A double-blind, randomized, placebo-controlled, active reference study of Lu AA21004 in patients with major depressive disorder

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Abstract
The efficacy, safety, and tolerability of Lu AA21004 vs. placebo using venlafaxine XR as active reference in patients with DSM-IV-TR major depressive disorder (MDD) were evaluated. Lu AA21004 is a novel antidepressant that is a 5-HT3 and 5-HT7 receptor antagonist, 5-HT1A receptor agonist, 5-HT1B receptor partial agonist and inhibitor of the 5-HT transporter in recombinant cell lines. In this 6-wk, multi-site study, 429 patients were randomly assigned (1:1:1:1) to 5 or 10 mg Lu AA21004, placebo or 225 mg venlafaxine XR. All patients had a baseline Montgomery–Åsberg Depression Rating Scale (MADRS) total score $\geq$ 30. The primary efficacy analysis was based on the MADRS total score adjusting for multiplicity using a hierarchical testing procedure starting with the highest dose vs. placebo. Lu AA21004 was statistically significantly superior to placebo ($n=105$) in mean change from baseline in MADRS total score at week 6 ($p<0.0001$, last observation carried forward), with a mean treatment difference vs. placebo of 5.9 (5 mg, $n=108$), and 5.7 (10 mg, $n=100$) points. Venlafaxine XR ($n=112$) was also significantly superior to placebo at week 6 ($p<0.0001$). In total, 30 patients withdrew due to adverse events (AEs) – placebo: four (4%); 5 mg Lu AA21004: three (3%); 10 mg Lu AA21004: seven (7%); and venlafaxine: 16 (14%). The most common AEs were nausea, headache, hyperhidrosis, and dry mouth. No clinically relevant changes over time were seen in the clinical laboratory results, vital signs, weight, or ECG parameters. In this study, treatment with 5 mg and 10 mg Lu AA21004 for 6 wk was efficacious and well tolerated in patients with MDD.

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Key words: Depression, Lu AA21004, MADRS, serotonin receptor affinity, venlafaxine.

Introduction
Lu AA21004 (1-[2-(2,4-dimethyl-phenylsulanyl)-phenyl]-piperazin) is a novel compound under development as an antidepressant (Bang-Andersen et al. 2011) with affinity for the human 5-HT1A, 5-HT1B, 5-HT3 and 5-HT7 receptors and the 5-HT transporter (SERT) (Moore et al. 2008). Based on preclinical data, these affinities are considered to be of clinical relevance and involved in the mechanism of action at therapeutic doses. In vivo, Lu AA21004 increases the extracellular levels of serotonin (5-HT), noradrenaline, dopamine, acetylcholine and histamine in rat prefrontal cortex and hippocampus (Moore et al. 2008).

Lu AA21004 is extensively metabolized in the liver and at least five cytochrome P450 isoenzymes appear to be involved. The metabolism of Lu AA21004 to its major metabolite (pharmacologically inactive) is mediated primarily by CYP2D6. In addition, Lu AA21004 does not seem to be a clinically relevant inhibitor or inducer of cytochrome P450 isoenzymes.
The terminal elimination half-life after multiple doses is estimated at \( \sim 60-70 \) h. The exposure (\( C_{\text{max}} \) and area under curve) increased linearly with dose (2.5–60 mg). The absorption of Lu AA21004 is independent of food intake (Wang et al. 2009) and maximum plasma concentrations are reached 3–16 h after dosing. The rationale for choosing the Lu AA21004 doses (5 and 10 mg) in this proof-of-concept study was based on non-clinical and phase I data. Approximately 60–80% occupancy of the human SERT is required to achieve a therapeutic effect with selective serotonin reuptake inhibitors (SSRIs) or serotonin noradrenaline reuptake inhibitors (SNRIs) (Meyer, 2007). In contrast, an occupancy level of 41% with Lu AA20004 in rats led to a significant increase in extracellular levels of 5-HT, perhaps due to the additional pharmacological activities of Lu AA21004, which may counteract negative feedback mechanisms operating at cellular and network levels. The dose of 5 mg/d corresponds to a SERT occupancy of \( \sim 40\% \) in human brain and was, therefore, expected to be an effective dose (Areberg et al. 2009).

The aim of this phase II clinical study was to investigate the efficacy, safety, and tolerability of two fixed doses (5 and 10 mg/d) of Lu AA21004 vs. that of placebo after 6 wk treatment in adult patients with major depressive disorder (MDD). Venlafaxine XR (225 mg/d) was used as the active reference.

**Method**

This randomized, double-blind, fixed-dose, placebo-controlled, active reference study recruited 429 randomized patients from 49 psychiatric settings in 11 countries (Australia, Austria, Canada, Czech Republic, Finland, France, Italy, Malaysia, Slovakia, Spain, Sweden). Outpatients with MDD were recruited from psychiatric settings from August 2006 to August 2007. Advertisements were used in Australia, Austria, Canada, Finland, Malaysia, and Sweden. The study was conducted in accordance with the principles of Good Clinical Practice (ICH, 1996) and the Declaration of Helsinki (WMA, 1964). Local ethics committees approved the study design and eligible patients gave their written informed consent before participating.

