A Randomised, Double-blind, Placebo-controlled Study of Escitalopram in Patients with Social Anxiety Disorder in Japan

Satoshi Asakura¹*, Taiji Hayano², Atsushi Hagino², Tsukasa Koyama³

¹ Health Care Center and Department of Psychiatry, Hokkaido University Graduate School of Medicine
² Mochida Pharmaceutical Co., Ltd.
³ Clinical Research Center, Oyachi Hospital

*Address for Correspondence:
Satoshi Asakura
North 16, West 7, Sapporo, Hokkaido 060-0816, Japan
e-mail: asakurap@academic.hokudai.ac.jp
TEL: +81-11-706-5418
FAX: +81-11-706-5081

Number of figures/tables: 5
Word count: approx. 3878
Type of article: Original article
Field: General topics in psychiatry and related fields

Neuropsychopharmacology
Abstract

Objective:
This randomised, double-blind placebo-controlled study compared the efficacy and tolerability of escitalopram (10 and 20mg/day) in Japanese patients with social anxiety disorder (SAD).

Research design and methods:
Patients aged 18-64 years with a primary diagnosis of DSM-IV-TR defined SAD, a Liebowitz Social Anxiety Scale Japanese version (LSAS-J) total score ≥60 and a Clinical Global Impression-Severity (CGI-S) score ≥4 at baseline were randomly assigned (1:1:1) to placebo, escitalopram 10mg or 20mg. The primary endpoint was change from baseline to Week 12 in the LSAS-J total score for both escitalopram 10mg and 20mg versus placebo (ANCOVA, FAS, LOCF), using a hierarchical testing procedure. Pre-specified secondary endpoints included LSAS-J sensitivity analyses.

Clinical trial registration:
This study has the www.japic.or.jp identifier: JapicCTI-121842.

Results:
For the primary efficacy endpoint, the difference from placebo in the LSAS-J was -3.9 (p=0.089) for escitalopram 10mg. Since the superiority of escitalopram 10mg over placebo was not confirmed, an analysis without multiplicity adjustment was made, which showed a difference for escitalopram 20mg versus placebo of -9.8 (p<0.001). In pre-specified sensitivity analyses, the difference versus placebo was -4.9 (p=0.035) (ANCOVA, FAS, OC) and -5.0 (p=0.028) (MMRM, FAS) (escitalopram 10mg) and
-10.1 (p<0.001) (ANCOVA, FAS, OC) and -10.6 (p<0.001) (MMRM, FAS) (escitalopram 20mg). Common adverse events (incidence ≥5% and significantly different from placebo) were somnolence, nausea and ejaculation disorder.

Conclusion:

Escitalopram was efficacious, safe and well tolerated by patients with SAD in Japan. Study limitations are discussed including patient characteristics. [240 words]

**Key words:** escitalopram, Japan, Liebowitz Social Anxiety Scale Japanese version (LSAS-J), randomised placebo-controlled study, social anxiety disorder (SAD)

**Short title:** Escitalopram in social anxiety disorder in Japan
Introduction

Escitalopram (ESC) is a selective serotonin reuptake inhibitor (SSRI). As of the end of December 2014, ESC had been approved in 100 countries. Depending on the specific country or region, ESC is approved for indications that include major depressive disorder, panic disorder, social anxiety disorder (SAD), generalised anxiety disorder, obsessive-compulsive disorder, and premenstrual dysphoric disorder.

SAD is a psychiatric disorder characterised by feelings of fear and severe strain stemming from interpersonal communication, with associated symptoms such as tremor, flushing, palpitations, and sweating\(^1\). Anxiety disorders, including SAD, are risk factors associated with suicidal ideation and attempted suicide. The associated risk is reported to increase with a concurrent incidence of a mood disorder and anxiety disorder\(^2\). Because SAD develops around the onset of adolescence and is more intractable than other anxiety disorders and more likely to become chronic\(^3,4\), it is well recognised that the condition requires treatment. In addition, because patients tend to be socially isolated due to the continuous avoidance of social relationships, SAD affects patients’ engagement in school, educational settings and workplaces, with subsequent negative impacts on their economic situation\(^3\). These effects may represent major losses not only to the patients themselves and their families, but also to society as a whole.

Treatments for SAD are broadly classified into pharmacotherapy and psychotherapy, the latter represented by psychotherapeutic interventions, including cognitive behaviour therapy. First-line pharmacotherapy include SSRIs or serotonin noradrenaline reuptake inhibitors (SNRIs)\(^1,5\).
Placebo-controlled studies conducted in Europe, Canada and South Africa support the
efficacy of ESC in the treatment of SAD. The 12-month prevalence of SAD is 2.3% in Japan, where only paroxetine and fluvoxamine have been approved for the
treatment of SAD.

The aim of this clinical study was to investigate the efficacy, safety, and tolerability of
two fixed doses (10 and 20mg/day) of ESC versus those of placebo after 12 weeks of
treatment in Japanese adult patients with SAD.
Patients and methods

Study design

This multicentre, randomised, double-blind, parallel-group, fixed-dose, placebo-controlled study included 588 randomised patients recruited from 86 medical institutions in Japan from June 2012 to March 2014. All participating medical institutions received approval to conduct the study from their local institutional review board prior to study initiation. All study procedures were conducted in compliance with the Declaration of Helsinki and the Ministerial Ordinance on Good Clinical Practice. In addition, the study investigator obtained written informed consent from all patients prior to their participation in the study.

After a 1-week screening period, eligible patients were randomly assigned (1:1:1) to placebo, ESC 10mg/day or ESC 20mg/day for 12 weeks of double-blind treatment. For the ESC 20mg group, patients were treated at an initial dose of 10mg/day for the first week, and then there was a mandatory increase to 20mg/day. Patients were seen at baseline and at Weeks 1, 2, 4, 6, 8 and 12. Patients who were withdrawn were seen as soon as possible after withdrawal. A safety follow-up contact was scheduled for 2 weeks after completion of the treatment period or after withdrawal from the study.

