Lorecivivint, a Novel Intraarticular CDC-like Kinase 2 and Dual-Specificity Tyrosine Phosphorylation-Regulated Kinase 1A Inhibitor and Wnt Pathway Modulator for the Treatment of Knee Osteoarthritis: A Phase II Randomized Trial

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Objective. To assess the safety and efficacy of a novel Wnt pathway modulator, lorecivivint (SM04690), for treating pain and inhibiting structural progression in moderately to severely symptomatic knee osteoarthritis (OA).

Methods. Subjects in this 52-week, phase IIa, multicenter, randomized, double-blind, placebo-controlled, dose-ranging trial received a single 2-ml intraarticular injection of lorecivivint (dose of 0.03 mg, 0.07 mg, or 0.23 mg) or placebo. Efficacy was assessed based on change from baseline on the Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC) score subscales for pain and function (scale 0–100 for each) and change from baseline in the radiographic medial joint space width (JSW). Baseline-adjusted analysis of covariance with multiple imputation was performed separately to evaluate efficacy. This proof-of-concept study evaluated the intent-to-treat population as well as a prespecified group of subjects with unilateral symptoms of knee OA (designated UNI) and an additional post hoc subgroup of subjects with unilateral symptoms but without widespread pain (designated UNI WP−).

Results. In this trial, 455 subjects were randomized to a treatment group. The primary end point, significant improvement in the WOMAC pain score compared with placebo at week 13, was not met by any lorecivivint dose group (mean ± SD change from baseline, −23.3 ± 2.2 in the 0.03 mg group, −23.5 ± 2.1 in the 0.07 mg group, −21.3 ± 2.2 in the 0.23 mg group, and −22.1 ± 2.1 in the placebo group; each P > 0.05 versus placebo). All groups (including placebo) demonstrated clinically meaningful (≥20-point) improvements from baseline in the WOMAC pain score. The durability of response was evaluated through week 52. In the prespecified UNI group and post hoc UNI WP− group at week 52, treatment with 0.07 mg lorecivivint significantly improved the WOMAC pain score (between-group difference versus placebo, −8.73, 95% confidence interval [95% CI] −17.44, −0.03 [P = 0.049] and −11.21, 95% CI −20.99, −1.43 [P = 0.025], respectively) and WOMAC function score (between-group difference versus placebo, −10.26, 95% CI −19.82, −0.69 [P = 0.036] and −13.38, 95% CI −24.33, −2.43 [P = 0.017], respectively). Relative to baseline, the mean change in the medial JSW at week 52 was −0.04 mm in the 0.03 mg cohort, −0.09 mm in the 0.07 mg cohort, −0.16 mm in the 0.23 mg cohort, and −0.14 mm in the placebo cohort; no treatment group achieved a significant change in medial JSW compared with placebo at week 52. In both unilateral symptom subgroups, the 0.07 mg lorecivivint dose significantly increased medial JSW compared with placebo at week 52 (medial JSW 0.39 mm, 95% CI 0.06, 0.72 in the UNI group [P = 0.021] and 0.42 mm, 95% CI 0.04, 0.80 in the UNI WP− group [P = 0.032]). Changes observed in the 0.03 mg and 0.23 mg dose groups were not significantly different from those in the placebo group for any of these measures. Lorecivivint appeared safe and well tolerated.

Conclusion. This phase IIa, proof-of-concept trial in patients with symptomatic knee OA did not meet its primary end point. Nevertheless, the study identified a target population in whom to evaluate the potential efficacy of lorecivivint for the treatment of knee OA.
INTRODUCTION

Knee osteoarthritis (OA) is a common, chronic disorder that is characterized by cartilage destruction, subchondral bone thickening, and osteophyte formation, leading to pain, functional limitation, and physical disability (1). The severity of knee OA is assessed by a combination of patient-reported outcome measures that include assessments of pain and function and objective structural measures such as radiologically assessed joint space narrowing (2). Pharmacologic interventions for knee OA management are symptom-alleviating treatments, including oral and topical nonsteroidal antiinflammatory drugs (NSAIDs), nonopioid analgesics, and intraarticular (IA) corticosteroid or hyaluronic acid injections (3,4). However, many of these treatments have limited short-term and long-term efficacy (4–7) and are associated with a high incidence of side effects. There remains a great unmet need for new treatments that provide symptom relief and even more of a need for disease-modifying OA drugs (DMOADs).

The Wnt pathway is integral for tissue homeostasis and regeneration (8,9) and is a key regulator of progenitor cell differentiation in the knee joint (10). Cartilage homeostasis requires a balance of Wnt pathway activity. While necessary for chondrocyte differentiation and function (11), aberrant Wnt pathway activity in OA directs progenitor cell differentiation in the joint toward development of osteoblasts instead of chondrocytes (12). Excessive activation of the Wnt pathway is known to increase OA susceptibility in animals and humans (13–16), whereas excessive inhibition of the Wnt pathway can cause cartilage and bone destruction (17–19). Therefore, a potential Wnt pathway–targeted DMOAD approach would need to maintain signaling within an optimal range.