Eligible patients were randomized equally (1:1:1:1) to one of the four treatment arms for a 6-wk double-blind treatment period. Randomized patients were given 1-wk wallet cards at each visit and were instructed to take two capsules per day, orally, at the same time every day (preferably in the morning). Lu AA21004 was dosed at 5 or 10 mg/d for 6 wk and venlafaxine at 75 mg/d for 4 d, 150 mg/d for the following 3 d, and 225 mg/d for the remainder of the treatment period. Efficacy and tolerability were assessed at screening, baseline and after 1, 2, 3, 4, 5, and 6 wk. Patients who completed the 6-wk double-blind treatment period entered a 2-wk double-blind taper period. During this period, patients on 5 mg/d Lu AA21004 switched to placebo; patients on 10 mg/d Lu AA21004 received 5 mg/d Lu AA21004 for the first week (week 7) and placebo for the second week (week 8); patients on placebo remained on placebo; patients on venlafaxine received 150 mg/d venlafaxine for the first week (week 7) and 75 mg/d for the second week (week 8). Patients were contacted for a safety follow-up 4 wk after the completion visit. Down-taper medication was also offered to patients who withdrew.

**Main entry criteria**

Patients with MDD presenting with a current major depressive episode according to DSM-IV-TR criteria (APA, 1994) were included in the study if they were an outpatient of either sex, aged from 18 yr to 65 yr, with a Montgomery–Åsberg Depression Rating Scale (MADRS) (Montgomery & Åsberg, 1979) total score \( \geq 30 \) at the baseline visit.

Patients were excluded if they had any current psychiatric disorder other than MDD as defined in DSM-IV-TR [assessed using the Mini International Neuropsychiatric Interview (MINI; Sheehan et al. 1998)], or if they had a current or past history of manic or hypomanic episode, schizophrenia or any other psychotic disorder, including major depression with psychotic features, mental retardation, organic mental disorders, or mental disorders due to a general medical condition, any substance abuse disorder within the previous 6 months, presence or history of a clinically significant neurological disorder (including epilepsy), any neurodegenerative disorder, or any Axis II disorder that might compromise the study.

Patients at serious risk of suicide, based on the investigator's clinical judgement, or who had a score of \( \geq 5 \) on item 10 of the MADRS scale (suicidal thoughts) were also excluded, as were those receiving formal behaviour therapy or systematic psychotherapy, or were pregnant or breastfeeding, had a known hypersensitivity or were non-response to venlafaxine, or whose current depressive symptoms were considered by the investigator to have been resistant to two adequate antidepressant treatments of at least 6 wk duration, or had previously been exposed to Lu AA21004.
Patients were also excluded if they were taking the following psychotropic drugs within 2 wk prior to baseline or during the study: Reversible or irreversible monoamine oxidase inhibitors, SSRIs (fluoxetine within 5 wk), SNRIs, tricyclic antidepressants, psychoactive herbal remedies, any drug used for augmentation of antidepressant action or any other antidepressant drugs, oral antipsychotic and antimanic drugs, or dopamine antagonists, any anxiolytics (including benzodiazepines); and any anticonvulsant drug, serotonergic agonists, narcotic analgesics or cough agents, anti-arrhythmics, oral anticoagulants, proton pump inhibitors, steroids, cisapride, macrolide antibiotics, antifungal agents, antihypertensives, all anti-inflammatory agents, anti-migraine agents, pseudoephedrine, hypolipidaemics, and episodic use of insulin. Occasional use of zolpidem, zopiclone and zaleplon for insomnia was allowed.

Patients were withdrawn if they became pregnant during the study, if the investigator considered it to be in the best interest of the patient for safety/efficacy reasons, if laboratory values were outside normal ranges and clinically significant, if they were considered to be at significant risk of suicide, if they scored ≥ 5 points on item 10 (suicidal thoughts) of the MADRS, if the randomization code for a patient was broken, if consent to participate was withdrawn, if they did not take study medication for more than 6 consecutive days, or if the patient was lost to follow-up. The patient could be withdrawn from the study if a serious adverse event (SAE) occurred. If adverse events (AEs) were contributory to withdrawal, they were always regarded as the primary reason for withdrawal.

Efficacy rating
Patients were evaluated using the MADRS from baseline to week 6. Rater training was undertaken to increase inter-rater reliability, and was chaired by an experienced investigator. Only those investigators who had actively participated in rater training sessions prior to inclusion of patients into the study and had received rater certification were allowed to rate patients. Patient ratings were assessed by the same investigator at each visit, whenever possible.

Allocation to treatment
The medication was given as capsules of identical appearance. Patients who met the selection criteria at the baseline visit were assigned to double-blind treatment according to a computer-generated randomization list. The details of the randomization series were unknown to any of the investigators and were contained in a set of sealed opaque envelopes. At each study site, sequentially enrolled patients were assigned the lowest randomization number available in blocks of four. All investigators, study personnel and participants were blinded to treatment assignment for the duration of the entire study. The randomization code was broken for one patient (accidentally) who had completed the study before this was discovered, and was therefore not withdrawn from the study.

