Pharmacokinetic Evaluation of a 1-Day Treatment Initiation Option for Starting Long-Acting Aripiprazole Lauroxil for Schizophrenia

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Abstract:  
Background: Aripiprazole lauroxil (AL), a long-acting injectable antipsychotic for the treatment of schizophrenia, requires 21 days of oral aripiprazole supplementation upon initiation. We report findings from a phase 1 study investigating a nanocrystalline milled dispersion of AL (ALNCD) as a potential 1-day initiation regimen. The 1-day initiation regimen is designed to enable rapid achievement of plasma aripiprazole concentrations that are comparable with the 21-day oral initiation regimen. Here, a 6-month pharmacokinetic study compared 2 different initiation regimens for starting AL.

Methods: Patients were randomized 1:1:1:1 to receive 1 of 4 treatments consisting of the 1-day (single ALNCD injection + one 30-mg dose of oral aripiprazole on day 1 only) or the 21-day (15-mg daily dose of oral aripiprazole for 21 days) initiation regimen, each combined with a starting AL dose of either 441 mg or 882 mg.

Results: In total, 133/161 patients completed the study. The pharmacokinetic profile of the 1-day initiation regimen was comparable to the 21-day initiation regimen; both achieved aripiprazole concentrations in the therapeutic range within 4 days and remained in a comparable concentration range during treatment initiation. Common adverse events (≥25.0%) were injection-site pain, headache, increased weight, insomnia, dyspepsia, and anxiety. Nine akathisia events occurred (4 events in 4 patients and 5 events in 2 patients in the 1-day and 21-day initiation regimen groups, respectively).

Conclusions: The 1-day initiation regimen resulted in plasma aripiprazole concentrations consistent with the 21-day initiation regimen. Therefore, a single dose of ALNCD with a single 30-mg oral dose of aripiprazole provides an alternative initiation regimen for starting AL.

Key Words: aripiprazole lauroxil, aripiprazole lauroxil nanocrystal dispersion, long-acting injectable, pharmacokinetics, schizophrenia

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of aripiprazole concentrations within the therapeutic range in a time frame similar to that of the 21-day oral aripiprazole supplementation and consistent with the 21-day oral regimen used in the pivotal ef-
ciency study of AL. The present study tested the PK projection
that a single dose of ALNCND along with a single 30-mg dose of
oral aripiprazole, could be used as part of a 1-day initiation reg-
imen that has a PK and safety profile comparable to that of the
21-day oral aripiprazole supplementation with proven efficacy.

MATERIALS AND METHODS

Study Design and Treatment Regimens

The study was conducted in accordance with the Declaration
of Helsinki and with Good Clinical Practice Guidelines agreed
by the International Conference on Harmonization in 1997. The
study protocols, amendments, and informed consent forms were
approved by an independent ethics committee/institutional review
board for each site. All patients provided written informed consent
before entering the study. The institutional review board and study
sites are listed in Supplemental Digital Content 1, http://links.lww.
com/JCP/A514.

This was a 6-month, double-blind, placebo-controlled, phase
1 study to assess the PK, safety, and tolerability of 2 initiation reg-
imen for starting treatment with AL in patients with schizophre-
nia (Fig. 1). The 1-day initiation regimen (comprising a single IM
662-mg ALNCND dose, a single 30-mg dose of oral aripiprazole,
and either AL 441 or 882 mg) was compared with the 21-day initia-
tion regimen (21 days of 15-mg oral aripiprazole with either AL
441 or 882 mg). A total of 160 patients were randomized 1:1:1:1 to 1 of 4
treatment groups. In all study groups, the order of administration
on day 1 was as follows: oral aripiprazole; IM injection of ALNCND
or placebo (<15 minutes after oral aripiprazole); IM injection of
AL (<30 minutes after IM injection of ALNCND or placebo).
Nanocrystalline milled dispersion of AL or placebo was admin-
istered as IM injections in the gluteal muscle. For AL doses, a
single 441-mg dose was given in the deltoid muscle or a single
882-mg dose in the gluteal muscle contralateral to the ALNCND
(oral placebo).

