −1.83 (25.02) | 208 |
| Blood glucose\textsuperscript{3} (mg/dL) | 93.23 (12.28) | −0.08 (14.71) | 205 | 92.92 (13.04) | 0.44 (12.84) | 211 |
| Hemoglobin A1c (%) | 17.57 (22.66) | −2.42 (22.93) | 222 | 20.50 (30.90) | −1.59 (29.60) | 234 |
| Prolactin, men     | 10.85 (10.35) | −0.16 (10.46) | 114 | 13.21 (11.69) | 0.21 (12.40) | 113 |
| Prolactin, women   | 24.71 (29.14) | −4.81 (30.98) | 108 | 27.40 (40.46) | −3.27 (39.39) | 121 |
| ECG: heart rate\textsuperscript{4} (beats/min) | 72.8 (12.8) | 1.1 (12.7) | 222 | 73.8 (12.2) | −1.6 (11.9) | 240 |
| ECG: RR interval\textsuperscript{4} (msec) | 851.2 (153.6) | −10.2 (143.3) | 222 | 837.2 (145.4) | 18.5 (137.6) | 240 |
| ECG: PR interval\textsuperscript{4} (msec) | 151.0 (20.2) | −0.2 (13.7) | 222 | 154.9 (21.9) | 0.5 (14.8) | 240 |
| ECG: QTcF interval\textsuperscript{4} (msec) | 401.3 (19.1) | 0.0 (16.9) | 222 | 403.9 (19.8) | 0.5 (14.9) | 240 |

\( P < 0.05. \) Comparisons of the lurasidone vs placebo at LOCF endpoint are based on a rank ANCOVA analysis.

\( ^{1} \)Significance testing was not pre-specified by statistical analysis plan.

\( ^{2} \)Fasting was required per protocol.

BMI, body mass index; ECG, electrocardiogram; HDL, high-density lipoprotein; LDL, low-density lipoprotein.

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significant benefit in the treatment of acute schizophrenia. The results of these RCTs may have been inconsistent due to different selection criteria, but conducting such a network meta-analysis suggests that lurasidone 40 mg/day is effective in patients with acute schizophrenia, including Japanese.

The pattern of adverse events found in the current study was similar to the AE profile observed in previous placebo-controlled trials that evaluated lurasidone 40 mg/day in the treatment of schizophrenia. The consistent minimal effects on weight and metabolic parameters observed in this and previous studies have been attributed to the lack of clinically relevant affinity of lurasidone for the receptors (H1-histamine and 5-HT2C) that are thought to be associated with weight gain.7,28 The minimal effect of lurasidone on metabolic parameters is especially important given the high risk for metabolic syndrome that has been found for individuals with schizophrenia.29

Within the subgroup of Japanese patients, the treatment effect size (0.42) comparing lurasidone with placebo on change from baseline to Week 6 on the PANSS total score was nearly identical to that found in the overall sample (effect size = 0.41). Effect sizes for the PANSS subscales and Lindenmayer scores in the Japanese subgroup were also similar to those found in the overall sample; and all treatment-by-subgroup (Japan vs others) terms were non-significant for primary and secondary measures. This suggests that lurasidone has similar efficacy for Japanese patients as it does for patients from other countries. This result, however, would need to be confirmed in a randomized, placebo-controlled trial that is fully powered to detect differences within a Japanese sample.

The findings of this study should be understood in the context of some limitations. First, the short-term (6-week) nature of the study leaves open the question of longer-term efficacy and safety. However, long-term safety/tolerability has been verified in previous long-term studies, including two 12-month studies30,31 and one 28-week double-blind maintenance study32 in USA. Second, the study was conducted using a sample of individuals with acute symptoms of schizophrenia but excluding individuals continuously hospitalized for >14 days for acute exacerbation of psychotic symptoms immediately prior to screening and who had been treated continuously with adequate doses of one or more antipsychotic agents for ≥4 weeks immediately prior to screening (as well as other exclusion criteria).

In summary, lurasidone 40 mg once daily dosing demonstrated efficacy in a patient population, including subjects from Japan, with acute schizophrenia and predominantly positive symptoms; there seems to be no significant difference in ethnicity or nationality, similar to the results of previous US trials. It was generally safe and well-tolerated.

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Author contributions
M.I, J.I and T.H: contributed to the conception and design of the study, acquisition and interpretation of results. M.N and Y.M: contributed to statistical data analysis of the study. R.S and J.T: contributed to conception and design of the study. K.O and A.F: contributed to acquisition of data and resource management of the study. T.N: contributed to conception and drafting of this manuscript. All authors contributed to and have approved the final manuscript.

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