Analysis sets
All safety analyses were based on the all-patients-treated set (APTS), comprising all randomized patients who took at least one dose of study medication. All efficacy analyses were based on a modified intent-to-treat set (ITT) – the full-analysis set (FAS), comprising all patients in the APTS who had at least one valid post-baseline MADRS total score assessment.

Power and sample size calculations
It was planned to randomize a minimum of 384 patients with a DSM-IV-TR diagnosis of a major depressive episode (MDE) into the double-blind period of the study. With 96 patients in each treatment group and a standard deviation (S.D.) of 9 points, the power to detect a true treatment effect of 3.7 points on the MADRS total score at week 6, using last observation carried forward (LOCF), would be 80%.

Primary efficacy analysis
Four hypotheses were part of the primary efficacy analysis, which was fully adjusted for multiplicity using a hierarchical testing procedure at the 5% level of significance as long as the previous hypothesis was rejected. The order of testing was: no difference between the 10 mg dose vs. placebo at week 6, no difference between 5 mg vs. placebo at week 6, no difference between 10 mg dose vs. placebo at week 1, and finally no difference between 5 mg dose vs. placebo at week 1. The statistical model was an analysis of covariance (ANCOVA) of the change from baseline in MADRS total score (FAS, LOCF) with treatment and site as fixed factors and the baseline MADRS score as a covariate. The primary efficacy analysis was repeated on observed cases (OC) data, using both an ANCOVA and a mixed model for repeated measurements (MMRM).

Secondary efficacy analysis
Prospectively defined secondary clinician-rated variables were: MADRS total score, 24-item Hamilton...
Depression (HAMD_24) total score (Hamilton, 1960), 
Clinical Global Impression – Improvement (CGI-I) 
and Clinical Global Impression – Severity (CGI-S) 
scores (Guy, 1976), Hamilton Anxiety (HAMA) total 
score (Hamilton, 1959), remission [defined as MADRS 
< 10, 17-item HAMD (HAMD_17) < 7 or as a CGI-S 
score < 2] and response (defined as 
< 50% decrease 
from baseline in MADRS or HAMD_24 total score, or a 
CGI-I score < 2) at all time points.

The change from baseline to each visit in all the 
secondary efficacy variables, except response and re-
mission, was analysed using an ANCOVA, adjusting 
for baseline score, site, and treatment, using both 
OC and LOCF data. For CGI-I, the baseline CGI-S 
score was used for adjustment. The change from 
baseline to each visit in all the secondary efficacy 
variables, except response and remission, was also 
analysed using MMRM to compare the treatment 
groups over all assessment points simultaneously 
using OC data.

Response and remission rates for each visit were 
evaluated using Fisher’s exact test. The CGI-S and 
CGI-I scores were analysed at the last visit (OC and 
LOCF) using ANCOVA. Unless otherwise stated, the 
terms ‘significant’ and ‘significantly’ refer to statistical 
significance at the 5% level, two-sided. Efficacy 
analyses that were not multiplicity-controlled were 
considered secondary. The principal statistical soft-
ware used was SAS® version 9.1 (SAS Institute Inc., 
USA).

Tolerability assessments

Each patient was asked a non-leading question 
such as, ‘how do you feel?’ at each visit, starting 
at baseline. All AEs (including any change in concur-
rent illnesses or new illnesses) either observed by 
the investigator or reported spontaneously by the 
patient were recorded. AEs were coded using the 
lowest level term according to the Medical Dictionary 
for Regulatory Activities, version 10.0. The time to 
withdrawal due to AEs was analysed using the Cox 
model. The incidences of individual AEs were com-
pared between the treatment groups using Fisher’s 
effect test.

As a post-hoc analysis, the safety database was 
searched at preferred-term and verbatim-term level 
for possible suicide-related AEs, as described by the 
FDA (Laughren, 2006).

Results

Patient baseline characteristics

The APTS comprised 426 patients (placebo, 105; 
venlafaxine, 113; 5 mg Lu AA21004, 108; 10 mg Lu 
AA21004, 100) (Fig. 1). Slightly more patients than 
planned were enrolled in the study, raising the power 
from 80% to 84%. There were no clinically relevant or 
statistically significant differences between the treat-
ment groups in patient demographics or clinical 
characteristics at baseline (Table 1). Patients had a

Fig. 1. Flow chart of patient disposition. AE, Adverse events, ITT, intention to treat; LoE, lack of efficacy; MADRS, 
Montgomery–Åsberg Depression Rating Scale; PBO, placebo; Ven 225, venlafaxine XR 225 mg.
The mean age (± S.D.) of 43.3 ± 11.5 yr, 62.7% were women, and 92.0% were Caucasian.

The mean baseline MADRS total score was 34.0, indicating a severely depressed patient population, consistent with the mean CGI-S score of 5.1. Patients were diagnosed with their first MDE 10 yr prior to enrolment. Between 74% and 80% of the patients in each treatment group had had a previous MDE and their current episode had started about 5 months prior to enrolment (Table 1). There was a substantial level of anxiety symptoms, as indicated by a mean baseline HAMA total score of 22.2. About 40% (range 36–41%) of the patients in each treatment group had a concurrent medical condition. The number of patients taking zolpidem, zopiclone, or zaleplon prescribed episodically for insomnia was similar for placebo (n = 3), venlafaxine (n = 6), 5 mg Lu AA21005 (n = 3), and 10 mg Lu AA21005 (n = 3). Between 21% and 33% of the patients took concomitant medication that they continued with, and 26–29% commenced concomitant medication during the study.