Lorecivivint (SMO4690) is a small-molecule Wnt pathway modulator currently in development as a potential DMOAD for the treatment of knee OA (20,21). Lorecivivint affects Wnt pathway activity via inhibition of 2 intranuclear targets, CDC-like kinase 2 (CLK2) and dual-specificity tyrosine phosphorylation-regulated kinase 1A (DYRK1A), through which it acts both independently and in combination to improve chondrocyte health and function while inhibiting inflammation (22). Preclinical studies, including repeat dosing in rats and dogs, have found the no-observed-adverse-effect level to be ~400 times the planned dose in humans (data on file). In vitro studies demonstrated that lorecivivint modulated Wnt signaling, reduced release of matrix-degrading enzymes from chondrocytes, demonstrated anabolic activity in chondrocytes, and reduced STAT3 signaling, NF-κB signaling, and inflammatory cytokine production in synoviocytes (20). In a rat model of anterior cruciate ligament transection and partial medial meniscectomy-induced knee OA, a single IA injection of lorecivivint protected chondrocytes from catabolic breakdown (20).

In a previous phase I, randomized, placebo-controlled trial (n = 61), a single IA injection of lorecivivint at a dose of 0.03 mg, 0.07 mg, or 0.23 mg administered into the target knee joint of subjects with moderately to severely symptomatic knee OA appeared safe and well tolerated and showed no evidence of systemic exposure. While all lorecivivint and placebo groups demonstrated improvements from baseline in the Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC) pain and function subscale scores at week 24, the 0.07 mg lorecivivint treatment group demonstrated more favorable reductions in both WOMAC indices as compared with placebo. Treatment with 0.07 mg lorecivivint also resulted in increased radiographic joint space width (JSW) beyond a minimum detectable difference of 0.13 mm (23), thus suggesting that lorecivivint may be a potential DMOAD for use in treating knee OA. Therefore, the objective of this phase IIa trial was to evaluate the safety and efficacy of lorecivivint among subjects with moderately to severely symptomatic knee OA.

SUBJECTS AND METHODS

Study design. This was a 52-week, phase IIa, multicenter, randomized, double-blind, placebo-controlled, dose-ranging trial of 3 different dose concentrations of lorecivivint injected into the target (most painful) knee joint of subjects with moderately to severely symptomatic knee OA. This study was conducted at 36 clinical sites in the United States between September 2015 and April 2017. Subjects participated in a screening period of up to 21 days and were periodically observed during a 52-week follow-up period. Visits were

witness on behalf of Williams & Connolly, LLP and Nix Patterson, LLP. Dr. Jones has received consulting fees from Samumed (less than $10,000) and receives royalties for the Journal of Bone and Joint Surgery. Dr. Bergfeld has received consulting fees from Samumed (less than $10,000). Dr. Hochberg has received consulting fees, speaking fees, and/or honoraria from Bone Therapeutics, Bristol Myers Squibb, EMD Serono, IBSA, Novartis Pharma AG, Regeneron, Samumed, Symic Bio, Theralogix, TissueGene, Vertex Pharmaceuticals, Vizu Health Sciences, Zynerva, Covance, Galapagos, ICON, and IQVIA (less than $10,000 each) and from Eli Lilly and Pfizer (more than $10,000), owns stock or stock options in Biori Biotech and Theralogix, and receives royalties from Wolters Kluwer for UpToDate and from Elsevier for Rheumatology 7th Edition.

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Submitted for publication August 29, 2019; accepted in revised form May 13, 2020.
scheduled at screening, treatment visit day 1, and at follow-up weeks 4, 13, 26, 39, and 52.

On day 1, subjects were randomized to receive a single 2-mL IA injection of lorcicrevint at a dose of 0.03 mg, 0.07 mg, or 0.23 mg or phosphate buffered saline as placebo. These 3 doses of lorcicrevint corresponded to the lower, middle, and upper therapeutic ranges that were established previously in preclinical studies (data on file). Randomization was accomplished using Medidata Balance (Medidata Solutions, Inc.) such that eligible subjects were randomized at a ratio of 1:1:1:1 using a permuted block design, with a block size of 8 and stratification by site. An unblinded pharmacist at each site mixed the working dose from a common stock solution bottle, and an unblinded injector performed the injection. Ultrasound guidance and joint aspiration (up to 0.5 cc) were allowed if these were considered part of the site’s standard IA injection protocol for joint placement. All unblinded site personnel were instructed to minimize any contact with study subjects and were not allowed to perform any study assessments. All study investigators and subjects were blinded with regard to group assignment, and subject blinding was maintained by not allowing any subject to witness the injection.

This study was conducted in accordance with the ethics principles of the Declaration of Helsinki, the International Conference on Harmonisation Guidelines for Good Clinical Practice, and applicable regulations. The study protocol was approved by each clinic site’s independent ethics committee or institutional review board. All subjects provided written informed consent prior to participation in any study-related procedures.

Subjects. Eligible subjects were ages 40–80 years with an established diagnosis of primary knee OA (established within ≥6 months prior to study start) and fulfilled American College of Rheumatology clinical and radiographic classification criteria (24). Enrolled subjects were required to have Kellgren/Lawrence (K/L) radiographic disease stage 2 or stage 3 OA in their target knee (defined at screening as the knee with greater pain based on the subject’s evaluation and the investigator’s clinical judgment) (25). Subjects were required to have a pain visual analog scale (VAS) score of 30–80 mm (on 100-mm VAS) (26) and a WOMAC total score of 72–192 (of 240) (27) at the target knee at screening. There were no limitations on contralateral knee pain. Subjects were considered eligible if they were in good general health and ambulatory; assistive devices (e.g., canes) were allowed if needed <50% of the time, whereas any use of a walker was excluded.