Study medication was given as placebo or ESC tablets of identical appearance. Patients were instructed to take two tablets per day, orally, after supper in the evening.

Main entry criteria

Eligible patients of either sex were aged ≥18 and ≤64 years, with a primary diagnosis of SAD according to the Diagnostic and Statistical Manual of Mental Disorders, 4th
edition, Text Revision (DSM-IV-TR). Patients were diagnosed using the
Mini-International Neuropsychiatric Interview (M.I.N.I.; Japanese version 5.0.0). In
addition, patients were required to have a total score ≥60 on the Japanese version of
the Liebowitz Social Anxiety Scale (LSAS-J) and ≥4 on the Clinical Global
Impression-Severity Scale (CGI-S) and to exhibit fear/anxiety or avoidance traits in at
least 4 items of the LSAS-J, of which ≥2 were social interaction items at screening
and baseline visits. Patients who met any of the following criteria were excluded from
the study: diagnosis of schizophrenia or another psychotic disorder; delirium;
dementia; amnestic disorder or another cognitive disorder; bipolar disorder;
obsessive-compulsive disorder; panic disorder; specific phobias; body dysmorphic
disorder; eating disorder; substance abuse or substance dependence (excluding
nicotine and caffeine); pervasive developmental disorder on Axis I of the
DSM-IV-TR; diagnosis with group A or group B personality disorder and mental
retardation on Axis II of the DSM-IV-TR; history of schizophrenia or another
psychotic disorder or bipolar disorder on Axis I of the DSM-IV-TR; primary diagnosis
with Axis I disorder other than SAD according to the DSM-IV-TR within 24 weeks of
the study; a total score ≥15 on the Montgomery Åsberg Depression Rating Scale
(MADRS); history or complication of convulsive disorder such as epilepsy (excluding
febrile seizure in childhood); patients with congenital long QT interval syndrome,
bleeding tendency, or haemorrhagic diathesis; patients at significant risk of suicide as
clinically judged by the investigator, or patients meeting the criteria of any of C4
through C6 in “C. Suicidality” of the M.I.N.I. or having a score ≥5 on Item 10 of the
MADRS (‘suicidal thoughts’) or with suicidal behaviour according to the Columbia
Suicide Severity Rating Scale Questionnaire (C-SSRS); patients who were pregnant
or breastfeeding, who might be pregnant, or who wanted to get pregnant during the
term of the study; and patients otherwise judged by investigators to be unsuitable for participation in this clinical study.

**Efficacy rating**

The effect of ESC *versus* placebo after 12 weeks of treatment was assessed using the LSAS-J total score. All raters underwent training in the LSAS-J, in order to maximise inter-rater reliability. Only trained raters (all of whom were either psychiatrists or psychologists) were allowed to rate patients.

**Allocation to treatment**

At each site, sequentially enrolled patients were assigned the lowest randomisation number available in blocks of 6. Each patient was assigned a randomisation number according to a randomisation list that was computer generated by the study medication allocation manager. All investigators, trial personnel and patients were blinded to treatment assignment for the duration of the study. The randomisation code was not broken for any patients during treatment.

**Analysis sets**

Safety analyses were based on the all-patients-treated set (APTS), comprising all randomised patients who took at least one dose of study medication. Efficacy analyses were based on the modified intent-to-treat set - the full-analysis set (FAS), comprising all patients in the APTS who had a valid baseline assessment and at least one valid post-baseline assessment of the primary efficacy variable (LSAS-J total score).

Statistical analyses were performed using SAS 9.2 and the level of statistical significance was defined as p<0.05 (two-sided).
Power and sample size calculations

Based on a previous ESC study in SAD\(^8\), with a mean difference to placebo of 7.27 on the change from baseline in the LSAS-J total score at Week 12 and a standard deviation of 24.85, calculations showed that with a power of \(\geq 80\%\), a total of 555 patients should be randomised to detect superiority of ESC to placebo, using a 5\% level of significance and a standard t-test.

Analysis of the primary efficacy endpoint

The prospectively defined primary efficacy analysis was an analysis of covariance (ANCOVA) of the change from baseline in the LSAS-J total score at Week 12 (FAS), with treatment as fixed factors and the baseline LSAS-J total score as a covariate, using last observation carried forward (LOCF). Sensitivity analyses to the primary efficacy analysis were performed using ANCOVA based on data from observed cases (OC), and mixed model repeated measures (MMRM). To control for a two-sided type I error in the primary efficacy endpoint, a closed testing procedure was adopted in which ESC 10mg/day \textit{versus} placebo was tested first and then ESC 20mg/day \textit{versus} placebo. Once an endpoint was non-significant, the formal testing procedure was stopped. For endpoints that occurred after the pre-specified statistical testing procedure was stopped or that were outside the testing procedure, nominal p-values with no adjustment for multiplicity are reported. A post-hoc analysis using ANCOVA (FAS, LOCF) was also made, in which patients who discontinued within 1 week after treatment initiation were excluded and patients with non-severe SAD and severe SAD.
Analysis of secondary efficacy endpoints

The following secondary analyses were prospectively defined: the change from baseline in the LSAS-J total score at other visits, the change from baseline to Week 12 in LSAS-J subscale scores and CGI-S scores, and CGI-I scores, LSAS-J response (≥30% decrease from baseline) and CGI-I response (CGI-I score ≤2) at Week 12.

Response rates were analysed using Fisher’s exact test, and CGI-I scores were analysed by an ANOVA. Changes from baseline were analysed in a manner similar to the primary analysis of the primary endpoint.