Withdrawals from the study

The withdrawal rate due to all reasons during the entire study was 15% (Fig. 1), ranging from 9% (5 mg Lu AA21004) to 18% (venlafaxine and 10 mg Lu AA21004). More than 80% of the patients in each treatment group completed the study (Fig. 1). There was a slightly larger proportion of patients who completed the study in the 5 mg Lu AA21004 group than in the placebo, 10 mg Lu AA21004, or venlafaxine groups. The proportions of patients who withdrew due to AEs was statistically significantly different between venlafaxine and placebo, but not between the Lu AA21004 groups and placebo. There was an even distribution of withdrawals for any reason over time and no statistically significant differences between the treatment groups, between men and women, or between patients aged ≤50 or >50 yr. The median compliance with study medication was 98%.

Efficacy

Primary endpoint

On the pre-defined primary efficacy endpoint, both doses of Lu AA21004 were statistically significantly (p < 0.0001) superior to placebo in mean change from baseline in MADRS total score at week 6 (FAS, LOCF), with mean treatment differences to placebo of 5.9 (5 mg) and 5.7 (10 mg) points (Table 2) in a multiplicity-controlled analysis. These differences to placebo correspond to a standardized effect size (Cohen’s d) of 0.56 (5 mg) and 0.54 (10 mg). Venlafaxine was also statistically significantly (p < 0.0001) superior to placebo at week 6, with a mean treatment difference to placebo of 6.4 points (LOCF). The estimated treatment differences and nominal p values at week 6 obtained from an analysis using MMRM were similar to those

Table 1. Baseline patient characteristics

|                  | Placebo (n = 105) | Lu AA21004 5 mg (n = 108) | Lu AA21004 10 mg (n = 100) | Venlafaxine 225 mg (n = 113) |
|------------------|-------------------|---------------------------|---------------------------|-----------------------------|
| Women            | 69 (65.7%)        | 70 (64.8%)                | 66 (66.0%)                | 62 (54.9%)                  |
| Age (yr)         |                   |                           |                           |                             |
| Mean ± S.D.      | 42.0 ± 10.9       | 43.8 ± 11.6               | 42.3 ± 13.1               | 45.0 ± 10.3                 |
| Range            | 20–61             | 20–64                     | 18–65                     | 21–63                       |
| Caucasian        | 93.3%             | 93.5%                     | 89.0%                     | 92.0%                       |
| Patients with first MDE | 20.0% | 22.2% | 26.0% | 25.7% |
| Years since first MDE ± S.D. | 10 ± 8 | 10 ± 8 | 9 ± 9 | 11 ± 9 |
| Days since start of current MDE ± S.D. | 176 ± 82 | 161 ± 60 | 163 ± 68 | 163 ± 68 |
| Efficacy scoresa (n = 105) |               | (n = 108)                | (n = 100)                 | (n = 112)                   |
| MADRS total score ± S.D. | 33.9 ± 2.7       | 34.1 ± 2.6                | 34.0 ± 2.8                | 34.2 ± 3.1                  |
| HAMDa ± S.D.     | 29.7 ± 5.0        | 29.9 ± 5.4                | 29.3 ± 5.6                | 29.4 ± 5.0                  |
| HAMA total score ± S.D. | 22.9 ± 5.9       | 21.7 ± 6.2                | 22.3 ± 5.6                | 22.0 ± 5.5                  |
| CGI-S ± S.D.     | 5.1 ± 0.7         | 5.2 ± 0.7                 | 5.1 ± 0.7                 | 5.2 ± 0.7                   |

* Based on the full-analysis set: CGI-S, Clinical Global Impression – Severity; HAMA, Hamilton Rating Scale for Anxiety; HAMDa, Hamilton Rating Scale for Depression (24 items); MADRS, Montgomery–Åsberg Depression Rating Scale; MDE, major depressive episode; S.D., standard deviation.
obtained in the ANCOVA analyses \(5.6 \pm 1.3\) (5 mg Lu AA21004), \(7.2 \pm 1.4\) (10 mg Lu AA21004), \(7.6 \pm 1.3\) (venlafaxine), all \(p < 0.0001\) (Table 2). As a sensitivity analysis, the non-parametric Kruskal–Wallis test showed a statistically significant difference between the active treatments and placebo. The assumption of homogeneity of variances across treatment groups was confirmed using Bartlett’s test \(p = 0.90\). At week 1, with a difference from placebo in the MADRS total score of 0.8 for 10 mg \(p = 0.2377\) and 0.2 for 5 mg \(p = 0.7489\), none of the active treatments separated significantly from placebo.