Key exclusion criteria included male subjects with female partners of childbearing potential who refused to use an effective contraceptive method and women who were of childbearing potential. Further exclusions included body mass index (BMI) >40 kg/m², history of partial or complete joint replacement in the target knee, previous exposure to lorcicrevint, a major surgery (e.g., interventional arthroscopy) in the target knee within 52 weeks prior to study medication injection, and any planned or elective surgery anywhere in the body during the study period. Additional exclusion criteria included having comorbid conditions that could affect pain assessment of the target knee or a history of malignancy (except for in situ cancer or basal or squamous cell skin cancer) <5 years prior to injection. Subjects could not receive any IA injections of glucocorticoids, hyaluronic acid, or other therapeutic agents into either knee during the study or within 2 months, 6 months, or 1 month prior to randomization, respectively. Electrotherapy or acupuncture for knee OA, chiropractic knee adjustments, or planned or elective surgery (e.g., arthroscopy) were also prohibited. Subjects could not take opioid analgesics or oral glucocorticoids during the study, although a stable background regimen of NSAIDs and acetaminophen was allowed provided that they were not taken within 24 hours prior to study visits.

Data collection. Subject characteristics, medical history, weight, and height were collected at screening. Unilateral or bilateral symptomatic knee OA status was designated by investigators at baseline based on the findings from history and physical examination. To assess comorbidity-related pain and symptoms, subjects completed the fibromyalgia diagnostic questionnaire, Widespread Pain Index (WPI) (total score range 0–19), and Symptom Severity Scale (total score range 0–12) at screening (28).

Efficacy assessments. Efficacy assessments administered at all study visits included the WOMAC questionnaire (version NRS 3.1) to assess pain (subscale range 0–100 [no pain–extreme pain]) and function (subscale range 0–100 [no difficulty with daily activities–extreme difficulty]) and the patient global assessment of disease activity (PtGA) (VAS score range 0–100 mm [“doing very well”–“doing very poorly”]). Fixed-flexion, posterior-anterior radiographs of the tibiofemoral compartments were obtained using a QuAP positioner at screening, week 26, and week 52. Quality control assessments of the radiographs were conducted at a central imaging laboratory (Medical Metrics Labs) in blinded manner (blinded with regard to treatment assignment), and medial JSW was measured using a landmark-based, fixed-location method.

The primary efficacy end point was the change from baseline in WOMAC pain score in the target knee at week 13 in the 0.07 mg lorcicrevint group compared with the placebo group. Key secondary end points included 1) change from baseline in WOMAC pain score in the target knee at week 26, 2) change from baseline in WOMAC function score in the target knee at weeks 13 and 26, 3) change from baseline in the PtGA score at weeks 13 and 26, and 4) change from baseline in medial JSW in the target knee at week 26. Exploratory end points included 1) change from baseline in WOMAC pain and function scores in the target knee at weeks 4, 39, and 52, 2) change from baseline in the PtGA score at
weeks 4, 39, and 52, and 3) change from baseline in medial JSW in the target knee at week 52.

**Safety.** Safety was assessed by evaluating the incidence, severity, and seriousness of treatment-emergent adverse events (TEAEs) and clinically significant changes in clinical laboratory measures and vital signs; no formal statistical analyses were planned for safety outcomes. Safety measures were summarized for all treatment groups as treated, not as randomized.

**Sample size.** A sample size of ~445 subjects was planned for this trial based on standard statistical practice to establish an acceptable level of precision with respect to treatment effect estimation (29); no formal calculation was used a priori to determine sample size. However, based on data from phase Iib trials and historical data, a Monte Carlo simulation was conducted to estimate the possible power of using the WOMAC pain score (estimated power of 95.8%) and WOMAC function score (estimated power of 78.5%) to estimate treatment effect given this sample size.

**Statistical analysis.** Statistical analyses were conducted using SAS version 9.4 (SAS Institute). Baseline characteristics of the subjects in each treatment group are presented as the mean ± SD for continuous variables and as the frequency (proportion of patients) for categorical variables. Efficacy outcome measures were evaluated using analysis of covariance models adjusted for baseline values under the intent-to-treat (ITT) analysis set (i.e., comprising all subjects as randomized). Multiple imputation under the missing-at-random assumption was performed for efficacy outcomes with missing values. The least-squares estimate of the difference in change in the outcome from baseline at each time point between each lorecivivint dose group and placebo, adjusted for baseline value, is reported along with the 95% confidence intervals (95% CIs). The familywise error rate for the efficacy analyses was controlled in the strong sense (i.e., using the Bonferroni method) to ensure that the test was controlled when any of the hypothesized differences were true. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mg lorecivivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, or placebo groups, respectively.