Safety assessments

All treatment-emergent adverse events (TEAEs) either observed by the investigator or reported spontaneously by the patient were recorded. Qualified personnel coded TEAEs using the preferred term according to the Medical Dictionary for Regulatory Activities/Japanese (MedDRA/J), Version 16.0. The incidence of individual TEAEs was compared between treatment groups using the Fisher’s exact test. Clinical safety laboratory tests, vital signs, weight, BMI, ECGs, and physical examination findings were also evaluated. The Wilcoxon 2-sample text was used to compare the change in the QTcF between the treatment groups. A safety follow-up contact was scheduled for 2 weeks after completion of the study or after withdrawal from the study. The C-SSRS was used to assess suicide risk in patients at screening, baseline, and weeks 1, 2, 4, 6, 8 and 12.
Results

Patient baseline characteristics

The APTS consisted of 587 patients (n=196 for placebo, n=198 for ESC 10mg, and n=193 for ESC 20mg) after the exclusion of 1 patient who did not take any study medication (Figure 1). Patients had a mean age of about 33 years, and approximately 56% were women. There were no apparent clinically relevant differences at baseline between treatment groups in demographic or baseline clinical characteristics (Table 1). The full-analysis set (FAS) comprised 587 patients.

The mean baseline LSAS-J total score was 94.4 ± 18.1, and the mean age of onset was 19 years (median of 17 years, range 5-61 years) before enrolment. The mean baseline MADRS total score was 3.7 ± 3.9.

Withdrawals from the study

The proportion of patients who discontinued from the study in the treatment period was 10.7% (21/196) in placebo, 10.1% (20/198) in ESC 10mg, and 11.9% (23/194) in ESC 20mg (Figure 1). The most common reason for discontinuation in the ESC group was TEAEs: the proportion of patients who discontinued from the study because of TEAEs was 3.6% (7/196) in placebo, 6.6% (13/198) in ESC 10mg, and 7.2% (14/194) in ESC 20mg.

Efficacy

Primary endpoint

In the primary efficacy analysis, the mean change from baseline in the LSAS-J total score at Week 12 (FAS, LOCF) was -23.1 (placebo), -26.9 (ESC 10mg) and -32.6
The mean difference from placebo for ESC 10mg was -3.9 [95% CI: -8.3, 0.6] (p=0.089) (Table 2). Pre-specified sensitivity analyses OC (FAS) and MMRM (FAS) resulted in p-values of 0.035 and 0.028, respectively (Table 2). Because the superiority of ESC 10mg to placebo (FAS, LOCF) was not statistically significant, a comparison of ESC 20mg to placebo using a closed testing procedure was not possible. However, in an analysis that did not take multiplicity into account (FAS, LOCF), the difference in the mean between placebo and ESC 20mg was -9.8 [95% CI: -14.5, -5.2] (P < 0.001). The estimated change from baseline in the LSAS-J total mean score plotted by visit is shown in Figure 2.

The number of patients who discontinued within 1 week after treatment initiation was 0.5% (1/196) for placebo, 4.5% (9/198) for ESC 10mg, and 2.6% (5/193) for ESC 20mg, although all patients in ESC groups received ESC 10mg/day during the first week. LSAS-J total scores measured at the withdrawal visit increased for these patients, indicating that they had discontinued treatment before a therapeutic effect had been seen. Based on this, a post-hoc efficacy analysis was made excluding these 15 patients. The mean difference to placebo was -5.1 [95% CI: -9.6, -0.7] (p=0.023) for ESC 10mg and -10.6 [95% CI: -15.2, -5.9] (p<0.001) for ESC 20mg (FAS, LOCF).

When stratified according to Cytochrome P450 (CYP) 2C19 phenotype, the mean difference to placebo on the MADRS for poor metabolisers (n=113) was +3.9 [95% CI: -7.0, 14.9] (p=0.477) for ESC 10mg and -11.4 [95% CI: -22.1, -0.7] (p=0.038) for ESC 20mg; and for extensive metabolisers (n=474), the mean difference to placebo was -5.5 [95% CI: -10.4, -0.7] (p=0.026) for ESC 10mg and -9.4 [95% CI: -14.6, -4.1]
(p<0.001) for ESC 20mg (FAS, LOCF).

A small proportion of patients had severe SAD (17.5%), defined as a baseline LSAS-J total score ≥100 and a CGI-S score ≥6. In post-hoc analyses of patients with non-severe SAD (n=484), the difference to placebo in the mean change from baseline in the LSAS-J total score was -5.4 [95% CI: -10.2, -0.7] (p=0.026) for ESC 10mg and -9.9 [95% CI: -14.8, -4.9] (p<0.001) for ESC 20mg. For patients with severe SAD (n=103), the difference to placebo was +2.9 [95% CI: -9.1, 14.9](p=0.635) for ESC 10mg and -9.9 [95% CI: -24.0, 4.2] (p=0.164) for ESC 20mg.

The two subscales of the LSAS-J were also analysed by ANCOVA (FAS, LOCF). The mean difference from placebo for the LSAS-J fear/anxiety subscale total score was -2.1 [95% CI: -4.3, 0.2] (p=0.069) for ESC 10mg and -4.9 [95% CI: -7.3, -2.5] (p<0.001) for ESC 20mg (FAS, LOCF). The mean difference from placebo for the LSAS-J avoidance subscale total score was -1.8 [95% CI: -4.2, 0.5] (p=0.124) for ESC 10mg and -5.0 [95% CI: -7.4, -2.6] (p<0.001) for ESC 20mg (FAS, LOCF) (Table 3).

**Secondary analyses**

Mean CGI-I and CGI-S scores improved throughout the 12-week treatment period in all treatment groups. The CGI-S scores improved from baseline to Week 12 from 4.8 ± 0.8 to 3.8 ± 1.0 (placebo), from 4.8 ± 0.9 to 3.7 ± 1.3 (ESC 10mg) and from 4.9 ± 0.8 to 3.5 ± 1.2 (ESC 20mg). The differences to placebo were -0.1 [95% CI: -0.4, 0.1] (p=0.178) (ESC 10mg) and -0.4 [95% CI: -0.6, -0.2] (p<0.001) (ESC 20mg) (FAS, LOCF).
The CGI-I scores improved from baseline to Week 12 to 2.8 ± 1.1 (placebo), to 2.6 ± 1.1 (ESC 10mg) and to 2.4 ± 1.1 (ESC 20mg). The differences to placebo were -0.2 [95% CI: -0.4, 0.0] (p=0.049) (ESC 10mg) and -0.4 [95% CI: -0.7, -0.2] (p<0.001) (ESC 20mg) (FAS, LOCF). The CGI-I response rates at Week 12 were 37.8% [95% CI: 30.9, 44.9] (placebo), 48.0% [95% CI: 40.8, 55.2] (ESC 10mg), and 54.9% [95% CI: 47.6, 62.1] (ESC 20mg). The differences in the response rate from placebo were 10.2% [95% CI: 0.5, 19.9] (p=0.042) (ESC 10mg) and 17.2% [95% CI: 7.4, 26.9] (ESC 20mg) (p<0.001) (FAS, LOCF).