Secondary efficacy analyses

MADRS

The mean MADRS total score decreased in all active treatment groups from 34.1 at baseline to \(~13.4\) in the LOCF analysis and to \(~9.7\) in the OC analysis at week 6. For Lu AA21004, a statistically significant difference compared to placebo in the change from baseline in MADRS total score, in favour of Lu AA21004, was seen from week 2 (10 mg) or week 3 (5 mg) onwards (LOCF and OC). For venlafaxine, a statistically significant difference to placebo was seen from week 2 (OC) or week 3 (LOCF) onwards (Fig. 2). At week 6, the proportion of MADRS responders (patients with \(~50\%\) decrease in MADRS total score) and remitters (MADRS score \(~10\) was statistically significantly higher in all active treatment groups than placebo (LOCF and OC) (Table 3). Single item analysis at week 6 showed a statistically significant advantage for both doses of Lu AA21004 for 9 out of the 10 items (except for ‘concentration difficulties’) relative to placebo.

HAMD

The mean HAMD\(_{24}\) total score decreased in all active treatment groups from 29.5 at baseline to \(~11.9\) in the LOCF analysis and \(~9.7\) in the OC analysis at week 6.

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**Table 2. Change from baseline in MADRS total score at week 6 (FAS)**

| Analysis | Treatment group | Mean ± S.E. | Difference to placebo | \(p\) value |
|----------|-----------------|-------------|-----------------------|-------------|
| LOCF, ANCOVA | Placebo \((n = 105)\) | \(-14.5 \pm 1.0\) | – | – |
| | Lu AA21004 5 mg \((n = 108)\) | \(-20.4 \pm 1.0\) | \(-5.9 \pm 1.4\) | \(< 0.0001\) |
| | Lu AA21004 10 mg \((n = 100)\) | \(-20.2 \pm 1.0\) | \(-5.7 \pm 1.4\) | \(< 0.0001\) |
| | Venlafaxine \((n = 112)\) | \(-20.9 \pm 1.0\) | \(-6.4 \pm 1.4\) | \(< 0.0001\) |
| OC, ANCOVA | Placebo \((n = 88)\) | \(-16.6 \pm 1.0\) | – | – |
| | Lu AA21004 5 mg \((n = 99)\) | \(-22.3 \pm 0.9\) | \(-5.7 \pm 1.3\) | \(< 0.0001\) |
| | Lu AA21004 10 mg \((n = 83)\) | \(-23.4 \pm 1.0\) | \(-6.8 \pm 1.3\) | \(< 0.0001\) |
| | Venlafaxine \((n = 95)\) | \(-24.2 \pm 0.9\) | \(-7.6 \pm 1.3\) | \(< 0.0001\) |
| MMRM | Placebo \((n = 88)\) | \(-15.7 \pm 1.0\) | – | – |
| | Lu AA21004 5 mg \((n = 99)\) | \(-21.3 \pm 0.9\) | \(-5.6 \pm 1.3\) | \(< 0.0001\) |
| | Lu AA21004 10 mg \((n = 83)\) | \(-22.9 \pm 1.1\) | \(-7.2 \pm 1.4\) | \(< 0.0001\) |
| | Venlafaxine \((n = 95)\) | \(-23.4 \pm 0.9\) | \(-7.6 \pm 1.3\) | \(< 0.0001\) |

FAS, Full-analysis set; LOCF, last observation carried forward; MADRS, Montgomery–Åsberg Depression Rating Scale; MMRM, mixed model repeated measures; OC, observed cases; S.E., standard error of the mean.

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**Fig. 2.** Mean change from baseline in Montgomery–Åsberg Depression Rating Scale (MADRS) total scores (ANCOVA, FAS, OC, over time) and LOCF (week 6). * \(p < 0.05\), ** \(p < 0.01\), *** \(p < 0.001\) vs. placebo. FAS, Full-analysis set; LOCF, last observation carried forward; OC, observed cases.
For Lu AA21004 (5 mg and 10 mg), a statistically significant difference to placebo was seen from week 1 onwards. For venlafaxine, a statistically significant difference to placebo was seen from week 2 (OC) or week 3 (LOCF) onwards. At week 6, the proportion of HAMD$_{24}$ responders (patients with ≥50% decrease in HAMD$_{24}$ total score) and remitters (HAMD$_{17}$ score ≤7) was statistically significantly higher in all active treatment groups compared to placebo (LOCF and OC) (Table 3).

### HAMA

The level of anxiety symptoms, as assessed by the mean HAMA total score decreased in all active treatment groups from ~22 at baseline to ~10.1 in the LOCF analysis (Table 4) and ~8.4 in the OC analysis at week 6. For Lu AA21004, a statistically significant difference to placebo was seen in change from baseline in HAMA total score from week 2 (10 mg, OC) or week 3 (LOCF and OC) onwards (Fig. 3). For venlafaxine, a statistically significant difference to placebo was seen from week 3 (OC) or week 4 (LOCF) onwards.

### CGI

The mean CGI-S score decreased in all active treatment groups from ~5.2 at baseline to ~2.6 in the LOCF analysis (Table 4) and ~2.3 in the OC analysis at week 6. The mean CGI-I score improved in all active treatment groups from ~3.7 at baseline to ~2.4 in the LOCF analysis (Table 4) and ~2.2 in the OC analysis at week 6.

