Abstract

**IMPORTANCE**  Vamorolone is a synthetic steroidal drug with potent anti-inflammatory properties. Initial open-label, multiple ascending dose-finding studies of vamorolone among boys with Duchenne muscular dystrophy (DMD) found significant motor function improvement after 6 months treatment in higher-dose (ie, ≥2.0 mg/kg/d) groups.

**OBJECTIVE**  To investigate outcomes after 30 months of open-label vamorolone treatment.

**DESIGN, SETTING, AND PARTICIPANTS**  This nonrandomized controlled trial was conducted by the Cooperative International Neuromuscular Research Group at 11 US and non-US study sites. Participants were 46 boys ages 4.5 to 7.5 years with DMD who completed the 6-month dose-finding study. Data were analyzed from July 2020 through November 2021.

**INTERVENTIONS**  Participants were enrolled in a 24-month, long-term extension (LTE) study with vamorolone dose escalated to 2.0 or 6.0 mg/kg/d.

**MAIN OUTCOMES AND MEASURES**  Change in time-to-stand (TTSTAND) velocity from dose-finding baseline to end of LTE study was the primary outcome. Efficacy assessments included timed function tests, 6-minute walk test, and NorthStar Ambulatory Assessment (NSAA). Participants with DMD treated with glucocorticoids from the Duchenne Natural History Study (DNHS) and NorthStar United Kingdom (NSUK) Network were matched and compared with participants in the LTE study receiving higher doses of vamorolone.

**RESULTS**  Among 46 boys with DMD who completed the dose-finding study, 41 boys (mean [SD] age, 5.33 [0.96] years) completed the LTE study. Among 21 participants treated with higher-dose (ie, ≥2.0 mg/kg/d) vamorolone consistently throughout the 6-month dose-finding and 24-month LTE studies with data available at 30 months, there was a decrease in mean (SD) TTSTAND velocity from baseline to 30 months (0.206 [0.070] rises/s vs 0.189 [0.124] rises/s), which was not a statistically significant change (−0.011 rises/s; CI, −0.068 to 0.046 rises/s). There were no statistically significant differences between participants receiving higher-dose vamorolone and matched participants in the historical control groups receiving glucocorticoid treatment (75 patients in DNHS and 110 patients in NSUK) over a 2-year period in NSAA total score change (0.22 units vs NSUK; 95% CI, −4.48 to 4.04); P = .92), body mass index z score change (0.002 vs DNHS SD/mo; 95% CI, −0.006 to 0.010; P = .58), or timed function test change. Vamorolone at doses up to 6.0 mg/kg/d was well tolerated, with 5 of 46 participants discontinuing prematurely and for reasons not associated with study drug. Participants in the DNHS treated with glucocorticoids had significant growth delay in comparison (continued)
Abstract (continued)

with participants treated with vamorolone who had stable height percentiles (0.37 percentile/mo; 95% CI, 0.23 to 0.52 percentile/mo) over time.

CONCLUSIONS AND RELEVANCE This study found that vamorolone treatment was not associated with a change in TTSTAND velocity from baseline to 30 months among boys with DMD aged 4 to 7 years at enrollment. Vamorolone was associated with maintenance of muscle strength and function up to 30 months, similar to standard of care glucocorticoid therapy, and improved height velocity compared with growth deceleration associated with glucocorticoid treatment, suggesting that vamorolone may be an attractive candidate for treatment of DMD.

TRIAL REGISTRATION ClinicalTrials.gov Identifier: NCT03038399

Introduction

Duchenne muscular dystrophy (DMD) is a rapidly progressive X-linked recessive disease, with symptom onset before age 6 years in boys. It is the most common form of muscular dystrophy in childhood, occurring among 1 in 3600 to 9300 male newborns worldwide. The disease is caused by variants in the DMD gene, which codes for the dystrophin protein. The lack of dystrophin results in membrane fragility, necrosis, inflammation, and progressive muscle atrophy. Aberrant intracellular signaling cascades also contribute to DMD pathophysiology. Upregulated inflammatory gene expression and activated immune cell infiltrates, at least partially mediated by nuclear factor κ light chain enhancer of activated B cells (NF-κB) activation, are evident during early disease stages and play a significant role in disease progression. NF-κB regulates expression of numerous inflammatory genes in immune cells and muscle fibers, and the infiltration and activation of these cells can trigger muscle cell death.

Despite scientific advances, the only medications that have consistently demonstrated efficacy in clinical trials for DMD are glucocorticoids (GCs), such as prednisone and deflazacort. Most gene-targeted disease-modifying technologies that are currently in development or have been recently approved focus on subsets of dystrophin variants and therefore do not address the unmet need among all individuals with DMD. Moreover, these treatments are prescribed in combination with and not as alternatives to GCs. The adverse outcomes associated with GCs, including weight gain, appearance associated with Cushing syndrome, short stature, behavior changes, low bone density, bone fractures, and cataracts, negatively impact quality of life among individuals with DMD. Other GC-associated safety concerns include adrenal suppression, delayed puberty, insulin resistance, and further muscle damage. The significant adverse growth and development outcomes associated with GCs hinder their routine administration among children aged younger than 2 years, even though earlier administration may be associated with better overall functional outcomes. A current goal of DMD research is to find a variant-independent treatment that matches or exceeds the efficacy of GC with a significantly better adverse outcome profile.

Vamorolone is a first-in-class steroidal anti-inflammatory drug that differs from conventional GCs in lacking an 11β hydroxy-carbonyl group. This difference removes a contact site with the target glucocorticoid receptor and significantly alters structure and activity associations. Additionally, unlike GC, vamorolone is not a substrate for 11β-hydroxysteroid dehydrogenase regulatory enzymes and is a potent antagonist of the mineralocorticoid receptor.