Patients were admitted as inpatients 1 day before their first
scheduled dose and were maintained as inpatients for the first
15 days. After discharge, patients returned for outpatient follow-
up assessments. Pharmacokinetic samples were taken daily on
days 1 to 15, every other day from days 17 to 25, on days 28
and 31, once weekly from days 35 to 85, and on days 113 and
141. On days 1 and 21, samples were collected at multiple time
points (as detailed in Pharmacokinetics).

Study Population

Main Inclusion Criteria

Eligible patients were adults aged 18 to 65 years with a di-
agnosis of chronic schizophrenia or schizoaffective disorder
based on the Diagnostic and Statistical Manual of Mental Disor-
ders, Fifth Edition, and a documented history of tolerability to
aripiprazole or a demonstrated tolerability to test doses during
screening. In addition, patients were required to have clinically
stable schizophrenia, defined as having no hospitalizations for
acute psychiatric exacerbations within 3 months before screen-
ing and a Clinical Global Impressions-Severity (CGI-S) score
of 3 or lower (mild) at screening and study initiation. Patients
were on a stable oral antipsychotic medication regimen (exclud-
ing aripiprazole and clozapine) for 2 months or more before
screening without any medication changes between screening and
randomization.

Main Exclusion Criteria

Key exclusion criteria included patients who had received
oral aripiprazole for 28 days or less before randomization or any
other LAI antipsychotic for 3 months or less before admission
and patients who were participating or who had participated in a
clinical trial involving any investigational product for 3 months
or less before admission. Patients who received AL or IM depot
aripiprazole for 6 months or less before inpatient admission were
excluded. A history of primary psychopathology other than
schizophrenia or schizoaffective disorder or a positive test result
for illicit drug use at screening or at admission was not permit-
ted. Patients deemed to be CYP2D6 poor metabolizers were ex-
cluded from this study as dose adjustments are required in this
patient population (see Table S1, Supplemental Digital Content
2, http://links.lww.com/JCP/A515).

Study Assessments

Pharmacokinetics

Blood samples for liquid chromatography–tandem mass
spectrometry were collected for analysis within 1 hour predose
and 1, 2, 3, 4, 5, 6, and 8 hours (±15 minutes) postdose on day
1. On postinitiation days 2 to 21, a single sample was collected

FIGURE 1. Study design. Patients were admitted to an inpatient facility on day –1 and were discharged after assessments on day 15. After
discharge, patients received outpatient follow-up assessments until day 141. *Patients naive to aripiprazole were administered 5-mg test doses
of oral aripiprazole on day –30 and day –29. †Patients in the 1-initiation groups were treated as follows: ALNCND intramuscular (gluteal) plus
30-mg oral aripiprazole plus either intramuscular AL 441 mg (deltoid) or AL 882 mg (contralateral gluteal) on day 1 followed by 20 days of oral
placebo. ‡Patients in the 21-initiation groups were treated as follows: placebo injection (gluteal) plus 15-mg oral aripiprazole plus either
intramuscular AL 441 mg (deltoid) or AL 882 mg (contralateral gluteal) on day 1 followed by 20 days of 15-mg oral aripiprazole.
before oral aripiprazole (or oral placebo) administration. As on day 1, after collection of the predose sample on day 21, additional samples were collected 1, 2, 3, 4, 5, 6, and 8 hours (±15 minutes) postdose. For days 23 to 85, a single sample was collected within ±2 hours of the day 1 oral dosing time, or as close to that time frame as possible. Single PK samples were collected on days 113 and 141. Concentrations of aripiprazole and dehydroaripiprazole were quantified in these plasma samples.