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**Table 3. Proportion (%) of responders and remitters at week 6 (FAS, mean)**

|                    | Placebo (n = 105) | Lu AA21004 5 mg (n = 108) | Lu AA21004 10 mg (n = 100) | Venlafaxine 225 mg (n = 112) |
|--------------------|-------------------|---------------------------|--------------------------|----------------------------|
| **Response**       |                   |                           |                          |                            |
| ≥50% MADRS        | 45                | 67**                      | 68**                     | 72**                       |
| ≥50% HAMD$_{24}$  | 40                | 72***                     | 69***                    | 72**                       |
| CGI-I ≤2          | 49                | 73***                     | 76**                     | 77***                      |
| **Remission**     |                   |                           |                          |                            |
| MADRS ≤10         | 27                | 49**                      | 49**                     | 57**                       |
| HAMD$_{17}$ ≤7    | 28                | 47**                      | 45*                      | 53**                       |
| CGI-S ≤2          | 26                | 45**                      | 50***                    | 54**                       |

**Table 4. Mean change from baseline in efficacy variables at week 6, difference to placebo (FAS)**

| Efficacy variable | LOCF, ANCOVA | OC, ANCOVA | MMRM |
|-------------------|--------------|------------|------|
|                   | Lu AA21004 5 mg | Lu AA21004 10 mg | Lu AA21004 5 mg |
| MADRS             | −5.9***      | −6.4***    | −7.6*** |
| HAMD$_{24}$       | −5.3***      | −5.1***    | −5.9*** |
| HAMA              | −3.3***      | −2.9**     | −3.5*** |
| CGI-S             | −0.9***      | −1.0***    | −1.2*** |
| CGI-I             | −0.6***      | −0.7***    | −0.9*** |
AEs than in the placebo group (statistically significantly more patients withdrew due to AEs: four (4%) in the placebo group, three (3%) in the 5 mg Lu AA21004 group, seven (7%) in the 10 mg Lu AA21004 group, and four (4%) in the venlafaxine group. Only in the venlafaxine group, did statistically significantly more patients withdraw due to AEs related to sexual dysfunction: one from each of the Lu AA21004 groups, and three (3%) in the 10 mg Lu AA21004 group. Of the men (n = 101), one from each of the Lu AA21004 groups, and three (3%) in the 10 mg Lu AA21004 group, and four (4%) in the venlafaxine group. Of the women (n = 104), all 11 were from the venlafaxine group. Of the men (n = 159), all 11 were from the venlafaxine group, in which the incidence of AEs related to sexual dysfunction was statistically significantly higher than placebo (12.4% vs. 1.9%, p = 0.0033, Fisher’s exact test). Two patients withdrew due to AEs related to sexual dysfunction; one due to anorgasmia and 1 due to delayed ejaculation, both from the venlafaxine group.

No possibly suicide-related AEs were found in the database search during the entire study. A decrease in MADRS item 10 score (suicidal thoughts) from treatment groups to ~2.0 at week 6 (LOCF, Table 4). For Lu AA21004, a statistically significant difference to placebo was seen in mean CGI-I score from week 1 (10 mg) or week 2 (5 mg) onwards (LOCF). For venlafaxine, a statistically significant difference to placebo was seen from week 3 onwards (LOCF). At week 6, the proportion of CGI responders (CGI-I ≤ 2) and CGI remitters (CGI-S ≤ 2) was statistically significantly higher in all active treatment groups than placebo (LOCF and OC) (Table 3).

### Tolerability and safety

**AEs**

Since Lu AA21004 is a compound with a new mode of action, its safety and tolerability profile is described in some detail below. During the 6-wk treatment period, approximately three-fifths of patients in the placebo (61%) and 5 mg Lu AA21004 (68%) groups and approximately three-quarters of the patients in the 10 mg Lu AA21004 (74%) and venlafaxine (75%) groups had one or more AE. A total of 30 (7%) patients withdrew due to AEs: four (4%) in the placebo group, three (3%) in the 5 mg Lu AA21004 group, seven (7%) in the 10 mg Lu AA21004 group, and 16 (14%) in the venlafaxine group. Only in the venlafaxine group, did statistically significantly more patients withdraw due to AEs than in the placebo group (p = 0.009). Seven patients withdrew from the study due to nausea: three (3%) in the 10 mg Lu AA21004 group and four (4%) in the venlafaxine group.

AEs reported by ≥5% of patients during the 6-wk treatment period are shown in Table 5. The most common AEs reported in the active treatment groups were nausea, headache, hyperhidrosis, and dry mouth. For Lu AA21004, nausea (5 and 10 mg), hyperhidrosis (10 mg), and vomiting (10 mg) were the only AEs reported with an incidence statistically significantly higher than placebo. For the majority of patients reporting nausea, it was transient and mild or moderate in intensity. In addition to nausea and hyperhidrosis, the incidence of dry mouth, constipation, and anorgasmia were statistically significantly higher in the venlafaxine group than placebo group.