A Phase Ila 2-week (VBPI5-002, NCT02760264) multiple ascending dose study of vamorolone (developmental code name, VBP) among boys with DMD who were GC naive and aged 4 years to less than 7 years was completed in 2017. Based on clinical safety and pharmacodynamic biomarker analysis, vamorolone showed dose-dependent anti-inflammatory activity.
Phase IIa 24-week (VBP15-003, NCT02760277) open-label dose-finding study suggested efficacy of vamorolone treatment at 2.0 and 6.0 mg/kg/d based on motor outcomes. Interim analysis at 18 months of the vamorolone open-label extension study (VBP15-LTE, NCT03038399) found ongoing improvement in motor function and a favorable safety profile. Here, we report on the long-term efficacy, safety, and tolerability of vamorolone among boys with DMD who completed the 6-month dose-finding and 24-month long-term extension trials, with a total of up to 30 months of vamorolone treatment.

Methods

This nonrandomized clinical study and all studies contributing patients were approved by ethics committees or institutional review boards as required by the participating international academic clinical recruitment sites. Written informed consent was obtained from parents or legal guardians, and assent was obtained from children recruited into each described trial. This report follows the Transparent Reporting of Evaluations With Nonrandomized Designs (TREND) reporting guideline for nonrandomized studies.

The Cooperative International Neuromuscular Research Group (CINRG) conducted 3 consecutive multicenter open-label vamorolone clinical trials. Participants were recruited from 11 participating centers in the United States, Canada, United Kingdom, Sweden, Israel, and Australia (Figure 1). The 2-week multiple-ascending-dose study enrolled 48 boys (ages 4 to <7 years) with DMD who were GC-naive from June 2016 to October 2017. All participants completed the initial study and were enrolled in a 24-week dose-finding trial; 2 participants subsequently withdrew for reasons unrelated to the study drug. The remaining 46 participants (ages 4.5-7.5 years) who completed the 6-month dose-finding study were eligible to enroll in a 24-month long-term extension (LTE) study, which started in February 2017 and was completed in April 2020.

Study Design

Throughout the studies, vamorolone was provided as a 4% flavored oral suspension taken daily at breakfast with a 240-mL glass of whole milk (approximately 8 g of fat) or equivalent high-fat food portion. Study participants were initially assigned to receive vamorolone at 1 of 4 dose levels (ie, 0.25, 0.75, 2.0, or 6.0 mg/kg/d) (eFigure 1 in Supplement 1) and remained at the same dose level in the 2-week multiple ascending dose and 24-week dose-finding trials. The last visit of the 24-week dose-finding trial coincided with the baseline visit of the 24-month LTE trial. Multiple dose escalations to the highest dose (ie, 6.0 mg/kg/d) were permitted in the LTE protocol (Supplement 2); de-escalations were also allowed in case of intolerability, at the discretion of investigators. Given the variable timing of dose escalations, prespecified analyses of drug-associated efficacy focused on 23 participants who were initially assigned and maintained on a 2.0 mg/kg/d or greater dose of vamorolone (ie, 2 mg/kg/d and 6 mg/kg/d dose groups, hereafter higher-dose groups) for up to 30 months.

Additionally, participants from the CINRG Duchenne Natural History Study (DNHS) treated with GC were selected to serve as a historical control group for timed function tests and anthropometric measurements. The DNHS (NCT00468832) included 440 participants with DMD, with data collected from December 2005 to November 2016. Participants in DNHS were first eligible for inclusion in the control group after they had experienced 6 months of continuous GC exposure, with similar ages (ie, 4.5-7.5 years) as participants in the LTE study who had 6 months of vamorolone exposure at baseline after completing the dose-finding trial. For statistical matching between participants receiving higher-dose vamorolone and those in the DNHS, prespecified criteria (eAppendix 1 in Supplement 1) were defined for coarsened exact matching (CEM).

In addition, participants from the NorthStar United Kingdom (NSUK) Network were selected for inclusion in a second historical control group to allow for comparison of NorthStar Ambulatory Assessment (NSAA) scores, given that NSAA was not assessed for most participants in DNHS.
Participants in NSUK included in this comparison were recruited from August 2005 to October 2019. Like participants from the DNHS, they were started on GC (any type or dose) for 6 months before the comparison baseline visit, with similar ages (4.5-7.5 years) at baseline compared with participants in the LTE, and were continuously maintained on GC (eAppendix 2 in Supplement 1).

**Measurements**

Study visits took place quarterly and included clinical and laboratory assessments with adverse event reporting. Standing height and weight were assessed at each study visit; height percentile, body mass index (BMI; calculated as weight in kilograms divided by height in meters squared) and BMI z score were calculated centrally. Clinical efficacy evaluations were performed at baseline (ie, multiple ascending dose entry), 6 months (ie, LTE baseline), 18 months (ie, LTE month 12), and 30 months (ie, LTE month 24) of treatment with vamorolone. Assessment of efficacy included timed function tests (ie, time to stand from supine [TTSTAND], time to run or walk 10 m [TTRW], time to climb 4 stairs [TTCLIMB]), distance covered in 6-minute walk test (6MWT), NSAA score (total NSAA score is made up of the sum of scores on 17 ordinal tests, scored as 0, 1, or 2, with a 2 indicating normal activity with no obvious modification) and quantitative muscle testing (QMT) using the CINRG Quantitative Measurement System (measurements in pounds done unilaterally using the dominant side, if known, for each muscle group [ie, knee extension, knee flexion, elbow extension, and elbow flexion]). TTSTAND change from the multiple ascending dose study baseline to LTE month 24 was the primary efficacy end point. For TTSTAND, TTRW, and TTCLIMB, results in seconds

**Figure 1. Flowchart of Vamorolone Long-term Extension (LTE) Study**

VBP indicates vamorolone.