**Safety and tolerability**

During the 12-week treatment period, approximately 70% of the patients in each ESC treatment group had one or more TEAEs. The most common TEAEs reported by at least 5% of patients for ESC and significantly more frequently than placebo were somnolence, nausea and ejaculation disorder (Table 4). The majority of TEAEs were mild or moderate in severity with the proportions of patients reporting severe TEAEs being 0% (0 of 196) (placebo), 1.5% (3 of 198) (ESC 10mg), and 1.0% (2 of 193) (ESC 20mg). During this period, 34 patients were withdrawn due to TEAEs (Figure 1). TEAEs leading to withdrawal of ≥2 patients in either group were anxiety (n=2) in placebo; headache and nausea (n=3 for each), social phobia, abdominal pain upper, and dizziness (n=2 for each) in ESC 10mg; and nausea (n=2) in ESC 20mg. The proportion of patients with severe TEAEs leading to withdrawal was 1.0% (2/198) (ESC 10mg) and 1.0% (2/193) (ESC 20mg) and the majority resolved after treatment discontinuation.
To investigate the development of possible withdrawal syndrome during the follow-up period, TEAEs in patients who did not concomitantly use an antidepressant during the follow-up period were examined.

Patients who did not receive antidepressants in the follow-up period accounted for 113 of 196 patients in placebo, 114 of 198 patients in ESC 10mg, and 112 of 193 patients in ESC 20mg. The proportion of patients with TEAEs were 10.6% (12/113) (placebo), 19.3% (22/114) (ESC 10mg), and 24.1% (27/112) (ESC 20mg). Dizziness was reported by 2.7% (3/113) (placebo), 6.1% (7/114) (ESC 10mg), and 10.7% (12/112) (ESC 20mg) of patients during follow-up (Table 5). All of these events were mild or moderate.

In the 12-week treatment period 8 serious AEs (SAEs) were reported by 6 patients, 3 patients in ESC 10mg and 3 patients in ESC 20mg. No SAE was reported by more than 1 patient and the types of SAEs were: convulsion, acute pyelonephritis, cervical vertebral fracture, lumbar vertebral fracture, and thoracic vertebral fracture (ESC 10mg) and appendicitis, osteoarthritis, and diabetes mellitus (ESC 20mg). Of the aforementioned events, the cervical vertebral fracture, lumbar vertebral fracture, and thoracic vertebral fracture in ESC 10mg were all events that occurred in the same patient, and the patient recovered from all of these events without discontinuing from study treatment. The other events all occurred in different patients and, except for diabetes, all were alleviated or recovered with study treatment discontinuation and therapy. The patient with diabetes was still receiving pharmacotherapy 4 months after discontinuing from study treatment, at which time the patient’s condition was judged to be following the natural course of diabetes by the investigator, and follow-up was therefore concluded.
No deaths occurred in this study.

The proportion of patients who reported sexual dysfunction-related TEAEs was 2.5% (5/198) in ESC 10mg and 2.6% (5/193) in ESC 20mg while none were reported in placebo. Ejaculation disorder (5 patients) and erectile dysfunction (1 patient) were reported in ESC 10mg, and ejaculation disorder (2 patients), libido decreased (2 patients), and ejaculation delayed (1 patient) and libido increased (1 patient) were reported in ESC 20mg. All sexual dysfunction-related TEAEs were rated by the investigator as mild in severity.

Suicide-related TEAEs were reported by 0.5% (1 of 196 placebo patients) (suicidal ideation), 1.5% (3 of 198 ESC 10mg patients) (2 suicidal ideation and 1 self-injurious behaviour), and 0% (0 of 193 ESC 20mg patients). Each suicide-related TEAE occurred once and there were no statistically significant differences between the placebo and ESC groups. All suicide-related TEAEs were rated by the investigator as mild or moderate in severity. Of the 3 patients in ESC 10mg, 1 of the patients with suicidal ideation and the patient with self-injurious behaviour recovered without therapy, and the other patient with suicidal ideation improved without therapy after discontinuation of the study treatment. These results were supported by the C-SSRS data.

No clinically relevant changes over time or differences between treatment groups were seen in clinical laboratory test results, vital signs, weight, or ECG parameters. Patients gained a mean of 0.24 kg and 0.29 kg (ESC 10mg or ESC 20mg, respectively) and 0.35 kg (placebo) compared to baseline at Week 12 or last assessment. No statistically significant differences were found between ESC 10mg
or ESC 20mg (p=0.751) versus placebo. The differences from placebo in the mean change from baseline in QTcF interval at the end of treatment were 3.0 ms [95% CI: 0.1 to 5.8] (ESC 10mg) and 5.0 ms [95% CI: 2.1 to 7.8] (ESC 20mg).