Plasma samples were prepared by a protein-precipitation extraction procedure and analyzed using high-performance liquid chromatography coupled to a tandem mass spectrometry detector (LC/MS/MS). The concentrations of all analytes were calculated using 1/x² linear regression with a lower limit of quantification of 1.00 ng/mL. Chromatographic separations were performed on a reversed phase column (UPLC SB-C8 1.8 µm, 2.1 x 100 m; from Agilent Technologies, Santa Clara, Calif.). The mobile phases were pH unadjusted 0.1% formic acid in 10 mM ammonium acetate (A) and acetonitrile (B). A gradient elution was used, starting at 30% B and ramping to 37% in 0.2 minutes, then increasing to 95% B in 2.9 minutes with a flow rate of 0.4 mL/min, holding at 95% B for 3.1 minutes with a flow rate of 0.8 mL/min, and then lowering back to 30% B in 0.2 minutes with a flow rate of 0.4 mL/min. The total run time was 7.0 minutes. The protonated analytes were quantified by selected reaction monitoring in the positive ionization mode by triple quadrupole mass spectrometer. The method was developed to detect AL, N-hydroxymethyl aripiprazole, aripiprazole, and dehydroaripiprazole, as well as their respective deuterated standards, for analyte transitions of aripiprazole, aripiprazole, and dehydroaripiprazole, as well as their respective deuterated standards, for analyte transitions of aripiprazole, aripiprazole, and dehydroaripiprazole, and dehydroaripiprazole and AUC₀₋₂₈ were summarized descriptively by treatment group.

A post hoc evaluation was conducted to compare aripiprazole concentration results from the present study with observed concentrations from the 12-week phase 3 efficacy study (that used the 21-day oral regimen).

Safety and tolerability parameters were estimated in the safety population. Adverse events that were new or that worsened from the time of administration of the first dose of study drug (ALNC₁₀₀) and a single 30-2 mg oral aripiprazole dose or placebo injection and 21 days of oral aripiprazole 15 mg plus either AL 441 or 882 mg) were summarized using descriptive statistics.

**RESULTS**

**Patient Disposition and Baseline Characteristics**

In total, 161 patients were enrolled, received one of the initiation regimens, and were included in the PK and safety populations (Fig. 1). Patients were randomized to receive a 1-day initiation regimen (n = 80) or a 21-day initiation regimen (n = 81), along with an AL starting dose of either 441 or 882 mg. Thirty-nine patients were enrolled in the AL 441 mg/1-day initiation group, and 41 patients were enrolled in the AL 882 mg/1-day initiation group. Of those enrolled in the 21-day initiation regimen groups, 40 patients were assigned to the AL 441 mg/21-day initiation group and 41 patients were assigned to the AL 882 mg/21-day initiation group.

A total of 133 patients (82.6%) completed the study. Among the 28 patients (17.4%) who did not complete the study, reasons for study withdrawal included lost to follow-up and withdrawal by the patient (each n = 10; 6.2%), AE (n = 5; 3.1%), protocol deviation (n = 2; 1.2%), and noncompliance with medication (n = 1; 0.6%). Patient demographics are summarized in Table 1. Mean age and body mass index of the patients were 44 years and 29.5 kg/m², respectively. All patients were either CYP2D6 extensive or intermediate metabolizers with the exception of 4 patients, whose status was considered to be inconclusive. All patients with an inconclusive result had at least 1 functional allele and were not poor metabolizers. Overall, the treatment groups were well balanced for demographic and baseline characteristics.

**Pharmacokinetic Results**

Results from the 1-day initiation regimen groups showed mean plasma aripiprazole concentrations and exposures within the first month that were comparable to those of the 21-day initiation regimen groups (Fig. 2). In the first 24 hours after initiation, higher aripiprazole concentrations were observed with the 1-day initiation regimen groups compared with the 21-day initiation regimen groups due to the higher dose of aripiprazole administered on day 1 with the 1-day initiation regimen (30 mg vs 15 mg). Plasma concentrations of postinitiation on day 4 were of particular interest because the 1-day initiation regimen was designed to replicate the 21-day initiation regimen in achieving aripiprazole concentrations in the therapeutic range within 4 days after the first AL dose. As shown in Figure 2, the 1-day regimen, like the 21-day initiation regimen, results in achievement of aripiprazole concentrations that are in the therapeutic range within 4 days.