* Treating physicians were permitted to up-titrated or down-titrated dose according to clinical judgement. There were 3 participants who ended the study at an intermediate dose of 4.0 mg/kg/d.
were converted to velocities (ie, rises per second for TTSTAND, meters per second for TTRW, and tasks per second for TTCLIMB); participants unable to perform tests owing to disease progression had a velocity score imputed as 0.25. Clinical evaluators were trained according to operating procedures that were standardized in the NSUK Network and vamorolone and DNHS participating centers.

Patient-reported outcomes and health-related quality of life were assessed by the Pediatric Outcomes Data Collection Instrument (PODCI) as an exploratory efficacy end point. The PODCI standardized Upper Extremity and Physical Function score is calculated using at least 4 items with valid nonmissing responses, with a possible range of 0 to 100. The standardized Transfer and Basic Mobility score is calculated using at least 7 items with valid nonmissing response, with a possible range of 0 to 100. Safety measurements included physical exam with weight and height, BMI, clinical laboratory tests, 12-lead electrocardiogram, bone age, lateral thoracolumbar spine x-ray, and adverse event (AE) reporting. All AEs were coded using the Medical Dictionary for Regulatory Activities (MedDRA, versions 19.0, 20.1, 21.0, and 21.1) system for reporting. Intensity of AEs was graded using the Common Terminology Criteria for Adverse Events version 4.03.

**Statistical Analysis**

A statistical analysis plan prespecified the longitudinal within-study (based on paired t test) analyses of motor outcomes, biomarkers, and safety analyses (Supplement 2). Summaries for TTSTAND, TTRW, TTCLIMB, 6MWT, NSAA, and QMT included descriptive statistics on observed values and changes from baseline values at each time.

Comparison of participants in the LTE study receiving higher doses with the historical control group consisting of participants from DNHS receiving GC was based on longitudinal outcome data using a mixed-effect model with repeated measures (MMRM). Owing to NSUK participant-level data sharing restrictions, NSAA data were analyzed separately by NSUK researchers and summaries of change were compared with the higher-dose LTE cohort using independent t tests and a relative risk analysis of NSAA scores. Nonparametric maximum likelihood estimation (NPMLE) was used to estimate the observed survival curve (median time to event and bootstrapped CIs) for the first TTSTAND milestone of 10 or more seconds to assess the risk associated with future loss of ambulation for each cohort.

For the post hoc delayed-start analysis, 23 participants initially assigned to the vamorolone low-dose groups (ie, 0.25 and 0.75 mg/kg/d) during the first 6 months of the dose-finding study were considered to have delayed starts. Outcomes for these participants were compared with outcomes among 23 participants who were initially assigned to and retained on higher-dose vamorolone (ie, 2.0 and 6.0 mg/kg/d) throughout their follow-up and were considered to have early starts. Estimates and standard errors were calculated using MMRM.

Data manipulation, tabulation of descriptive statistics, and graphical representations were performed primarily using statistical software SAS version 9.4 or higher (SAS Institute) and R version 4.1.0 (R Project for Statistical Computing). The following R packages were used: interval, lme4, lmerTest, sjPlot, visreg, and vistime. All statistical tests were 2-sided, and a resultant P value ≤ .05 was considered statistically significant. No adjustments for multiplicity on inferential statistics were made as per the statistical analysis plan (Supplement 2). Data were analyzed from July 2020 through November 2021. More details regarding statistical modeling, as well as real-world comparator populations (ie, historical control groups with different GCs [prednisone or deflazacort], and regimen [daily or intermittent]) are provided in eAppendix 3 in Supplement 1.

**Results**

Among 46 boys with DMD who completed the dose-finding study, 41 participants (89.1%; baseline mean [SD] age, 5.33 [0.96] years) completed the 2-year LTE treatment period; 5 participants withdrew for reasons unrelated to the study drug (Figure 1). At the LTE month 24 visit, 11 participants,
3 participants, and 27 participants were being treated at 2.0 mg/kg/d, 4.0 mg/kg/d, and 6.0 mg/kg/d, respectively (eFigure 1 in Supplement 1). Analysis of treatment efficacy focused on 23 participants receiving higher-dose vamorolone who were initially assigned and maintained on 2.0 mg/kg/d or more for up to 30 months during the dose-finding study (6 months) and LTE study (24 months). Demographic and clinical characteristics of participants in the LTE study receiving higher doses are summarized in Table 1. The mean (SD) baseline age was 5.83 (0.88) years in the higher-dose LTE group. In the higher-dose LTE group, 3 participants (13.0%) withdrew before the LTE month 24 study visit. The subsets of participants receiving higher doses who completed all planned outcomes are summarized in eAppendix 3 in Supplement 1.

Changes in Functional Outcomes Among Participants in LTE With Higher Doses

Among 23 participants started on higher-dose vamorolone, 21 individuals were treated throughout the dose-finding and LTE studies and had available data at 30 months (3 individuals dropped out, but for 1 individual, terminal visit data was available close to 30 months and counted as such based on procedures). In these individuals, there was a decrease from baseline to 30 months in mean (SD) TTSTAND velocity (0.206 [0.070] rises/s vs 0.189 [0.124] rises/s), which was not a statistically significant change (−0.011 rises/s; 95% CI, −0.068 to 0.046 rises/s). Similarly, there were no significant changes from baseline to 30 months in TTCLIMB velocity score among 18 participants with TTCLIMB data (0.035 tasks/s; 95% CI, −0.051 to 0.121 tasks/s), TTRW velocity score among 18 participants with TTRW data (0.061 m/s; 95% CI, −0.272 to 0.394 m/s), 6MWT total distance among 15 participants with 6MWT data (32.0 m; 95% CI, −18.95 to 82.95 m), and total NSAA score among 18 participants with NSAA data (1.6; 95% CI, −2.92 to 6.14). There was no statistically significant change in PODCI transfer and basic mobility scores from baseline to 30 months (eTable 2 in Supplement 1), and interpretation of QMT was limited by few observations after 30 months (eTable 3 in Supplement 1).