**RESULTS**

**Subject disposition and baseline characteristics.** Overall, 1,033 subjects were screened and 455 (44.0%) were randomized; 3 subjects were removed from the study prior to administration of a study drug injection (Figure 1). Cohorts of 112 subjects, 117 subjects, 109 subjects, and 114 subjects were randomized to receive 0.03 mg lorecivivint, 0.07 mglorexivint, 0.23 mglorexivint, or placebo, respectively. For subjects in the UNI group (n = 164), cohort sizes were 45 subjects, 35 subjects, 45 subjects, and 39 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, 0.23 mglorexivint, and placebo groups, respectively. For subjects in the UNI WP− group (n = 128), cohort sizes were 34 subjects, 33 subjects, and 32 subjects in the 0.03 mg lorecivivint, 0.07 mglorexivint, 0.23 mglorexivint, and placebo groups, respectively. Among the subjects who completed the study, 103 (92%) were in the 0.03 mg cohort, 107 (91.5%) were in the 0.07 mg cohort, 95 (86.4%) were in the 0.23 mg cohort, and 97 (83.6%) were in the placebo cohort. At enrollment, the mean ± SD age of the subjects was 60.3 ± 8.7 years and the mean ± SD BMI was 29.9 ± 4.6 kg/m². Overall, 268 (58.9%) of the enrolled subjects were women, 392 (86.2%) were white, 292 (64.2%) had a K/L radiographic OA severity grade of 3 in the target knee, and 164 (36.0%) were classified as having unilateral symptomatic knee OA but without widespread pain (designated the UNI WP− subgroup), defined as having a WPI score of ≤4 and Symptom Severity Scale question 2 score of ≤2 (disregarding question 3). A post hoc concordance analysis was conducted to estimate the ability of one outcome to predict another outcome. It employed within-group logistic regression to estimate the likelihood of baseline-adjusted changes in medial JSW being associated with positive clinical responses (i.e., achieving both WOMAC pain and WOMAC function score improvements of ≥50% [relative change] and ≥20 points of 100). The area under the curve (AUC) of receiver operator characteristic curves represented the concordance between change in medial JSW and clinical response. Concordance was defined as "acceptable" when the AUC was >0.7 and "excellent" when the AUC was >0.8 (30); an AUC of 0.5 represents concordance that is no better than statistical chance.
Clinical outcomes. All subjects. The differences in change from baseline in WOMAC pain scores between the lorecivint dose groups and the placebo group were not statistically significant at week 13 (mean ± SD change from baseline, −23.3 ± 2.2 in the 0.03 mg group, −23.5 ± 2.1 in the 0.07 mg group, −21.3 ± 2.2 in the 0.23 mg group, and −22.1 ± 2.1 in the placebo group; each \( P > 0.05 \) versus placebo); thus, the primary end point was not met, and all analyses were considered.

Table 1. Demographic and clinical characteristics of the eligible subjects at baseline, by treatment group*

| Characteristic                              | Lorecivint | Lorecivint | Lorecivint | Placebo  |
|---------------------------------------------|------------|------------|------------|----------|
|                                             | 0.03 mg    | 0.07 mg    | 0.23 mg    | Placebo  |
|                                             | (n = 112)  | (n = 117)  | (n = 109)  | (n = 114) |
| Age, mean ± SD years                        | 59.0 ± 9.0 | 60.0 ± 8.2 | 61.3 ± 8.7 | 60.7 ± 8.9 |
| Body mass index, mean ± SD kg/m²            | 29.77 ± 4.81 | 30.81 ± 4.74 | 29.64 ± 4.45 | 29.17 ± 4.40 |
| Female, no. (%)                             | 68 (60.7)  | 60 (51.3)  | 68 (61.8)  | 72 (62.1) |
| Race/ethnicity, no. (%)                     |            |            |            |          |
| White                                       | 92 (82.1)  | 102 (87.2) | 96 (87.3)  | 102 (87.9) |
| African American                            | 18 (16.1)  | 14 (12.0)  | 12 (10.9)  | 10 (8.6)  |
| Hispanic or Latino                          | 20 (17.9)  | 23 (19.7)  | 17 (15.5)  | 21 (18.1) |
| Other                                       | 2 (1.8)    | 1 (0.9)    | 2 (1.8)    | 4 (3.4)   |
| K/L grade 3 radiographic OA, no. (%)        | 74 (66.1)  | 74 (63.2)  | 70 (63.6)  | 74 (63.8) |
| Unilateral symptomatic knee OA, no. (%)     | 45 (40.2)  | 35 (29.9)  | 45 (40.9)  | 39 (33.6) |
| WPI ≤4 and SS Scale score ≤2, no. (%)       | 73 (65.2)  | 79 (67.5)  | 76 (69.1)  | 75 (64.7) |

* K/L = Kellgren/Lawrence; OA = osteoarthritis; WPI = Widespread Pain Index; SS = Symptom Severity (question 2).
exploratory. However, subjects in all of the lorecivivint dose groups and the placebo group achieved at least a 20-point mean improvement from baseline in the WOMAC pain and function subscale scores at week 13 through week 52 and in the PtGA score at all time points postinjection (Figures 2A and 3A, and Supplementary Figure 1A, available on the *Arthritis & Rheumatology* website at http://onlinelibrary.wiley.com/doi/10.1002/art.41315/abstract). Treatment with 0.07 mg lorecivivint led to numerically larger improvements from baseline in the pain and function scores as compared with treatment with either the 0.03 mg or 0.23 mg dose of lorecivivint. This was apparent starting at 13 weeks postinjection and continued through 52 weeks.