The proportions of patients with TEAEs were 52.6% (placebo), 85.3% (ESC 10mg), and 65.9% (ESC 20mg) for CYP2C19 poor metabolisers and 57.0% (placebo), 59.8% (ESC 10mg), and 65.1% (ESC 20mg) for extensive metabolisers. The differences from placebo in the mean change from baseline to the end of treatment in the QTcF interval were 4.6 ms (ESC 10mg) (p=0.241) and 3.5 ms (ESC 20mg) (p=0.282) for CYP2C19 poor metabolisers and 2.6 ms (ESC 10mg) (p=0.143) and 5.3 ms (ESC 20mg) (p=0.004) for extensive metabolisers. The completion rate in poor metabolisers was 94.7% (36/38)(placebo), 85.3% (29/34)(ESC 10mg), and 92.7% (38/41)(ESC 20mg). The completion rate in extensive metabolisers was 88.0% (139/158)(placebo), 90.9% (149/164)(ESC 10mg), and 87.5% (133/152)(ESC 20mg).
Discussion

This is the first randomised placebo-controlled clinical study for the treatment of SAD with ESC in Japan. Apart from this study, three clinical studies of ESC have been conducted in patients with SAD in countries outside of Japan. In these studies, the short-term (12 weeks) and long-term (24 weeks) efficacy of ESC treatment and relapse-prevention effect (24 weeks) were established, and its safety and tolerability were also demonstrated\textsuperscript{6,8,9}.

The primary study endpoint in this placebo-controlled study in Japan, the change in the LSAS-J total score at Week 12, was not established for ESC 10mg versus placebo. The primary analysis was performed on data with the missing values imputed by LOCF. The International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH) -E9 guidelines\textsuperscript{10} state that “Unfortunately, no universally applicable methods of handling missing values can be recommended. An investigation should be made concerning the sensitivity of the results of analysis to the method of handling missing values, especially if the number of missing values is substantial.” Thus, we performed prospectively defined sensitivity analyses based on OC and MMRM. The results of these sensitivity analyses demonstrated the superiority of ESC 10mg over placebo. Since different results were obtained from the sensitivity analyses, it was considered likely that the results in the primary efficacy endpoint were affected by the handling of missing values. According to the European Medicines Agency guidelines from 2010\textsuperscript{11}, it is suggested that, in the case of diseases that tend to improve spontaneously over time (such as depression), efficacy is evaluated conservatively with data imputation by LOCF analysis if there are a large number of patients in the active treatment group.
who discontinue treatment at an early stage. Since SAD, like depression, also tends to improve over time, it was suggested that efficacy should be evaluated conservatively by carrying out a LOCF analysis.

The proportion of discontinued patients in each treatment group, which would affect the LOCF analysis, was investigated. Although the dose was 10mg/day for both ESC 10mg and ESC 20mg through Week 1 of the treatment period, the rate of patient discontinuation within this first week of treatment was 0.5% (1 of 196 patients in placebo), 4.5% (9 of 198 patients in ESC 10mg), and 2.6% (5 of 193 patients in ESC 20mg); i.e., a higher rate was observed in the “ESC 10mg”. In the ESC groups, main reason for discontinuation within the first week is TEAE; the number of patients who discontinued due to TEAE was 8 patients in ESC 10mg and 4 patients in ESC 20mg. For the patients who discontinued treatment within Week 1 of treatment, the mean change in the LSAS-J total score at treatment discontinuation indicated a slight worsening in all treatment groups. The period for efficacy evaluation was defined as 12 weeks using the treatment algorithm of Stein et al.\textsuperscript{12} and the British Association for Psychopharmacology's guidelines\textsuperscript{13}, whereas the Canadian Psychiatric Association's guidelines\textsuperscript{1} specify that an early response to pharmacotherapy is usually observed in the first 6-8 weeks, but that it may take 12 weeks or longer for pharmacotherapy to exert its full effects in some cases. Since the primary endpoint included patients who had discontinued treatment before the first week of treatment, the efficacy of ESC 10mg was underestimated by including the values imputed by LOCF.

When the change in the LSAS-J total score at Week 12 of treatment was analysed, significant improvement was demonstrated for ESC 20mg versus placebo using both
the primary analysis method and the sensitivity analyses. In addition, statistically significant improvements were also demonstrated for all of the pre-defined secondary endpoints for ESC 20mg versus placebo using the LOCF analysis.

Patients were also analysed by baseline severity, whereby severe SAD was defined as a LSAS-J total baseline score $\geq 100$ and a CGI-S score $\geq 6$. Analysis demonstrated statistically improvement of symptoms in non-severe patients taking ESC 10mg. ESC 10mg appeared to be an insufficient dose for patients with severe SAD, whereas ESC 20mg was equally efficacious in both severe and non-severe patients with SAD. These findings suggest that ESC is expected to be effective for non-severe SAD patients at a dose of 10mg/day and that a dose increase to 20mg/day is appropriate for severe SAD patients who do not respond to ESC 10mg/day.

In a placebo-controlled, randomised, double-blind, parallel-group, fixed-dose study in which placebo, ESC 5, 10, or 20mg/day or paroxetine 20mg/day was administered for 24 weeks that was conducted in countries outside of Japan in SAD (Lader et al., 2004)$^8$, the change from treatment initiation in the LSAS total score at week 12 (LOCF, mean) was -29.5 in placebo, -38.7 in ESC 5mg, -34.6 in ESC 10mg, -39.8 in ESC 20mg, and -39.3 in paroxetine. There were no major differences between the studies conducted in Japan and the studies conducted outside Japan in the differences between placebo and either ESC 10mg or ESC 20mg in the change in the LSAS total score at week 12.

There was no marked difference between the ESC doses in the incidence of common TEAEs during treatment, and thus no indication of any dose effect. Somnolence, nausea and ejaculation disorder were reported by a greater proportion of patients treated with ESC 10mg and ESC 20mg than with placebo. Almost all of the TEAEs
that resulted in study treatment discontinuation in all of the ESC groups were mild or moderate, and no major differences were found among the groups in the incidences thereof.

All of the suicide-related TEAEs that occurred were mild or moderate in severity, and no major differences were found between the ESC group and the placebo group in the incidence of suicide-related TEAEs.