In a delayed start analysis, participants initiated on higher-dose vamorolone (ie, those with early starts) had better clinical outcomes at 6 months compared with those initially treated with low doses (ie, those with delayed starts). At 6, 18, and 30 months, the means of all 5 motor outcome measures were increased for participants with early starts compared with those with delayed starts, although

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**Table 1. Participant Demographic and Baseline Characteristics for External Comparisons**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Participants Higher-dose VBP15-LTE (N = 23)b</th>
<th>GC in CINRG DNHS (N = 75)</th>
<th>GC in NorthStar UK Network (N = 110)c</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at baseline comparison visit, y</td>
<td>Mean (SD) 5.83 (0.88)</td>
<td>6.08 (0.81)</td>
<td>6.00 (0.77)</td>
</tr>
<tr>
<td></td>
<td>Median (range) 5.65 (4.60-7.29)</td>
<td>5.97 (4.50-7.48)</td>
<td>5.94 (NA)</td>
</tr>
<tr>
<td>Steroid exposure at baseline comparison visit, mean (SD), d</td>
<td>200.57 (7.54)</td>
<td>227.73 (61.91)</td>
<td>264.84 (57.39)</td>
</tr>
<tr>
<td>Duration of follow up from baseline visit, mean (SD), y</td>
<td>1.85 (0.46)</td>
<td>1.36 (0.49)</td>
<td>NA</td>
</tr>
<tr>
<td>Participants with &gt;18 mo follow-up after initial 6 mo of steroid exposure, No. (%)</td>
<td>21 (91.3)</td>
<td>30 (40.0)</td>
<td>NA</td>
</tr>
<tr>
<td>Weight (kg), No.</td>
<td>23</td>
<td>73</td>
<td>NA</td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>21.98 (3.78)</td>
<td>20.35 (3.55)</td>
<td>NA</td>
</tr>
<tr>
<td>Median (range)</td>
<td>22.50 (16.30-32.60)</td>
<td>20.10 (14.80-32.60)</td>
<td>NA</td>
</tr>
<tr>
<td>Height (cm), No.</td>
<td>22</td>
<td>73</td>
<td>NA</td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>111.80 (6.94)</td>
<td>109.86 (6.86)</td>
<td>NA</td>
</tr>
<tr>
<td>Median (range)</td>
<td>113.05 (100.60-123.00)</td>
<td>110.00 (96.10-126.30)</td>
<td>NA</td>
</tr>
<tr>
<td>Body Mass Index, No.</td>
<td>22</td>
<td>72</td>
<td>NA</td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>17.68 (1.23)</td>
<td>16.68 (1.55)</td>
<td>NA</td>
</tr>
<tr>
<td>Median (range)</td>
<td>17.65 (15.90-21.50)</td>
<td>16.56 (12.19-20.44)</td>
<td>NA</td>
</tr>
</tbody>
</table>

Abbreviations: CINRG, Cooperative International Neuromuscular Research Group; DNHS, Duchenne Natural History Study; GC, glucocorticoid; LTE, long-term extension; NA, not applicable; VBP, vamorolone.

a Based on baseline comparator visit, which corresponds to approximately 6 months of steroid exposure for participants in the NorthStar UK Network and CINRG DNHS.

b The higher-dose cohort refers to participants assigned to vamorolone 2.0 and 6.0 mg/kg/d from initial dose-finding (VBP15-002/-003) studies and maintained at 2.0 mg/kg/d or more from long-term extension (VBP15-LTE) baseline.

c Owing to NorthStar UK Network participant-level data-sharing restrictions, limited summary information was available.

d Reported for time to stand velocity.
not all differences were statistically significant. At 30 months, no means were statistically significantly different (Figure 2).

**Efficacy of Higher-Dose Vamorolone Compared With GC Treatment in Historical Control Groups**

The mean (SD) baseline age was 6.08 (0.81) years among 75 participants in the DNHS control group and 6.00 (0.77) years among 110 participants in the NorthStar UK Network control group. An

![Figure 2. Efficacy of Vamorolone in Motor Function Assessments by High vs Low Starting Dose](https://jamanetwork.com/)

6MWT indicates 6-minute walk test; high, a starting dose of 2.0 mg/kg/d or greater; low, a starting dose of less than 2.0 mg/kg/d; points, means; whiskers, standard error of the mean. Mean estimated differences between participants treated at higher dose and low dose, as well as P values, are provided at 6 months (ie, prior to LTE) and 30 months (ie, at conclusion of LTE).

**Figure 2. Efficacy of Vamorolone in Motor Function Assessments by High vs Low Starting Dose**

- **A** Change in time to stand velocity
  - High vs low: 0.067 (P < .001)
  - High vs low: 0.049 (P = .14)
  - High vs low, change from month 6 to 30: −0.018 (P = .52)

- **B** Time to climb velocity
  - High vs low: 0.033 (P < .001)
  - High vs low: 0.045 (P < .001)
  - High vs low, change from month 6 to 30: −0.013 (P = .79)

- **C** Time to run or walk
  - High vs low: 0.081 (P = .07)
  - High vs low: 0.224 (P = .28)
  - High vs low, change from month 6 to 30: 0.081 (P = .63)

- **D** 6MWT
  - High vs low: 33.4 (P = .01)
  - High vs low: 15.0 (P = .28)
  - High vs low, change from month 6 to 30: −6.4 (P = .55)