**UNI group.** Among the subjects in the UNI subgroup, those receiving the 0.07 mg dose of lorecivivint demonstrated improvements in baseline-adjusted change in WOMAC pain and function scores compared with placebo at week 13, which continued through week 52 (Figures 2B and 3B). At week 52, the 0.07 mg lorecivivint group had significantly lower scores on the WOMAC pain subscale compared with the placebo group (between-group difference, −8.73, 95%
CI −17.44, −0.03; \( P = 0.049 \)) and significantly lower scores on the WOMAC function subscale compared with the placebo group (between-group difference, −10.26, 95% CI −19.82, −0.69; \( P = 0.036 \)). In this subgroup of patients with unilateral symptomatic knee OA, there were no significant differences in either the WOMAC pain score or WOMAC function score between the 0.03 mg or 0.23 mg lorecivivint dose groups compared with the placebo group (Figures 2B and 3B). The 0.03 mg treatment group showed significant improvement in the PtGA score compared with the placebo group at weeks 13 and 26 (see Supplementary Figure 1B [http://onlinelibrary.wiley.com/doi/10.1002/art.41315/abstract]).

**UNI WP− group.** Among the subjects in the UNI WP− subgroup, those receiving 0.07 mg lorecivivint demonstrated significant improvements in the WOMAC pain and function scores compared with the placebo group. The between-group difference in the WOMAC pain score was −9.11 (95% CI −17.75, −0.47) (\( P = 0.039 \)) at week 26, −11.83 (95% CI −23.23, −0.42) (\( P = 0.042 \)) at week 39, and −11.21 (95% CI −20.99, −1.43) (\( P = 0.025 \)) at week 52. The between-group difference in the

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**Figure 3.** Mean scores on the Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC) function subscale over time (left) and ladder plots (mean and 95% confidence intervals) of baseline-adjusted change from baseline in the WOMAC function scores (right), comparing the lorecivivint (LOR) dose groups and the placebo group over time in the intent-to-treat (ITT) analysis set (A), subjects with unilateral symptomatic knee osteoarthritis (OA) (B), and subjects with unilateral symptomatic knee OA but without widespread pain (WP−) (C).
WOMAC function score was −9.62 (95% CI −18.14, −1.10) ($P = 0.027$) at week 26, −11.57 (95% CI −22.31, −0.82) ($P = 0.035$) at week 39, and −13.38 (95% CI −24.33, −2.43) ($P = 0.017$) at week 52 (Figures 2C and 3C). In this subgroup of patients with unilateral symptomatic knee OA but without widespread pain, there were no significant differences in change in pain or function scores between the 0.03 mg or 0.23 mg lorecivivint dose groups and the placebo group (Figures 2C and 3C). However, the 0.03 mg treatment group showed a significant improvement in the PtGA score compared with the placebo group at week 13, and both the 0.03 mg and 0.07 mg treatment groups showed significant improvements in the PtGA score compared with the placebo group at week 26 in this post hoc analysis (see Supplementary Figure 1C [http://onlinelibrary.wiley.com/doi/10.1002/art.41315/abstract]).

**Radiographic outcomes.** All subjects. Compared with the values at baseline, the mean change in medial JSW was −0.07 mm at week 26 and −0.04 mm at week 52 in the 0.03 mg cohort, −0.11 mm at week 26 and −0.09 mm at week 52 in the 0.07 mg cohort, −0.02 mm at week 26 and −0.16 mm at week 52 in the 0.23 mg cohort, and −0.20 mm...
at week 26 and −0.14 mm at week 52 in the placebo cohort (Figure 4A). At week 26, the mean change in medial JSW in those receiving the 0.23 mg dose of lorecivivint was significantly different from that in the placebo group (between-group difference, 0.19 mm, 95% CI 0.02, 0.36; \( P = 0.032 \)). At week 52, the mean change in medial JSW in the 0.03 mg and 0.07 mg lorecivivint dose groups was similar to that seen at week 26, whereas the mean change in the 0.23 mg lorecivivint dose group and placebo group had declined (Figure 4A). Among all subjects, both treatment with 0.03 mg lorecivivint (at week 52, change from baseline −0.04 mm) and treatment with 0.07 mg lorecivivint (at week 52, change from baseline −0.09 mm), but not treatment with 0.23 mg lorecivivint, maintained the medial JSW when compared with that in the placebo group (at week 52, change from baseline −0.14 mm); however, the differences were not statistically significant.

**UNI group.** Subjects in the UNI subgroup treated with 0.07 mg lorecivivint showed improvements in the medial JSW at weeks 26 and 52 (mean change from baseline 0.26 mm and 0.19 mm, respectively), whereas subjects in the UNI subgroup treated with placebo showed worsening of medial JSW (mean change from baseline −0.26 mm and −0.21 mm, respectively) (Figure 4B). The differences between the 0.07 mg lorecivivint group and the placebo group were significant at both time points; the mean change in medial JSW was 0.52 mm (95% CI 0.15, 0.89) at week 26 (\( P = 0.006 \)) and 0.39 mm (95% CI 0.06, 0.72) at week 52 (\( P = 0.021 \)). Among these subjects with unilateral symptomatic knee OA, there were no significant differences in the change in medial JSW when comparing the 0.03 mg or 0.23 mg lorecivivint group with the placebo group (Figure 4B).