ESC is primarily metabolised by CYP2C19, and approximately 20% of the Japanese people are CYP2C19 poor metabolisers. In this study, the percentage of CYP2C19 genotype poor metabolisers was 19.3%. Because it has been shown that the $\text{AUC}_{0-\infty}$ of plasma ESC in CYP2C19 poor metabolisers is approximately twice that of extensive metabolisers\(^{14}\), the safety of ESC according to CYP2C19 genotype was examined. There were no safety or tolerability issues for poor metabolisers compared to extensive metabolisers, and this included those patients with TEAEs or QTcF interval changes. Collectively, these data demonstrate that there were no major clinical problems regarding the safety findings obtained in this study.

There are several limitations to be noted regarding this study. The inclusion and exclusion criteria may limit the generalizability of the study, and the case comorbidity may sometimes visit the medical institution by an actual clinical situation.

In a 12-week, placebo-controlled, randomised, double-blind, parallel-group, variable-dose study of ESC (10 or 20mg/day) study of patients with SAD in countries outside of Japan (Kasper et al., 2005)\(^6\), the following TEAEs occurred in $\geq 5\%$ of patients in the ESC group: headache, nausea, fatigue, somnolence, diarrhoea, insomnia, dizziness, rhinitis, sweating increased, ejaculation failure, and libido decreased and in the placebo group the following TEAEs occurred in $\geq 5\%$ of patients: headache, nausea, fatigue, somnolence, diarrhoea, insomnia, dizziness and rhinitis.
There were no major differences in the TEAEs reported by patients in the two studies. The proportion of patients who discontinued due to TEAEs in the Lader et al. study was 4.5% in the placebo group and 8.8% in the ESC group, and it thus appears that there were no major differences in safety or tolerability between studies conducted in Japan or other countries.

Conclusions
This study demonstrated the efficacy of ESC 10mg/day and 20mg/day in patients with SAD in Japan, as well as the safety and good tolerability of both doses of ESC.

Transparency
Declaration of funding
Mochida Pharmaceutical Co., Ltd sponsored the study and was involved in the study design, in the collection, analysis and interpretation of data, in the writing of the report, and in the decision to submit the paper for publication.

Statement of interest
All authors designed the original study and wrote the protocol. TH monitored study progress. AH undertook the statistical analysis. All authors contributed to and have approved the final manuscript.

Declaration of financial/other relationships
TH and AH are employees of Mochida Pharmaceutical Co., Ltd. SA and TK served as medical experts for this clinical study. The expenses in preparing this paper were met by Mochida Pharmaceutical Co., Ltd.
Acknowledgments

We thank the following investigators for their participation in this study: Dr. Hiroki Ishikawa, Dr. Masahiko Fujita, Dr. Kunihiro Kawamura, Dr. Kyoko Kawaharata, Dr. Kimihiro Nakajima, Dr. Takeshi Fujita, Dr. Michihiro Shimode, Dr. Motomichi Shimizu, Dr. Ken-ichi Harada, Dr. Hideo Maeda, Dr. Junko Shiraki, Dr. Keiji Kaneta, Dr. Naofumi Kusaka, Dr. Shun-ichi Munakata, Dr. Hirofumi Kanome, Dr. Yoshinori Watanabe, Dr. Masatoshi Yaginuma, Dr. Toshiaki Shichijo, Dr. Hiroshi Nagao, Dr. Masaharu Asakawa, Dr. Hisashi Miyake, Dr. Masataka Noguchi, Dr. Gaku Ishikawa, Dr. Shojiro Takahashi, Dr. Il Bong Lee, Dr. Kenji Shigemori, Dr. Satoshi Nakamura, Dr. Eiji Yoshida, Dr. Hiroko Sakamoto, Dr. Jun Matsushima, Dr. Takuji Hishinuma, Dr. Masami Tanaka, Dr. Naoki Harikae, Dr. Nobutoshi Kariya, Dr. Isao Fukunishi, Dr. Jun-ichi Aoki, Dr. Naohiro Yokoyama, Dr. Masaki Kishiro, Dr. Ken-ichi Goto, Dr. Yasutaka Tawara, Dr. Kotaro Kudo, Dr. Takashi Hayama, Dr. Hiroaki Harai, Dr. Keiichi Tanaka, Dr. Tokuji Hokuto, Dr. Yuji Ikeuchi, Dr. Kiyoshi Kohduki, Dr. Takashi Taguchi, Dr. Kenji Tatsuta, Dr. Masanori Kyō, Dr. Seigen Gon, Dr. Norihide Ensako, Dr. Osamu Yamamoto, Dr. Seiichi Tsuji, Dr. Tokumi Fujikawa, Dr. Soichiro Hirayama, Dr. Akio Mantani, Dr. Norio Yokota, Dr. Hiroshi Yamaguchi, Dr. Masamiki Kimura, Dr. Nobuhiko Imato, Dr. Hiromitsu Kaku, Dr. Naoyuki Hamada, Dr. Kazuhide Takada, Dr. Hideyo Sugahara, Dr. Masaru Yoshimura, Dr. Junko Hatakeyama, Dr. Hikari Mizuma, Dr. Hiroaki Furui, Dr. Kaori Hamada, Dr. Masahiro Matsunaga, Dr. Tetsuya Tachiyama, Dr. Joji Kobayashi, Dr. Naoki Kojima, Dr. Hideaki Sakai, Dr. Kunio Kato, Dr. Tetsuo Abe, Dr. Masanobu Takeuchi, Dr. Taihei Fukuhara, Dr. Akihiro Kakishima, Dr. Yoko Ueda, Dr. Masashi Yoshida, Dr. Takashi Furune, Dr. Kahori Ito, and Dr. Naoshige Matsuguchi.
References
10. ICH Harmonised Tripartite Guideline E9: Statistical Principles for Clinical Trials 1998. Available at:


Table 1. Baseline patient characteristics (mean ± SD)