- **E** NSAA total score
  - High vs low: 4.18 (P = .06)
  - High vs low: 1.47 (P = .11)
  - High vs low, change from month 6 to 30: 2.71 (P = .23)
increased proportion of participants in the higher-dose LTE group had more than 18 months of follow-up after the initial 6 months of steroid exposure compared with participants in the DNHS control group (21 participants [91.3%] vs 30 participants [40.0%]). As noted previously, 75 participants from DNHS and 23 participants from the LTE study receiving higher doses were criteria-matched and subjected to CEM, yielding a smaller subset of matched comparator group (including 29 participants from DNHS and 20 participants from the LTE study). However, participants from DNHS who were CEM-matched had limited follow-up data, and the baseline characteristics of these participants with approximately 2 years of long-term follow-up (which would inform longitudinal analysis) were dissimilar to those from participants in the LTE study who were CEM-matched, resulting in unsuccessful CEM for longitudinal comparison (eTable 1 in Supplement 1). Hence, we conducted a prespecified sensitivity analysis among 75 participants from DNHS and 23 participants from LTE; we present these comparisons in Table 1 and Table 2.

In longitudinal comparison of mean TTSTAND, TTRW, and TTCLIMB velocity from baseline to end of follow-up, the LTE group and DNHS group were not significantly different (Table 2, Figure 3; eFigure 2 in Supplement 2). Mean TTSTAND velocity trajectories were not significantly different (estimates of difference from interaction of orthogonal polynomial terms with cohort: quadratic term coefficient estimate, 0.001; 95% CI, −0.30 to 0.30; P > .99; linear term coefficient estimate, −0.30; 95% CI, −0.66 to 0.05; P = .10). There was no significant difference in the NPMLE estimate of time to reach a TTSTAND milestone of 10 seconds or more between 22 participants in the higher-dose LTE

### Table 2. Comparison of Study Primary Efficacy and Safety Outcome Measures

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Unique participants (longitudinal samples used), No. (No.)</th>
<th>Baseline mean (SD)</th>
<th>End of follow up mean (SD)</th>
<th>Statistical model or test</th>
<th>Value (2-sided 95% CI)</th>
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</thead>
<tbody>
<tr>
<td><strong>Efficacy</strong></td>
<td></td>
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<tr>
<td>TTSTAND velocity, rises/s</td>
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<tr>
<td>Higher-dose vamorolone LTE&lt;sup&gt;c&lt;/sup&gt;</td>
<td>23 (66)</td>
<td>0.25 (0.10)</td>
<td>0.20 (0.13)</td>
<td>vs DNHS (MMRM)</td>
<td>&gt; .99 (−0.30 to 0.30); linear comparison: .10 (−0.66 to 0.05)</td>
</tr>
<tr>
<td>GC in DNHS</td>
<td>75 (311)</td>
<td>0.25 (0.10)</td>
<td>0.25 (0.13)</td>
<td>Quadratic comparison&lt;sup&gt;a&lt;/sup&gt;</td>
<td>&gt; .99 (−0.30 to 0.30); linear comparison: .10 (−0.66 to 0.05)</td>
</tr>
<tr>
<td>With ≥1.5 y follow-up&lt;sup&gt;b&lt;/sup&gt;</td>
<td>30 (153)</td>
<td>0.24 (0.09)</td>
<td>0.24 (0.12)</td>
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<tr>
<td>TTCLIMB velocity, tasks/s</td>
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<tr>
<td>Higher-dose vamorolone LTE&lt;sup&gt;c&lt;/sup&gt;</td>
<td>23 (63)</td>
<td>0.31 (0.13)</td>
<td>0.32 (0.19)</td>
<td>vs DNHS (MMRM)</td>
<td></td>
</tr>
<tr>
<td>GC in DNHS</td>
<td>75 (311)</td>
<td>0.32 (0.14)</td>
<td>0.33 (0.18)</td>
<td>Quadratic comparison&lt;sup&gt;a&lt;/sup&gt;</td>
<td>&gt; .99 (−0.36 to 0.36); linear comparison: .93 (−0.41 to 0.44)</td>
</tr>
<tr>
<td>With ≥1.5 y follow-up&lt;sup&gt;b&lt;/sup&gt;</td>
<td>30 (153)</td>
<td>0.31 (0.13)</td>
<td>0.32 (0.16)</td>
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<tr>
<td>TTRW velocity, meters/s</td>
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<td></td>
</tr>
<tr>
<td>Higher-dose vamorolone LTE&lt;sup&gt;c&lt;/sup&gt;</td>
<td>23 (63)</td>
<td>1.90 (0.34)</td>
<td>1.87 (0.63)</td>
<td>vs DNHS (MMRM)</td>
<td></td>
</tr>
<tr>
<td>GC in DNHS</td>
<td>75 (317)</td>
<td>1.91 (0.52)</td>
<td>1.89 (0.71)</td>
<td>Quadratic comparison&lt;sup&gt;a&lt;/sup&gt;</td>
<td>&gt; .99 (−1.24 to 1.41) Linear comparison: .94 (−1.57 to 1.46)</td>
</tr>
<tr>
<td>With ≥1.5 years follow-up&lt;sup&gt;b&lt;/sup&gt;</td>
<td>30 (153)</td>
<td>1.85 (0.37)</td>
<td>1.85 (0.64)</td>
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<tr>
<td>6MWT, m walked</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Higher-dose vamorolone LTE&lt;sup&gt;c&lt;/sup&gt;</td>
<td>20 (37)</td>
<td>377.9 (64.77)</td>
<td>369.9 (77.81)</td>
<td>Paired sample (within group) t test: .20 (−18.95 to 4.04)</td>
<td></td>
</tr>
<tr>
<td>NSAA score&lt;sup&gt;d&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Higher-dose vamorolone LTE&lt;sup&gt;c&lt;/sup&gt;</td>
<td>23 (41)</td>
<td>22.3 (4.72)</td>
<td>21.78 (7.86)</td>
<td>vs NorthStar UK cohort (independent t test on change over 2 y)&lt;sup&gt;f&lt;/sup&gt;</td>
<td>90 (−1.24 to 1.41) Linear comparison: .94 (−1.57 to 1.46)</td>
</tr>
<tr>
<td>GC in NSUK Network</td>
<td>110 (159)</td>
<td>26.63 (5.65)&lt;sup&gt;e&lt;/sup&gt;</td>
<td>26.24 (7.2)</td>
<td>.92 (−4.48 to 4.04)</td>
<td></td>
</tr>
<tr>
<td>Height percentile</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Higher-dose vamorolone LTE&lt;sup&gt;c&lt;/sup&gt;</td>
<td>23 (65)</td>
<td>32.26 (26.87)</td>
<td>37.03 (31.14)</td>
<td>vs DNHS (MMRM)</td>
<td></td>
</tr>
<tr>
<td>GC in DNHS</td>
<td>75 (312)</td>
<td>19.88 (21.70)</td>
<td>13.42 (18.62)</td>
<td>8.94 × 10&lt;sup&gt;−7&lt;/sup&gt; (0.23 to 0.52)</td>
<td></td>
</tr>
<tr>
<td>BMI z score</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Higher-dose vamorolone LTE&lt;sup&gt;c&lt;/sup&gt;</td>
<td>23 (65)</td>
<td>1.28 (0.51)</td>
<td>1.52 (0.66)</td>
<td>vs DNHS (MMRM)</td>
<td></td>
</tr>
<tr>
<td>GC in DNHS</td>
<td>75 (309)</td>
<td>0.65 (1.03)</td>
<td>0.79 (1.11)</td>
<td>.58 (−0.01 to 0.01)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: DNHS, Duchenne Natural History Study; GC, glucocorticoid; LTE, long-term extension; MMRM, mixed-effect model with repeated measures; NSAA, NorthStar Ambulatory Assessment; NSUK, NorthStar United Kingdom; TTCLIMB, time to climb 4 stairs; TTRW, time to run or walk 10 m; TTSTAND, time to stand from supine position; 6MWT, 6-minute walk test distance.