**UNI WP− group.** In the UNI WP− subgroup, the 0.07 mg lorecivivint treatment group demonstrated improved medial JSW at week 26 (mean change from baseline 0.28 mm) and week 52 (mean change from baseline 0.17 mm), whereas the placebo treatment group had worsening of medial JSW at both time points (mean change from baseline −0.26 mm and −0.26 mm, respectively) (Figure 4C). The differences between the 0.07 mg lorecivivint group and the placebo group were significant at both time points (between-group difference, 0.53 mm, 95% CI 0.10, 0.97 at week 26 \( P = 0.016 \)) and 0.42 mm, 95% CI 0.04, 0.80 at week 52 \( P = 0.032 \)). Subjects with unilateral symptomatic OA but without widespread pain in the 0.03 mg or 0.23 lorecivivint treatment groups showed no significant differences in change in medial JSW compared with the placebo group (Figure 4C).

### Table 2. TEAEs with a reported frequency of >1%, by treatment group*

| Reported TEAEs | Lorecivivint | Placebo | All subjects† |
|---------------|-------------|---------|---------------|
|               | 0.03 mg (n = 111) | 0.07 mg (n = 114) | 0.23 mg (n = 104) | Placebo (n = 108) | All subjects (n = 452) |
| Total TEAEs   | 142/61 (55.0%) | 147/65 (57.0%) | 107/47 (45.2%) | 117/53 (49.1%) | 547/237 (52.4%) |
| Arthralgia    | 16/13 (11.7%) | 14/13 (11.4%) | 13/9 (8.7%) | 12/10 (9.3%) | 61/49 (10.8%) |
| Back pain     | 0/0 (0.0%) | 2/2 (1.8%) | 1/1 (1.0%) | 2/2 (1.9%) | 5/5 (1.1%) |
| Bronchitis    | 2/2 (1.8%) | 3/2 (1.8%) | 3/2 (1.9%) | 3/2 (1.9%) | 7/6 (1.3%) |
| Cystitis      | 0/0 (0.0%) | 3/3 (2.6%) | 2/1 (1.0%) | 1/1 (0.9%) | 6/5 (1.1%) |
| Fall          | 2/2 (1.8%) | 2/2 (1.8%) | 0/0 (0.0%) | 1/1 (0.9%) | 5/5 (1.1%) |
| Gastroenteritis | 3/3 (2.7%) | 0/0 (0.0%) | 1/1 (1.0%) | 1/1 (0.9%) | 5/5 (1.1%) |
| Headache      | 0/0 (0.0%) | 6/3 (2.6%) | 2/2 (1.9%) | 4/4 (3.7%) | 13/10 (2.2%) |
| Hypertension  | 0/0 (0.0%) | 4/4 (3.5%) | 4/4 (3.8%) | 3/3 (2.8) | 11/11 (2.4) |
| Increased AST level | 2/2 (1.8%) | 1/1 (0.9%) | 0/0 (0.0%) | 2/2 (1.9%) | 5/5 (1.1) |
| Influenza     | 4/4 (3.6%) | 0/0 (0.0%) | 2/2 (1.9%) | 0/0 (0.0%) | 6/6 (1.3) |
| Joint effusion| 5/4 (3.6%) | 2/2 (1.8%) | 1/1 (1.0%) | 2/2 (1.9%) | 10/9 (2.0) |
| Joint injury  | 2/2 (1.8%) | 0/0 (0.0%) | 1/1 (1.0%) | 1/1 (0.9%) | 6/6 (1.3) |
| Joint swelling| 5/3 (2.7%) | 4/4 (3.5%) | 2/2 (1.9%) | 6/5 (4.6) | 17/14 (3.1) |
| Meniscus injury | 2/2 (1.8%) | 2/2 (1.8%) | 0/0 (0.0%) | 0/0 (0.0%) | 5/5 (1.1) |
| Nasopharyngitis| 4/4 (3.6%) | 3/3 (2.6%) | 3/3 (2.9%) | 3/3 (2.9%) | 11/11 (2.4) |
| Nausea        | 2/2 (1.8%) | 1/1 (0.9%) | 2/2 (1.9%) | 1/1 (0.9%) | 6/6 (1.3) |
| Noncardiac chest pain | 1/1 (0.9%) | 2/2 (1.8%) | 1/1 (1.0%) | 1/1 (0.9%) | 6/6 (1.3) |
| Osteoarthritis | 4/3 (2.7%) | 2/2 (1.8%) | 3/3 (2.9) | 5/5 (3.8) | 14/11 (2.4) |
| Sinusitis     | 1/1 (0.9%) | 2/2 (1.8%) | 1/1 (1.0%) | 1/1 (0.9%) | 6/6 (1.3) |
| Tendinitis    | 3/3 (2.7%) | 1/1 (0.9%) | 1/1 (1.0%) | 1/1 (0.9%) | 6/6 (1.3) |
| Upper respiratory tract infection | 5/5 (4.5%) | 2/2 (1.8%) | 1/1 (1.0%) | 3/3 (2.8) | 12/12 (2.7) |
| Urinary tract infection | 2/2 (1.8%) | 2/2 (1.8%) | 3/2 (1.9) | 3/3 (2.8) | 10/9 (2.0) |