As seen in Figure 2, mean concentrations appear visually lower in the 1-day initiation regimen groups than in the 21-day initiation regimen groups from approximately day 4 to day 14; error bars around the plasma concentration means show complete overlap in the range of concentrations across the treatment groups. As expected for the 21-day oral initiation regimen group, aripiprazole concentrations declined after day 21 after discontinuation of the active oral medication. In contrast, for the 1-day initiation regimen
groups, plasma aripiprazole concentrations did not show any meaningful changes until after postinitiation on day 30, when mean aripiprazole concentrations began to decline (Fig. 3), indicating that the 1-day initiation regimen provides continuous coverage over a longer period than the 21-day initiation regimen.

Values of AUC_{0–28} were comparable across the 4 treatment groups (Fig. 4). Comparison of the range of values across groups indicated similar exposure within the first month of treatment regardless of the initiation regimen used.

Post hoc comparison of aripiprazole concentrations resulting from the present study with those observed in the 12-week phase 3 efficacy study showed consistent and reproducible results across studies with the 21-day oral aripiprazole initiation regimen (see Fig. S1, Supplemental Digital Content 3, http://links.lww.com/JCP/A516). The 1-day initiation regimen from the present study resulted in aripiprazole concentrations within the concentration range observed with the 21-day oral aripiprazole supplementation used in the phase 3 efficacy study (see Fig. S2, Supplemental Digital Content 4, http://links.lww.com/JCP/A517).

The plasma dehydroaripiprazole concentration–time profile followed that of aripiprazole in each treatment group through day 21 (the last day of oral administration of aripiprazole or placebo). Thereafter, plasma dehydroaripiprazole concentrations continued to persist in the systemic circulation in each 1-day initiation regimen group, whereas concentrations decreased more rapidly in each 21-day initiation regimen group (see Fig. S3, Supplemental Digital Content 5, http://links.lww.com/JCP/A518). Mean ± SD AUC_{0–28} values were comparable in each treatment group: ALNCD initiation regimen + AL 441 mg 441 mg/1-day initiation regimen, 1222.4 (455.7) day × ng/mL; ALNCD initiation regimen + AL 882 mg 21-day initiation regimen + AL 441 mg (deltoid) or AL 882 mg (contralateral gluteal) on day 1 followed by 20 days of oral placebo.

### TABLE 1. Patient Baseline Characteristics

| Treatment Group | ALNCD Initiation Regimen + AL 441 mg | ALNCD Initiation Regimen + AL 882 mg | 21-day Oral Initiation Regimen + AL 441 mg | 21-day Oral Initiation Regimen + AL 882 mg |
|-----------------|-------------------------------------|-------------------------------------|------------------------------------------|------------------------------------------|
| n               | 161                                 | 39                                  | 41                                       | 40                                       |
| Mean age (SD), y| 44.0 (10.6)                         | 44.4 (10.0)                        | 42.3 (12.4)                              | 44.2 (9.7)                              |
| Male, n (%)     | 118 (73.3)                          | 30 (76.9)                           | 29 (70.7)                                | 27 (67.5)                               |
| Race, n (%)     |                                    |                                    |                                          |                                          |
| Black or African American | 125 (77.6) | 31 (79.5) | 33 (80.5) | 26 (65.0) |
| White           | 35 (21.7)                           | 8 (20.5)                            | 8 (19.5)                                 | 14 (35.0)                               |
| Asian           | 1 (0.6)                             | 0 (0.0)                             | 0 (0.0)                                  | 0 (0.0)                                 |
| Mean BMI (SD), kg/m² | 29.5 (5.4) | 28.1 (5.5) | 29.8 (4.7) | 30.3 (5.3) |
| CYP2D6 metabolizer status, n (%) | Extensive | 106 (65.8) | 26 (66.7) | 29 (70.7) |
| Intermediate    | 51 (31.7)                           | 13 (33.3)                           | 10 (24.4)                                | 13 (32.5)                               |
| Inconclusive    | 4 (2.5)                             | 0 (0.0)                             | 2 (4.9)                                  | 2 (5.0)                                 |

All values are mean values, unless otherwise indicated.