<table>
<thead>
<tr>
<th>APTS</th>
<th>Placebo (n=196)</th>
<th>Escitalopram 10mg (n=198)</th>
<th>Escitalopram 20mg (n=193)</th>
<th>Total (n=587)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender (% female)</td>
<td>55.6</td>
<td>56.6</td>
<td>54.9</td>
<td>55.7</td>
</tr>
<tr>
<td>Mean weight (kg)</td>
<td>58.21 ± 11.79</td>
<td>58.45 ± 11.33</td>
<td>59.06 ± 13.04</td>
<td>58.57 ± 12.05</td>
</tr>
<tr>
<td>Mean height (cm)</td>
<td>163.85 ± 7.99</td>
<td>164.08 ± 8.39</td>
<td>163.96 ± 8.48</td>
<td>163.96 ± 8.28</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>21.56 ± 3.37</td>
<td>21.65 ± 3.50</td>
<td>21.86 ± 3.86</td>
<td>21.69 ± 3.58</td>
</tr>
<tr>
<td>Mean age, range (years)</td>
<td>33.0 (18-63)</td>
<td>33.6 (18-62)</td>
<td>32.5 (18-64)</td>
<td>33.0 (18-64)</td>
</tr>
<tr>
<td>Age at SAD onset (years)</td>
<td>18.8 ± 9.6</td>
<td>18.8 ± 9.0</td>
<td>18.8 ± 8.8</td>
<td>18.8 ± 9.1</td>
</tr>
<tr>
<td>Duration of SAD (years)</td>
<td>14.2 ± 10.5</td>
<td>14.7 ± 10.1</td>
<td>13.7 ± 10.6</td>
<td>14.2 ± 10.4</td>
</tr>
<tr>
<td>History of pharmacotherapy for SAD (%)</td>
<td>57.1</td>
<td>58.6</td>
<td>56.5</td>
<td>57.4</td>
</tr>
<tr>
<td>1 drug* (%)</td>
<td>21.4</td>
<td>30.8</td>
<td>27.5</td>
<td>26.6</td>
</tr>
<tr>
<td>2 drugs* (%)</td>
<td>7.7</td>
<td>5.1</td>
<td>3.1</td>
<td>5.3</td>
</tr>
<tr>
<td>3 drugs* (%)</td>
<td>3.6</td>
<td>2.5</td>
<td>3.1</td>
<td>3.1</td>
</tr>
<tr>
<td>4 drugs* (%)</td>
<td>0.5</td>
<td>1.0</td>
<td>0</td>
<td>0.5</td>
</tr>
<tr>
<td>CYP2C19 genotype (% EM)</td>
<td>80.6</td>
<td>82.8</td>
<td>78.8</td>
<td>80.7</td>
</tr>
<tr>
<td>Comorbid psychiatric disorder (%)</td>
<td>14.3</td>
<td>18.2</td>
<td>20.2</td>
<td>17.5</td>
</tr>
<tr>
<td>Mean baseline LSAS-J total score</td>
<td>95.3 ± 18.5</td>
<td>94.5 ± 18.2</td>
<td>93.4 ± 17.8</td>
<td>94.4 ± 18.1</td>
</tr>
<tr>
<td>Fear/anxiety subscale</td>
<td>51.4 ± 9.1</td>
<td>51.1 ± 9.3</td>
<td>50.5 ± 8.9</td>
<td>51.0 ± 9.1</td>
</tr>
<tr>
<td>Avoidance subscale</td>
<td>43.9 ± 10.7</td>
<td>43.4 ± 10.5</td>
<td>42.8 ± 10.5</td>
<td>43.4 ± 10.5</td>
</tr>
<tr>
<td>Mean baseline MADRS total score</td>
<td>3.6 ± 4.0</td>
<td>3.6 ± 3.9</td>
<td>3.9 ± 3.8</td>
<td>3.7 ± 3.9</td>
</tr>
<tr>
<td>Mean baseline CGI-S</td>
<td>4.8 ± 0.8</td>
<td>4.8 ± 0.9</td>
<td>4.9 ± 0.8</td>
<td>4.8 ± 0.8</td>
</tr>
</tbody>
</table>

* Number of selective serotonin reuptake inhibitors (SSRIs) or serotonin noradrenaline reuptake inhibitors (SNRIs)

anxiety disorder
Table 2. Summary of LSAS-J efficacy assessments (mean ± SD) (FAS)

<table>
<thead>
<tr>
<th>Group</th>
<th>N</th>
<th>LSAS-J total score</th>
<th>Comparison with placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Baseline</td>
<td>Week 12</td>
</tr>
<tr>
<td><strong>ANCOVA (LOCF)</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>196</td>
<td>95.3 ± 18.5</td>
<td>72.2 ± 27.4</td>
</tr>
<tr>
<td>10mg/day</td>
<td>198</td>
<td>94.5 ± 18.2</td>
<td>67.6 ± 29.0</td>
</tr>
<tr>
<td>20mg/day</td>
<td>193</td>
<td>93.4 ± 17.8</td>
<td>60.7 ± 28.0</td>
</tr>
<tr>
<td><strong>ANCOVA (OC)</strong>&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>175</td>
<td>94.7 ± 18.3</td>
<td>69.4 ± 26.6</td>
</tr>
<tr>
<td>10mg/day</td>
<td>177</td>
<td>93.5 ± 18.1</td>
<td>63.4 ± 27.2</td>
</tr>
<tr>
<td>20mg/day</td>
<td>171</td>
<td>93.6 ± 17.8</td>
<td>58.4 ± 27.7</td>
</tr>
<tr>
<td><strong>MMRM</strong>&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>196</td>
<td>95.3 ± 18.5</td>
<td>69.4 ± 26.6</td>
</tr>
<tr>
<td>10mg/day</td>
<td>198</td>
<td>94.5 ± 18.2</td>
<td>63.4 ± 27.2</td>
</tr>
<tr>
<td>20mg/day</td>
<td>193</td>
<td>93.4 ± 17.8</td>
<td>58.4 ± 27.7</td>
</tr>
<tr>
<td><strong>ANCOVA (LOCF) excluding patients discontinued in the first week</strong>&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>195</td>
<td>95.2 ± 18.5</td>
<td>72.0 ± 27.4</td>
</tr>
<tr>
<td>10mg/day</td>
<td>189</td>
<td>94.3 ± 18.3</td>
<td>66.0 ± 28.4</td>
</tr>
<tr>
<td>20mg/day</td>
<td>188</td>
<td>93.6 ± 17.8</td>
<td>60.0 ± 27.9</td>
</tr>
</tbody>
</table>