* Values are the number of reported treatment-emergent adverse events (TEAEs)/number of unique subjects reporting the event (% of treatment group). AST = aspartate aminotransferase.
† The group of all subjects includes those who received a dose of lorecivivint or placebo that was not specified per protocol (n = 15).
Concordance between change in medial JSW and clinical response. In the all-subjects analysis, no treatment group achieved an AUC of >0.7 (a measure of concordance between change in medial JSW and clinical response) (see Supplementary Figure 2A, available on the Arthritis & Rheumatology website at http://onlinelibrary.wiley.com/doi/10.1002/art.41315/abstract). Among subjects receiving the 0.07 mg dose of lorecivivint, concordance was “acceptable” (AUC 0.783) in the UNI subgroup and “excellent” (AUC 0.825) in the UNI WP− subgroup (Supplementary Figures 2B and C [http://onlinelibrary.wiley.com/doi/10.1002/art.41315/abstract]). No other doses in either subgroup achieved an AUC >0.7.

Safety. No clinically significant safety concerns with respect to vital signs, clinical laboratory results, or AEs were observed; rates were comparable between the lorecivivint and placebo groups. No deaths were reported during the study. Fifteen subjects incorrectly received a study injection that diverged from that prescribed by the protocol; these subjects are described as “Other” in the safety analysis.

In total, 547 TEAEs were reported by 237 subjects (52.4% of 452 subjects), of which 40 AEs in 32 subjects (7.1%) were deemed related to the study drug by the investigator (Table 2). Among the treatment groups, 142 TEAEs were reported by 61 subjects in the 0.03 mg cohort (55.0% of 111 treated subjects), 147 TEAEs by 65 subjects in the 0.07 mg cohort (57.0% of 114 treated subjects), 107 TEAEs by 47 subjects in the 0.23 mg cohort (45.2% of 104 treated subjects), 117 TEAEs by 53 subjects in the placebo cohort (49.1% of 108 treated subjects), and 34 TEAEs by 11 subjects who were classified in the “Other” dose group (73.3% of 15 subjects). Arthralgia, defined for this study as an exacerbation (increase in frequency, severity, or specificity) of an existing condition, was the most common AE reported across sources. These results inform the design of future lorecivivint trials (36); in fact, the slowing of joint space narrowing has been recommended as an appropriate structural end point for DMOAD trials (36). In the all-subjects analysis, both 0.03 mg lorecivivint and 0.07 mg lorecivivint would be found to produce clinically meaningful improvements from baseline in the WOMAC pain and function subscales.

Analysis of the prospec ted UNI subgroup showed greater improvements in WOMAC pain scores, WOMAC function scores, and medial JSW for the 0.07 mg cohort compared with the placebo cohort. These differences appeared to be further enhanced in the post hoc analysis of subjects in the UNI WP− subgroup receiving 0.07 mg lorecivivint. Pain reporting by subjects with bilateral symptomatic knee OA is known to be complicated, not only because of the presence of contralateral knee pain, but also because other joints may be affected by OA (33,34), nociceptive biomechanical factors may be involved, and other centralized pain conditions (e.g., fibromyalgia) may be present (35). Therefore, the improvements compared with placebo observed in both the UNI and the UNI WP− subgroups versus the all-subjects group after an IA injection of lorecivivint into the target knee may be attributable to a predominance of subjects with unilateral OA symptoms being able to discriminate their target knee pain from other pain sources. These results inform the design of future lorecivivint trials by identifying a target population in whom potential symptomatic efficacy could be more clearly delineated.

In addition to symptom improvements, inhibition of structural progression is a key goal of disease modification in OA (36); in fact, the slowing of joint space narrowing has been recommended as an appropriate structural end point for DMOAD trials (36). In the all-subjects analysis, both 0.03 mg lorecivivint (at week 52, change from baseline in medial JSW −0.04 mm) and 0.07 mg lorecivivint (at week 52, change from baseline in medial...
JSW −0.09 mm) maintained, at least numerically, the medial JSW at week 52, but neither dose achieved a significant difference when compared with placebo (at week 52, change from baseline in medial JSW −0.14 mm). Subjects with unilateral symptomatic OA (i.e., both the UNI and the UNI WP− subgroups) treated with 0.07 mg lorecivivint showed mean medial JSW increases beyond a 0.13-mm minimum detectable difference (23), whereas subjects who received placebo showed decreases (narrowing) in mean medial JSW from baseline. Joint space narrowing has also been correlated with clinical outcomes, including an increased risk of total knee replacement in those with joint space narrowing of >0.5 mm over 2 years (an outcome indicative of treatment failure) (36,37). Knee OA is associated with typical joint space narrowing >0.5 mm over 2 years (an outcome indicative of treatment failure) (36,37). Knee OA is associated with typical joint space narrowing of 0.1–0.3 mm per year (38,39); although such changes require

Knee OA is associated with typical joint space narrowing of 0.1–0.3 mm per year (38,39); although such changes require precise and reproducible measurement methods, such as the positioned, fixed-flexion radiography technique employed herein, the accuracy of the knee radiographic measurement of medial JSW can range from 0.04 mm to 0.5 mm (23,40–42). The relative improvement in medial JSW in the unilateral symptomatic subject subgroups may also be related to a more favorable local biomechanical environment in individuals with unilateral knee pain (43,44).