*Patients in the 1-day initiation groups were treated as follows: ALNCD intramuscular (gluteal) plus 30-mg oral aripiprazole plus either intramuscular AL 441 mg (deltoid) or AL 882 mg (contralateral gluteal) on day 1 followed by 20 days of oral placebo.†Patients in the 21-day initiation groups were treated as follows: placebo injection (gluteal) plus 15-mg oral aripiprazole plus either intramuscular AL 441 mg (deltoid) or AL 882 mg (contralateral gluteal) on day 1 followed by 20 days of 15-mg oral aripiprazole.
Aripiprazole Lauroxil 1-Day Initiation: PK and therefore was evaluated in greater detail. All ISRs could not be estimated for aripiprazole, by starting AL dose/initiation regimen. The boxes represent the 25th and 75th percentiles of in con-

The most common description of ISRs was injection-site pain, which was reported in 12 (15.0%) of 80 patients who received ALNCD injection compared with 4 (4.9%) of 81 patients who received placebo IM injection. ISRs Associated With AL Injection

Eighteen (22.8%) of 79 patients who received AL 441 mg in the deltoid muscle and 15 (18.3%) of 82 patients who received AL 882 mg in the gluteal muscle experienced ISRs (Table 2). The most common description of ISRs was injection-site pain, which was reported in 16 (20.3%) of 79 patients who received AL 441 mg and 15 (18.3%) of 82 patients who received AL 882 mg. Akathisia

Akathisia was an AE of interest because it is commonly reported in association with oral aripiprazole used for the treatment of patients with schizophrenia. The overall incidence of akathisia in all groups was low, with a total AE rate of 6 (3.7%) of 161 patients. Among patients treated with the 1-day initiation regimen, akathisia was reported in 4 (5.0%) of 80 patients. Two of these patients reported mild akathisia during the first week of treatment. One was assessed as having mild akathisia probably not related to the study drug and the other was assessed as having mild akathisia definitely related to the study drug. The other 2 patients experienced akathisia in the third week of treatment, one of which was rated as mild and the other as moderate in severity, assessed as probably related and definitely related to treatment, respectively. Among patients treated with the 21-day oral regimen, mild akathisia was reported in 2 (2.5%) of 81 patients. One experienced the first akathisia event in the second week and the other in the third week of treatment, assessed as possibly related and probably related to treatment, respectively.

DISCUSSION

A 1-day initiation regimen was evaluated as an alternative to the current 21-day initiation regimen. In this study, ALNCD in conjunction with 30-mg oral aripiprazole and either a 441-mg or an

FIGURE 4. Box plot of AUC0–28 for aripiprazole, by starting AL dose/initiation regimen. The boxes represent the 25th and 75th percentiles of aripiprazole concentration, the line within each box marks the median, the asterisk indicates the mean, and the whiskers indicate the 10th and 90th percentiles. The squares represent individual observations beyond the 10th and 90th percentiles. *AUC0–28 values could not be estimated for all patients because some patients discontinued the study before day 28.

AL 441 mg/21-day initiation regimen, 1435.5 (654.8) day × ng/mL; AL 882 mg/21-day initiation regimen, 1315.9 (439.6) day × ng/mL.

Safety Results

Adverse Events

Throughout the study period, small and similar mean changes (≤0.1) from baseline (score of 3.0, mild) in CGI-S score were seen in each initiation regimen group at all time points, indicating no change in disease severity. All patients had a score of 0 (no suicidal behavior or ideation) for Columbia-Suicide Severity Rating Scale throughout the study.

In the AL 441 mg/1-day initiation and the AL 882 mg/1-day initiation groups, 26 patients (66.7%) and 28 patients (68.3%) experienced AEs, respectively. In addition, 24 patients (60.0%) and 28 patients (68.3%) in the AL 441 mg and the AL 882 mg/21-day initiation groups experienced AEs, respectively (see Table S2, Supplemental Digital Content 6, http://links.lww.com/JCP/A519). Most AEs were mild or moderate in intensity. Serious AEs were reported in 6 patients: 3 each in the 1-day initiation regimen (road traffic accident, status epilepticus, psychotic disorder, and tricyclic disorder) and 2 in the 21-day initiation regimen (road traffic accident and status epilepticus were assessed as possibly related to treatment. A total of 5 patients discontinued the study because of AEs: 3 in the 1-day initiation regimen (road traffic accident, upper gastrointestinal hemorrhage, cellulitis, road traffic accident, and accidental overdose). Of these, schizoaffective disorder) and the 21-day initiation regimen (sustained in a road traffic accident (considered not related to study treatment). A total of 5 patients discontinued the study because aripiprazole concentration, the line within each box marks the median, the asterisk indicates the mean, and the whiskers indicate the 10th and 90th percentiles. The squares represent individual observations beyond the 10th and 90th percentiles. The squares represent individual observations beyond the 10th and 90th percentiles. The squares represent individual observations beyond the 10th and 90th percentiles. The squares represent individual observations beyond the 10th and 90th percentiles.