In all treatment groups, the majority of patients who had AEs, had mild or moderate AEs. The incidence of severe AEs was 4% in the placebo group, 6% in the Lu AA21004 groups, and significantly higher at 12% in the venlafaxine group (p = 0.026, Fisher’s exact). Severe AEs reported by at least two patients in any Lu AA21004 treatment group included: severe headache by three patients (3%) in the 10 mg Lu AA21004 group and two patients (1.9%) in the placebo group, and two patients (1.9%) in the 5 mg Lu AA21004 group had severe fatigue. In addition, severe AEs reported by at least two patients in the venlafaxine group were: severe nausea and severe vomiting, each reported by two patients (1.8%), severe insomnia in four patients (3.5%), severe dizziness in three patients (2.7%), and severe hyperhidrosis in two patients (1.8%).

For patients treated with both Lu AA21004 doses, the incidence of AEs related to sexual dysfunction (anorgasmia, delayed ejaculation, erectile dysfunction, decreased libido, impotence, abnormal orgasm, abnormal sexual function) was at placebo level [1.9% (5 mg) and 1.0% (10 mg) vs. 1.9% (placebo)]. In total, 23 AEs related to sexual dysfunction were reported by 18 patients, comprising seven women and 11 men. Of the women (n = 267), two were in the placebo group, one from each of the Lu AA21004 groups, and three from the venlafaxine group. Of the men (n = 159), all 11 were from the venlafaxine group, in which the incidence of AEs related to sexual dysfunction was statistically significantly higher than placebo (12.4% vs. 1.9%, p = 0.0033, Fisher’s exact test). Two patients withdrew due to AEs related to sexual dysfunction; one due to anorgasmia and 1 due to delayed ejaculation, both from the venlafaxine group.

Fig. 3. Mean change from baseline in Hamilton Rating Scale for Anxiety (HAMA) total scores (ANCOVA, FAS, OC, over time) and LOCF (week 6). * p < 0.05, ** p < 0.01, *** p < 0.001 vs. placebo. FAS, Full-analysis set; LOCF, last observation carried forward; OC, observed cases. Some patients were excluded due to the use of a non-validated scale in France.
baseline was seen in all treatment groups at all weeks. A numerical superiority over placebo was seen in all active treatment groups from week 2 onwards.

**SAEs**

No deaths occurred during the study. Three patients had SAEs: two in the 10 mg Lu AA21004 group (one patient with worsening of MDD, and one patient with Varicella zoster infection) and one in the venlafaxine group (brain tumour).

**Vital signs, weight, clinical laboratory values, ECGs**

No consistent trends were observed for vital signs, weight, clinical laboratory values or ECG in the active treatment groups, and there were no marked differences between patients receiving active treatment and those receiving placebo. The incidence of potentially clinically significant (PCS) values was generally low and evenly distributed among the treatment groups for vital signs, weight or clinical laboratory values, and no patients withdrew due to a PCS value.

All mean vital signs were within the reference ranges and the mean changes from screening were generally small [≤2 mmHg (supine diastolic blood pressure), ≤5 mmHg (supine systolic blood pressure), or ≤4 bpm (supine pulse)]. The mean weight change from baseline to week 6 was ±0.3 kg in the Lu AA21004 and placebo groups and −0.8 kg in the venlafaxine group, which was not considered to be clinically relevant. Weight gain (≥7%) was recorded for one placebo patient and three patients in the 10 mg Lu AA21004 group, whereas weight loss (≥7%) was recorded for two patients in the 10 mg Lu AA21004 group, and one patient in the venlafaxine group. No patients withdrew due to weight change.

The mean changes in clinical laboratory values were small and similar between treatment groups and the incidence of PCS values was generally <2% in any treatment group for any laboratory test. No clinically relevant abnormalities in ECG values were found after administration of Lu AA21004.

**Discussion**

This is the first double-blind, randomized, placebo-controlled study to evaluate the efficacy, safety and tolerability of Lu AA21004 in patients with MDD. The active reference, venlafaxine XR (225 mg), was included with the purpose of validating the study methodology and patient population, and was effective on the primary efficacy analysis. Both doses of Lu AA21004 resulted in a significant improvement.
compared to placebo on the primary efficacy analysis. It has been suggested (Moncrieff & Kirsch, 2005) that the difference in total scores for an active treatment vs. placebo can be driven by a few single individual items in a rating scale. However, this is not the case in the present study, in which both doses of Lu AA21004 showed significantly greater efficacy than placebo on nine of the 10 MADRS items.

There is a large difference to placebo for all active treatment groups of about 5–6 points on the HAMD\textsubscript{24} (which translates to ~4 points on the HAMD\textsubscript{11}), which is more than the ~2 points on the HAMD\textsubscript{17} seen in FDA pivotal antidepressant studies (Kirsch et al. 2002). This also confirms the assay sensitivity of the studied population, who were not only severely depressed, but also had a substantial level of anxiety symptoms at baseline. At week 6, the proportion of MADRS responders (patients with ≥50% decrease in MADRS total score) and remitters (MADRS score ≤10) was statistically significantly greater in all active treatment groups than in placebo (LOCF and OC). The difference between active treatment and placebo of ~6 points on the MADRS translates into a clinically relevant difference in response rates of between 22% and 32% units, compared to an average of 16% units for antidepressants approved by the competent European authorities (Melander et al. 2008). The robustness of the results was also confirmed by the significantly better outcome than placebo on HAMD\textsubscript{24}, HAMA, CGI-I and CGI-S.