<sup>a</sup> Baseline values after approximately 6 months of continual treatment with vamorolone or GC.

<sup>b</sup> Last visit of participants is used. One participant in vamorolone LTE had 1 measurement and was included in baseline comparator and end of follow-up summaries.

<sup>c</sup> Participants assigned to higher-dose vamorolone (ie, 2.0 and 6.0 mg/kg/d) at start of study and maintained at 2.0 mg/kg/d or more at follow up.

<sup>d</sup> The multivariable model included a quadratic term in addition to model trajectories (ie, Duchenne muscular dystrophy natural history during the age range of interest includes periods of improvement followed by stability and deterioration); P values are provided for comparisons between comparator cohorts.

<sup>e</sup> DNHS is a natural history study with different participants followed up for different durations in the age range of comparison. Summary characteristics are also provided for participants in DNHS with 1.5 years or more of follow-up in the age range of interest.

<sup>f</sup> Owing to NorthStar UK Network participant-level data sharing restrictions, NSAA data was analyzed separately by UK Network researchers and summaries were shared for comparison with the vamorolone LTE cohort. NSAA data had a possible range from 0 to 34.

<sup>g</sup> Owing to the use of independent t test for change in NSAA over 2 years, the baseline mean is provided for 49 participants with 2 years of follow-up.
not beyond 10 seconds at baseline (median age, >9.31 years; 95% CI, 7.51 to 8.29 years) and 74 participants in the DNHS group not beyond 10 seconds at baseline (median age, >9.31 years; 95% CI, 7.51 to 10.09 years) (asymptotic log-rank 2-sample test \( P = .74 \)).

Change in mean total NSAA score for 18 participants receiving higher-dose vamorolone with NSAA data (−0.61; 95% CI, −4.65 to 3.43) vs 49 participants in NSUK receiving GC with NSAA data (−0.39; 95% CI, −1.91 to 1.13) at 2-years of follow-up was not statistically significantly different (0.22 units; 95% CI, −4.48 to 4.04; \( P = .92 \)) (Table 2). Additionally, body mass index z score change (0.002 SD/mo; 95% CI, −0.006 to 0.010; \( P = .58 \)) was not significantly different between the higher-dose vamorolone group vs the DNHS group. The NPMLE estimate of time to reach a TTSTAND milestone

Figure 3. Outcome Comparisons of Vamorolone Long-term Extension (LTE) and Duchenne Natural History (DNHS) Study Cohorts

A Time to stand velocity

B Time to run or walk 10 m

C Time to climb velocity

D BMI

E Height percentile

BMI indicates body mass index. Trajectories based on model estimates, along with 95% CIs, are plotted for an individual with fixed baseline values.
of 10 seconds or more for 108 participants in NSUK who had not experienced this event by baseline (median age, 9.55 years; 95% CI, 8.87 to 10.09 years) was similar to that for participants in DNHS and participants in the LTE study receiving higher doses. Additionally, a relative risk analysis of NSAA scores showed no significant difference in the risk of losing a motor function (ie, change from a function score of 1 or 2 at baseline to a 0 at the end of follow-up) between participants in NSUK and participants in the LTE study receiving higher doses (139 of 1734 functions lost [8.0%] vs 26 of 335 functions lost [7.8%]; relative risk, 1.033 [95% CI, 0.691 to 1.544]).