Adverse Events of Interest

Injection-site reaction is a common AE associated with LAIs

and therefore was evaluated in greater detail. All ISRs associated with the ALNCD placebo IM, AL 441 mg, and AL 882 mg injections were mild to moderate in severity. ISRs Associated With ALNCD or Placebo IM Injection

Fourteen (17.5%) of 80 patients who received ALNCD injection experienced ISRs compared with 5 (6.2%) of 81 patients who received placebo injection (Table 2). The most common description of ISRs was injection-site pain, which was reported in 12 (15.0%) of 80 patients who received ALNCD injection compared with 4 (4.9%) of 81 patients who received placebo IM injection.

Akathisia

Akathisia was an AE of interest because it is commonly reported in association with oral aripiprazole used for the treatment of patients with schizophrenia. The overall incidence of akathisia in all groups was low, with a total AE rate of 6 (3.7%) of 161 patients. Among patients treated with the 1-day initiation regimen, akathisia was reported in 4 (5.0%) of 80 patients. Two of these patients reported mild akathisia during the first week of treatment. One was assessed as having mild akathisia probably not related to the study drug and the other was assessed as having mild akathisia definitely related to the study drug. The other 2 patients experienced akathisia in the third week of treatment, one of which was rated as mild and the other as moderate in severity, assessed as probably related and definitely related to treatment, respectively. Among patients treated with the 21-day oral regimen, mild akathisia was reported in 2 (2.5%) of 81 patients. One experienced the first akathisia event in the second week and the other in the third week of treatment, assessed as possibly related and probably related to treatment, respectively.
882-mg dose of AL resulted in aripiprazole concentrations within the therapeutic range by the fourth day and a range of aripiprazole concentrations that overlapped with those produced by the 21-day period of oral aripiprazole. Results also showed that the 1-day initiation regimen provided continuous exposure within the first month of treatment initiation that was comparable with the 21-day initiation regimen. In addition, total aripiprazole exposure during the first 28 days after treatment initiation was comparable between the 2 initiation regimens. Dehydroaripiprazole concentrations followed a pattern generally similar to that seen for aripiprazole, and total exposure was similar between treatment groups. The 1-day initiation regimen was shown to be suitable as an alternative option to the current 21 days of concomitant oral aripiprazole for starting AL.

Comparison of aripiprazole concentrations for the 21-day initiation regimen groups in the present study with those obtained from the 2 active AL arms in the pivotal 12-week phase 3 efficacy study that used the same dose of oral aripiprazole (21 days of 15 mg/d) (NCT01469039) showed that the aripiprazole concentrations achieved were consistent and reproducible. In addition, aripiprazole concentrations resulting from the 1-day initiation regimen in the present study were within the concentration range observed with oral initiation in the phase 3 study. A to-be-marketed dose strength of 675 mg is proposed to distinguish ALNCD from AL in the clinical setting. The dose of 675 mg is within the range of acceptable variance for the 662-mg dose.

Overall, treatment with ALNCD in combination with a single dose of 30-mg oral aripiprazole and either AL 441 mg or AL 882 mg was well tolerated in patients with schizophrenia. The safety profile of the 1-day initiation regimen was generally consistent with the known safety profile of aripiprazole and AL, with most AEs being mild to moderate in intensity and a low rate of discontinuations occurring due to drug-related AEs. A low incidence of akathisia was reported, and analysis of the time to akathisia occurrence and its severity did not show any differences between the 1-day and the 21-day groups.