<sup>a</sup>Bre-specified primary endpoint, <sup>b</sup>Pre-specified sensitivity analysis, <sup>c</sup>Post-hoc analysis, 95% CI: 95% confidence interval, ANCOVA: analysis of covariance, FAS: full-analysis set, LOCF: last observation carried forward, LSAS-J: Liebowitz social anxiety scale - Japanese version, MMRM: mixed model repeated measures, OC: observed cases.
Table 3. Summary of LSAS-J subscale efficacy assessments (mean ± SD) (FAS, LOCF)

<table>
<thead>
<tr>
<th>Group</th>
<th>N</th>
<th>LSAS-J subscale total score</th>
<th>Comparison with placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Baseline</td>
<td>Week 12</td>
</tr>
<tr>
<td>Fear/anxiety subscale</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>196</td>
<td>51.4 ± 9.1</td>
<td>39.9 ± 13.8</td>
</tr>
<tr>
<td>10mg/day</td>
<td>198</td>
<td>51.1 ± 9.3</td>
<td>37.6 ± 14.8</td>
</tr>
<tr>
<td>20mg/day</td>
<td>193</td>
<td>50.5 ± 8.9</td>
<td>34.3 ± 14.6</td>
</tr>
<tr>
<td>Avoidance subscale</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>196</td>
<td>43.9 ± 10.7</td>
<td>32.3 ± 14.5</td>
</tr>
<tr>
<td>10mg/day</td>
<td>198</td>
<td>43.4 ± 10.5</td>
<td>30.0 ± 15.2</td>
</tr>
<tr>
<td>20mg/day</td>
<td>193</td>
<td>42.8 ± 10.5</td>
<td>26.4 ± 14.5</td>
</tr>
</tbody>
</table>

* ANCOVA: analysis of covariance

95% CI: 95% confidence interval, FAS: full-analysis set, LOCF: last observation carried forward, LSAS-J: Liebowitz social anxiety scale - Japanese version.
Table 4. Treatment-emergent adverse events (TEAEs) in ≥5% of patients in any treatment group in the 12-week treatment period (APTS)

<table>
<thead>
<tr>
<th>Preferred Term</th>
<th>Placebo (n=196)</th>
<th>Escitalopram 10mg (n=198)</th>
<th>Escitalopram 20mg (n=193)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients with TEAEs</td>
<td>110 (56.1%)</td>
<td>127 (64.1%)</td>
<td>126 (65.3%)</td>
</tr>
<tr>
<td>Somnolence</td>
<td>17 (8.7%)</td>
<td>36 (18.2%)*</td>
<td>43 (22.3%)*</td>
</tr>
<tr>
<td>Nasopharyngitis</td>
<td>35 (17.9%)</td>
<td>33 (16.7%)</td>
<td>32 (16.6%)</td>
</tr>
<tr>
<td>Nausea</td>
<td>12 (6.1%)</td>
<td>29 (14.6%)*</td>
<td>31 (16.1%)*</td>
</tr>
<tr>
<td>Abdominal discomfort</td>
<td>4 (2.0%)</td>
<td>4 (2.0%)</td>
<td>11 (5.7%)</td>
</tr>
<tr>
<td>Headache</td>
<td>15 (7.7%)</td>
<td>10 (5.1%)</td>
<td>9 (4.7%)</td>
</tr>
<tr>
<td>Ejaculation disorder (men)</td>
<td>0</td>
<td>5 (5.8%)*</td>
<td>2 (2.3%)</td>
</tr>
</tbody>
</table>

APTS: all-patients-treated set

*p<0.05 versus placebo (Fisher’s exact test)
**Table 5.** Treatment-emergent adverse events (TEAEs) in ≥2% of patients who did not receive antidepressants in any treatment group in the follow-up period (APTS)

<table>
<thead>
<tr>
<th>Preferred Term</th>
<th>Placebo (n=113)</th>
<th>Escitalopram 10mg (n=114)</th>
<th>Escitalopram 20mg (n=112)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients with TEAEs</td>
<td>12 (10.6%)</td>
<td>22 (19.3%)</td>
<td>27 (24.1%)*</td>
</tr>
<tr>
<td>Dizziness</td>
<td>3 (2.7%)</td>
<td>7 (6.1%)</td>
<td>12 (10.7%)*</td>
</tr>
<tr>
<td>Nasopharyngitis</td>
<td>3 (2.7%)</td>
<td>3 (2.6%)</td>
<td>4 (3.6%)</td>
</tr>
<tr>
<td>Nausea</td>
<td>1 (0.9%)</td>
<td>1 (0.9%)</td>
<td>3 (2.7%)</td>
</tr>
<tr>
<td>Headache</td>
<td>5 (4.4%)</td>
<td>2 (1.8%)</td>
<td>2 (1.8%)</td>
</tr>
</tbody>
</table>

APTS: all-patients-treated set

*p<0.05 versus placebo (Fisher’s exact test)
Figure Legends

Figure 1. Flow chart of patient disposition. ESC: escitalopram, FAS: full-analysis set.

Figure 2. Estimated change in Liebowitz Social Anxiety Scale (LSAS-J) total scores from baseline to Week 12 (FAS, OC by visit) and FAS, LOCF at Week 12. FAS: full-analysis set, LOCF: last observation carried forward, OC: observed cases. Patient numbers at each visit are shown below the x-axis for each treatment group. The pre-specified primary endpoint is at Week 12 (FAS, ANCOVA, LOCF). *p<0.05 versus placebo.
*The adverse events occurred during the taking of placebo
**Multiple answers are available for a reason of discontinuation
***Patient took no study medication

Figure 1.
Figure 2.