Safety Analysis of LTE Cohort
During the 24-month LTE treatment period, all 46 participants experienced at least 1 treatment-emergent AE (eTable 4 in Supplement 1). Among 41 participants receiving vamorolone at 6.0 mg/kg/d, 10 participants (24.4%) deescalated to 2.0 mg/kg/d owing to a treatment-emergent AE of weight gain. The AE abated among 6 participants after dose reduction. Among the 46 LTE participants, 6 participants (13.0%) were observed to have a total of 7 clinical fracture events according to local site AE reporting, including 1 participant with a vertebral fracture and a foot fracture on 2 separate occasions, 3 participants with an upper limb fracture, 1 participant with a vertebral compression fracture, and 1 participant with multiple vertebral fractures. There were 2 serious treatment-emergent AEs: moderate pneumonia in 1 participant and severe myoglobinuria, which occurred twice in 1 participant; 1 participant withdrew from the study owing to moderate muscle weakness. These treatment-emergent AEs were considered by the investigators to not be associated with the study drug.

Mean changes in hematology, chemistry, liver function, lipid profile, urinalysis, and electrocardiogram parameters were minimal and clinically unremarkable. There was no significant difference in changes to mean BMI z scores between participants receiving higher-dose vamorolone and those from DNHS (Figure 3D). In comparison of participants receiving higher-dose vamorolone with participants from DNHS treated with GC, there was a statistically significant difference in change in mean height percentile change (0.37 percentile/mo; 95% CI, 0.23 to 0.52 percentile/mo), with no evidence of growth deceleration in the vamorolone group (Figure 3E; eFigure 3 in Supplement 1). Among 41 participants in LTE who finished the 2-year LTE period, 39 participants (95.1%) had a hand and wrist x-ray at month 24; the mean bone age to chronologic age difference was −1.1 years (95% CI, −1.5 to −0.7 years). Results on serum biomarkers of safety including insulin resistance, adrenal suppression, and bone turnover will be presented in a separate report.

Discussion
The results of this nonrandomized clinical trial extend the findings of previous reports on open-label vamorolone treatment among boys with DMD to a total treatment duration of 30 months.25-27 We found that vamorolone treatment was not associated with a change in TTSTAND velocity from baseline to 30 months among boys with DMD aged 4 to 7 years at enrollment. However, based on clinical and laboratory results, long-term vamorolone treatment at doses up to 6.0 mg/kg/d appeared to be safe and well-tolerated. While there was an overall decline after initial improvement in TTSTAND velocity observed among boys treated with vamorolone over 2 years, vamorolone showed similar treatment efficacy as GC in DMD,48 and the decline was slower compared with that of untreated patients.26,27 Moreover, participants receiving higher doses (ie, ≥2 mg/kg/d) of vamorolone had persistent improvement in motor function as measured by TTCLIMB, TTRW, NSAA, and 6MWT distance over a 30-month treatment period.

Vamorolone treatment was associated with fewer adverse outcomes compared with GC therapy. Continuous treatment with vamorolone for up to 30 months was not associated with linear growth deceleration, whereas growth delay is a well-recognized adverse outcome associated with traditional GC, which can be distressing to boys with DMD as they mature through adolescence.49-51 We observed that boys treated with vamorolone had minimal bone age delay relative to
chronological age. In this setting of improved growth velocity with vamorolone, the slight bone age delay is anticipated to be favorable for these patients to attain increased adult height associated with greater residual growth potential. Although some participants receiving high-dose vamorolone had dose reduction to address undesired weight gain, the change in mean BMI z score was not significantly different between participants receiving vamorolone compared with participants in DNHS receiving GC. To date, pharmacodynamic biomarker data suggest that vamorolone is not associated with short-term detrimental bone outcomes. Additionally, vamorolone does not have the same association with insulin resistance as classic GC therapy; however, long-term vamorolone treatment may be associated with adrenal suppression.

Limitations
This study has several limitations. The open-label study design and use of prospectively collected real-world observational control data sets with different GCs (ie, prednisone or deflazacort), dose, and regimen (ie, daily or intermittent as per clinical practice) require cautious interpretation of the study results. The comparator data sets included patients with DMD who had been on GC for a variable amount of time before the baseline comparator visits. There was also considerable attrition in the data of participants in comparator groups over time (eg, 91.3% of participants in LTE vs 40.0% of participants in DNHS had >18 months of follow-up data for TTSTAND velocity). Despite strict subset enrollment criteria and adjustment for baseline characteristics, confounding is still possible. We could not look for association between height and motor function among patients treated with vamorolone and GC owing to the nonrandomized study design and small sample size; the risk of adrenal insufficiency will require additional studies.

Conclusions
We found that vamorolone, a novel dissociative steroidal anti-inflammatory drug, was not associated with a change in TTSTAND velocity from baseline to 30 months among boys with DMD aged 4 to 7 at enrollment. However, vamorolone was associated with similar efficacy and maintenance of muscle function comparable to historical cohorts of patients with DMD treated with GC after treatment for 30 months. Vamorolone appeared to be well-tolerated, with fewer safety concerns that are typically seen with standard-of-care GC treatment for ambulatory patients with DMD. A randomized double-blind vamorolone study (VBPI5-004; NCT03439670) is near completion; results from this study will provide class I evidence for vamorolone efficacy and safety.
Author Contributions: Dr Dang 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.

Concept and design: Clemens, Guglieri, Finkel, McDonald, Schwartz, Mengle-Gaw, Leinonen, Hoffman, Dang.

Acquisition, analysis, or interpretation of data: All authors.

Drafting of the manuscript: Mah, McDonald, Leinonen, Hoffman, Dang.

Critical revision of the manuscript for important intellectual content: Mah, Clemens, Guglieri, Smith, Finkel, Tulinius, Nevo, Ryan, Webster, Castro, Kunz, Damsker, Schwartz, Mengle-Gaw, Jackowski, Stimpson, Ridout, Ayyar-Gupta, Baranello, Manzur, Muntoni, Gordish-Dressman, Leinonen, Ward, Hoffman, Dang.