One patient experienced a serious AE of status epilepticus. This patient experienced peak aripiprazole concentrations on day 7 and by day 17, and when the event occurred, aripiprazole concentrations were declining. The postseizure magnetic resonance imaging findings were consistent with preexisting seizure diathesis and focal structural lesion was excluded. Although these details were revealed through the follow-up information, the causality remained conservatively the same as possibly related. The patient was also prescribed multiple other psychotropic medications. Although rated as “possibly related” to the AL and ALNCD intervention, there are multiple confounders in this case for a plausible relationship to the study drug.

The results demonstrate that a 1-day initiation regimen was comparable to the current 21-day initiation regimen and may offer an alternative for patients starting AL therapy. In addition, a 1-day initiation regimen may be beneficial for patients in whom poor adherence is anticipated, and may offer another option for patients who may prefer a regimen with lower pill burden or who are interested in reducing the duration of the oral initiation regimen from 21 days to 1 day.

In conclusion, a key step in the management of schizophrenia is successful completion of the initiation phase for the LAI antipsychotic treatment. In the present study, we report the development of a 1-day initiation regimen for AL that uses a single injection of ALNCD and a single oral dose of aripiprazole. The proposed 1-day initiation regimen could provide clinicians and patients with the option of an LAI initiation that can be completed in a single day as an alternative to 21 consecutive days of oral aripiprazole in conjunction with the first dose of AL.

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AUTHOR DISCLOSURE INFORMATION

Dr Hard is a former employee of Alkermes and is presently an employee of Nuvotena Pharma Sciences. Drs Wehr, Weiden, and von Moltke are employees of Alkermes. Dr Walling has received grants from Alkermes, Janssen, Otsuka, Forum, Lundbeck, Sunovion, Acadia, Allergan, IntraCellular, Noven, Merck, AbbVie, and Roche.

Lisa von Moltke has declared a relationship with a member of the Editorial Board of the Journal of Clinical Psychopharmacology; which may be considered a possible conflict of interest. An alternative independent review mechanism has been utilized

### Table 2. ISRs by Treatment Group Associated With ALNCD or Placebo Injection and Those Associated With AL (441 mg or 882 mg)

| Injection, n (%) | ALNCD | Placebo Injection | AL 441 mg | AL 882 mg |
|----------------|-------|-------------------|-----------|-----------|
| n = 80         | n = 81| n = 79            | n = 82    |
|Patients with at least 1 ISR | 14 (17.5) | 5 (6.2) | 18 (22.8) | 15 (18.3) |
|Injection-site pain | 12 (15.0) | 4 (4.9) | 16 (20.3) | 15 (18.3) |
|Injection-site induration | 3 (3.8) | 0 | 4 (5.1) | 2 (2.4) |
|Injection-site swelling | 0 | 0 | 2 (2.5) | 1 (1.2) |
|Injection-site discomfort | 0 | 0 | 1 (1.3) | 0 |
|Injection-site erythema | 1 (1.3) | 1 (1.2) | 1 (1.3) | 0 |

The injection site and surrounding area were evaluated with each injection (separately for the ALNCD or placebo injection site and the AL site). Any observed ISRs were monitored until resolution.

*Patients received a single ALNCD injection (gluteal) on day 1 as part of the 1-day initiation regimen.
†Patients received a placebo injection (gluteal) on day 1 as part of the 21-day initiation regimen.
‡Patients received an AL 441 mg injection (deltoid) on day 1 as part of the 1-day or 21-day initiation regimen.
§Patients received an AL 882 mg injection (contralateral gluteal) on day 1 as part of the 1-day or 21-day initiation regimen.

*A single patient could have more than 1 ISR.
according to the Journal of Clinical Psychopharmacology procedures for dealing with any potential appearance of bias.

The study was conducted in accordance with the Declaration of Helsinki and with Good Clinical Practice Guidelines agreed by the International Conference on Harmonization in 1997. Study protocols, amendments, and informed consent forms were approved by the independent ethics committee/institutional review board for each site.

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

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