A post hoc analysis of both the UNI and the UNI WP− subgroups demonstrated that the radiographic findings (medial JSW) and clinical findings (WOMAC pain and function scores) in the 0.07 mg dose group were concordant (i.e., the change in the former is associated with change in the latter). This suggested a connection between improvement in structural measures and improvement in clinical responses. The 2018 draft guidance from the US Food and Drug Administration on OA structural end points suggests that additional data are needed to support the relationship between structural measurements and clinical outcomes; this analysis sought to contribute to this growing evidence base.

This phase IIa study has several limitations, including no formal, preplanned sample size or power calculation and considerable placebo responses for patient-reported outcomes, similar to those demonstrated in other OA trials (5,45). Although trials investigating IA therapies for knee OA commonly use saline as a placebo comparator arm, evidence suggests that IA saline might actually be therapeutic (46). Therefore, further studies of lorecivivint in larger clinical trials with refined inclusion criteria (e.g., focusing on subjects with unilateral symptomatic OA, as lorecivivint is administered into the single most painful knee) are needed to disentangle the active treatment effects from the placebo effects. Although radiographic medial JSW represents an objective measure for assessing structural progression, the evidence supporting the usefulness of this measurement is not definitive and other imaging modalities, such as magnetic resonance imaging, may also be considered. Larger and longer studies are needed to determine the best methods for assessing the disease-modifying abilities of drugs in knee DMOAD trials.

Finally, the primary statistical analysis Type 1 error control strategy was not achieved, leading to all statistical results being considered exploratory. Since the (prespecified) UNI and (post hoc) UNI WP− groups were small with respect to the number of subjects, the results of these exploratory analyses in both groups are considered to be hypothesis generating, and thus require validation in a prospective trial.

In summary, although the primary end point in all subjects was not met, treatment of subjects with moderately to severely symptomatic knee OA in the UNI and UNI WP− subgroups with an IA injection of 0.07 mg lorecivivint resulted in numerical improvements in pain, function, and medial JSW compared with placebo. Furthermore, treatment with 0.07 mg of lorecivivint demonstrated the greatest improvements in WOMAC pain and function scores and the highest concordance between symptom relief and structural changes. This study identified a target group of subjects with unilateral symptomatic knee OA and a potentially optimal dose of lorecivivint (0.07 mg). The clinical and radiographic outcomes warrant additional studies of the potential of lorecivivint for both analgesia and disease-modifying activity in knee OA.

**AUTHOR CONTRIBUTIONS**

All authors were involved in drafting the article or revising it critically for important intellectual content, and all authors approved the final version to be published. Dr. Yazici had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

**Study conception and design.** Yazici, Swearingen, DiFrancesco, Simsek, Tambiah.

**Acquisition of data.** Yazici, Swearingen, DiFrancesco.

**Analysis and interpretation of data.** Yazici, McAlindon, Gibofsky, Lane, Clauw, Jones, Swearingen, DiFrancesco, Bergfeld, Simsek, Tambiah, Hochberg.

**ROLE OF THE STUDY SPONSOR**

Samumed, LLC designed, funded, and monitored the study and also conducted the data management and statistical analysis. The authors independently interpreted the results and had the final decision to submit the manuscript for publication. Publication of this article was not contingent upon approval by Samumed, LLC.

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The patient, a 66-year-old man, presented with a 1-week history of left elbow pain and swelling. He denied experiencing any prior trauma or fever. He had a history of diabetes mellitus, hypertension, hyperlipidemia, and chronic kidney disease. Five years earlier, he had developed podagra and was diagnosed as having gout, which was treated with indomethacin. He did not take any urate-lowering medication. At current presentation, he reported experiencing multiple episodes of gout per year, mostly affecting the feet. Physical examination of the left elbow revealed swelling, tenderness, warmth, erythema, and decreased range of motion. Examination of synovial fluid by polarized light microscopy revealed cholesterol crystals and monosodium urate monohydrate (MSU) crystals. In A and B, cholesterol appears as plate-like structures with notched corners and MSU appears as needle-shaped crystals with negative birefringence (appearing yellow or blue when viewed with a first-order red plate compensator parallel or perpendicular to the axis, respectively; arrows depict the axis of the compensator) (original magnification × 200 in A; × 400 in B). Gram stain and bacterial cultures were negative. Symptoms resolved after a 5-day course of oral glucocorticoid treatment; after which urate-lowering therapy was started. Cholesterol crystals are occasionally seen in chronically inflamed joints in rheumatoid arthritis (1,2) and bursa effusions (2), and there have been rare reports of their presence in chronic tophaceous gout (3). They are extracellular and can be present in association with Maltese cross-shaped fat droplets. Cholesterol crystals typically appear as large, flat, rectangular plates and have notched corners with varying birefringence. These features enable differentiation from MSU crystals, which are needle-shaped and strongly negatively birefringent.

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