Obtained funding: Clemens, Guglieri, Hoffman.

Administrative, technical, or material support: Mah, Clemens, Finkel, Ryan, McDonald, Damsker, Schwartz, Mengle-Gaw, Jackowski, Ayyar-Gupta, Manzur, Ward, Hoffman.

Supervision: Mah, Clemens, Tulinius, Ryan, Castro, McDonald, Ridout, Baranello, Hoffman.

Conflict of Interest Disclosures: Dr Mah reported receiving grants from ReveraGen BioPharma during the conduct of the study; grants from Pfizer, Ifalfarmaco, Sarepta, Catabasis, Roche, Biogen, Novartis, NS Pharma, PTC Therapeutics, and Alberta Children’s Hospital Foundation; and personal fees from PTC Therapeutics, Biogen, and Roche outside the submitted work. Dr Clemens reported serving as a board member for Therapeutic Research in Neuromuscular Disorders Solutions (TRINDS); receiving grants from the National Institutes of Health (NIH), NS Pharma, Amicus, Sanofi, Spark, and Muscular Dystrophy Association; receiving research support through a donation from the Foundation to Eradicate Duchenne to the University of Pittsburgh; and receiving personal fees from Epiprim during the conduct of the study outside the submitted work. Dr Guglieri reported receiving data management support from North Star Network during the conduct of the study; grants from the NIH, European Committee H2020, Duchenne Muscular Dystrophy UK, and Sarepta (funding through Translational Research in Europe: Assessment and Treatment of Neuromuscular Disorders); personal fees from Sarepta; and travel fees from Santhera outside the submitted work and serving as principal or chief investigator or clinical investigator for clinical trials sponsored by Pfizer, Ifalfarmaco, Summit, Santhera, Roche, and PTC Therapeutics. Dr Smith reported receiving grants for partial salary support from Reveragen BioPharma as principal investigator during the conduct of the study. Dr Finkel reported receiving grants from ReveraGen BioPharma during the conduct of the study and grants from Catabasis and Sarepta outside the submitted work. Dr Webster reported receiving research funding from Reveragen BioPharma during the conduct of the study. Dr Kunz reported receiving grants from Astellas Gene Therapies, Biogen, Genentech, Novartis, and Sarepta; personal fees from Biogen, Genentech, Novartis, and Sarepta; and data safety monitoring board services from Sarepta during the conduct of the study. Dr McDonald reported receiving grants from the NIH National Institute of Arthritis and Musculoskeletal and Skin Diseases (NIAMS), US Department of Defense, National Institute on Disability Independent Living and Rehabilitation Research, and Parent Project Muscular Dystrophy and personal fees from Astellas, BioMarin, Epirium Bio (formerly Capricor), Catabasis, Entrada Therapeutics, Avidity Therapeutics, Edgewise Therapeutics, FibroGen, Hoffmann La Roche, Marathon, Pfizer, Santhera Pharmaceuticals, Sarepta Therapeutics, and PTC Therapeutics outside the submitted work. Dr Damsker reported being an employee and shareholder of ReveraGen BioPharma. Dr Schwartz reported receiving personal fees from ReveraGen Biopharma during the conduct of the study and personal fees from RegenXBio and TRINDS outside the submitted work. Dr Mengle-Gaw reported receiving personal fees from ReveraGen BioPharma during the conduct of the study and personal fees from RegenXBio outside the submitted work. Dr Baranello reported receiving personal fees from Roche, serving as principal investigator in clinical trials for Pfizer and Reveragen, receiving grants from Sarepta via the Dubowitz Neuromuscular Centre, and serving as an investigator in a clinical trial for NS Pharma outside the submitted work. Dr Manzur reported receiving grants from NorthStar Clinical Network during the conduct of the study and serving as clinical lead for Northstar Clinical Network UK. Dr Muntoni reported receiving grants from Muscular Dystrophy UK North Star Consortium during the conduct of the study; grants from the NIH Research Great Ormond Street Hospital-ICH Biomedical Research Centre, Genethon, and Sarepta; and personal fees from Dyne Therapeutics, Pfizer, and Sarepta outside the submitted work. Dr Gordish-Dressman reported receiving personal fees from TRINDS during the conduct of the study. Dr Mah reported receiving grants from ReveraGen BioPharma during the conduct of the study; grants from Pfizer, Ifalfarmaco, Sarepta, Catabasis, Roche, Biogen, Novartis, NS Pharma, PTC Therapeutics, and Alberta Children’s Hospital Foundation; and personal fees from PTC Therapeutics, Biogen, and Roche outside the submitted work.
study and co-owning TRINDS outside the submitted work. Dr Ward reported receiving grants from ReveraGen BioPharma via the Children's Hospital of Eastern Ontario (CHEO) Research Institute during the conduct of the study and grants from Amgen, Novartis, and PTC Therapeutics via the CHEO Research Institute outside the submitted work. Dr Hoffman reported receiving salary as chief executive officer and owning stock from ReveraGen BioPharma, serving as a board member for TRINDS, receiving grants from the NIH National Institute of Neurological Disorders and Stroke during the conduct of the study; serving as vice president and owning stock for Agada BioSciences outside the submitted work; and owning a patent issued to ReveraGen BioPharma. Dr Dang reported receiving grants from the Foundation to Eradicate Duchenne and personal fees from ReveraGen BioPharma during the conduct of the study and grants from the NIH NIAMS outside the submitted work. No other disclosures were reported.

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Group Information: NorthStar UK Network and CINRG NHS Investigators are listed in Supplement 3.


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REFERENCES


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SUPPLEMENT 2.
Trial Protocol

SUPPLEMENT 3.
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SUPPLEMENT 4.
Data Sharing Statement