Several pharmacological mechanisms are likely to account for the multimodal antidepressant action of Lu AA21004. It has been estimated that an 80% occupancy of the human SERT is achieved at standard doses of SSRIs or SNRLs (Meyer, 2007). However, 5 mg Lu AA21004 occupies ~40% of SERT sites, suggesting the existence of additional mechanisms involved in its therapeutic activity. Hence, SERT blockade by SSRIs evokes a series of negative feedback mechanisms that attenuate the increase in extracellular (synaptic) concentration of 5-HT, including the activation of 5-HT\textsubscript{1A} and 5-HT\textsubscript{1B} autoreceptors on serotonergic neurons (Artigas et al. 1996, 2001). The partial agonist activity of Lu AA21004 at 5-HT\textsubscript{1B} receptors may therefore help counteract the inhibition of terminal 5-HT synthesis and release evoked by 5-HT\textsubscript{1B} receptor activation. Likewise, its full agonist activity at human 5-HT\textsubscript{1A} receptors expressed in cell lines is predicted to evoke a rapid desensitization of 5-HT\textsubscript{1A} autoreceptors (Haddjeri et al. 2009), thereby normalizing serotonergic cell firing and 5-HT release. On the other hand, given the presence of excitatory 5-HT\textsubscript{3} receptors in GABAergic interneurons in cortical and limbic areas (Morales et al. 1996; Puig et al. 2004), their activation by 5-HT may induce a GABA-mediated inhibition of neurotransmitter release. In support of this view, the 5-HT\textsubscript{3} receptor antagonist ondansetron augments the increase of extracellular 5-HT in the ventral hippocampus induced by the SSRI paroxetine (Mørk et al. 2009). Moreover, blockade of 5-HT\textsubscript{1A} receptors has been shown to produce rapid antidepressant-like effects in the rat in behavioural and electrophysiological experimental paradigms (Mnie-Filali et al. 2011). In addition to these effects on the 5-HT system, the systemic administration of Lu AA21004 increases the extracellular concentration of dopamine, noradrenaline and acetylcholine (Haddjeri et al. 2009), an effect probably contributing to its antidepressant activity.

Due to the profile of Lu AA21004, a hierarchical procedure was used to test for onset of action at week 1. Although not significant on the MADRS, Lu AA21004 displayed onset of antidepressant action, with significant improvement vs. placebo at week 1 onwards for both doses on HAMD\textsubscript{24} and for the 10 mg dose on CGI-I.

The proportion of patient withdrawals has been used in recent years as an indirect index of drug effectiveness in the real world (Kahn et al. 2008; Lieberman et al. 2005; Trivedi et al. 2006). The analysis of withdrawal rates in patients treated with Lu AA21004 indicates a better tolerability profile compared to the active reference, venlafaxine. Compared to placebo, significantly more patients withdrew due to AEs only in the venlafaxine group.

The most common AEs reported in the active treatment groups were nausea, headache, hyperhidrosis, and dry mouth. No possibly suicide-related AEs were found. No consistent trends were observed for vital signs, weight, clinical laboratory values or ECG in the active treatment groups, and there were no marked differences between patients receiving active treatment and those receiving placebo.

Sexual dysfunction during antidepressant treatment is one of the main reasons for the lack of compliance (Kennedy & Rizvi, 2009). According to the present data, the incidence of spontaneously reported AEs related to sexual dysfunction was similar to placebo in patients treated with either dose of Lu AA21004. In the venlafaxine group, the incidence of AEs related to sexual dysfunction was significantly higher than that of placebo (12.4% vs. 1.9%). Unlike SSRIs, Lu AA21004 also displays moderate to high affinity for 5-HT\textsubscript{1A}, 5-HT\textsubscript{1B}, 5-HT\textsubscript{3}, and 5-HT\textsubscript{1B} receptors (see above). There is limited information on the role of these receptors on sexual drive, although the increase
in plasma testosterone levels evoked in male rats by the proximity of female rats is further enhanced by the selective 5-HT₃ antagonist ondansetron, which suggests that 5-HT₃ receptor blockade may lead to an enhanced sexual drive (Amstislavskaya & Popova, 2004).

The generalizability of results from this study to the broad population of depressed patients, like most randomized controlled trials, is limited by the inclusion and exclusion criteria. Patients aged <18 yr or >65 yr were not included, nor were patients with specified psychiatric or medical comorbidities, or patients at risk of suicidal behaviour, nor those with treatment-resistant depression or with mild to moderate depression. The titration of venlafaxine XR, from 75 mg to 225 mg over 7 d, was according to the manufacturer’s instructions. Only one patient treated with venlafaxine withdrew in the first week of treatment, indicating that there was no bias due to early withdrawals in this treatment arm.

In conclusion, treatment with 5 mg and 10 mg Lu AA21004 for 6 wk in this proof-of-concept study was well tolerated and efficacious in reducing depressive and anxious symptoms in patients with MDD.

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