A 2-part, randomized, double-blind, placebocontrolled study in participants with Duchenne muscular dystrophy amenable to exon 44 skipping to evaluate the safety and efficacy of ENTR-601-44 (ELEVATE-44)

Submission date	Recruitment status Recruiting	[X] Prospectively registered		
26/11/2024		Protocol		
Registration date	Overall study status Ongoing Condition category Genetic Diseases	Statistical analysis plan		
12/05/2025		Results		
Last Edited		Individual participant data		
22/08/2025		[X] Record updated in last yea		

Plain English summary of protocol

Background and study aims

This study aims to determine if ENTR-601-44 is safe, identify any side effects, and see how well it works on Duchenne muscular dystrophy (by increasing the amount of dystrophin protein that is produced). Dystrophin helps muscles function properly. ENTR-601-44 is an investigational medication, meaning it has not been approved by the Medicines and Healthcare products Regulatory Agency (MHRA), the health authority that gives approval for new medications in the United Kingdom.

Who can participate?

This study is for ambulatory minors and adults aged 4 to 20 years old, inclusive, who were assigned male at birth, have a confirmed diagnosis of Duchenne muscular dystrophy, and have a variant of the dystrophin gene that allows ENTR-601-44 to skip exon 44.

What does the study involve?

Participants will receive either ENTR-601-44 or a placebo (a substance with no active ingredients). At the end of the 25 weeks of the study, all participants (including those who received placebo) may be able to receive ENTR-601-44 in a long-term study. All participants will be screened to confirm eligibility for participation. This involves providing biological samples (e.g., blood and urine) and undergoing additional physical procedures at study visits. Participants will have 2 muscle biopsies over the course of the study. Muscle biopsies are important because they allow researchers to compare whether there have been changes in the muscle as a result of the study drug. The study requires regular health check visits, which will be completed according to a schedule. During these visits, various tests will be conducted, including physical exams, heart rate, temperature, blood pressure, electrocardiogram, echocardiogram, and muscle

function tests.

What are the possible benefits and risks of participating? Participants may or may not benefit from this study.

Participation could help increase knowledge about DMD and the study medication. The possible benefit of receiving ENTR-601-44 for participants is dystrophin production may increase and improve muscle function, but there is no guarantee this will happen. This is an early study of ENTR-601-44 in individuals with DMD, and therefore information on its potential positive effects in people is limited.

Since this is an early study with the medication in humans, there is limited information on side effects. There have been studies done in laboratories on animals, there has been one other study on healthy men, and there have been studies with similar medicines. From this research, possible side effects might include issues with kidney function, blood clotting, blood cell count, and liver enzymes. There could also be side effects from the medication administration and muscle biopsies. However, researchers do not know all the side effects that could happen.

Where is the study run from? Entrada Therapeutics, Inc. (USA)

When is the study starting and how long is it expected to run for?

Who is funding the study?

The Sponsor, Entrada Therapeutics, Inc., is providing financial support and materials for this study. The study site is being paid by the Sponsor to do this study. Otherwise, the site staff including the study doctor have no financial ties to the Sponsor.

Who is the main contact?

Contact information

Type(s)

Public, Scientific

Contact name

Dr . Entrada Therapeutics Clinical Trials

Contact details

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Type(s)

Principal investigator

Contact name

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Additional identifiers

Clinical Trials Information System (CTIS)

2024-517584-23

Integrated Research Application System (IRAS)

1010840

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

ENTR-601-44-201, CPMS 65821

Study information

Scientific Title

A 2-part, randomized, double-blind, placebo-controlled study in participants with Duchenne muscular dystrophy amenable to exon 44 skipping with an initial multiple ascending dose part A to assess the safety, tolerability, pharmacokinetics and pharmacodynamics of ENTR-601-44, followed by Part B to evaluate the safety and efficacy of ENTR-601-44 (ELEVATE-44)

Acronym

ELEVATE-44

Study objectives

Key Objectives (Part A)

- 1. To evaluate the safety and tolerability of ENTR-601-44 in participants with Duchenne muscular dystrophy (DMD)
- 2. To characterize the pharmacokinetics of ENTR-601-44 in participants with DMD
- 3. To characterize the pharmacodynamics of ENTR-601-44 in participants with DMD
- 4. To evaluate the immune response to ENTR-601-44 in participants with DMD

Ethics approval required

Ethics approval required

Ethics approval(s)

approved 27/01/2025, South Central - Oxford A Research Ethics Committee (Ground Floor Temple Quay House 2 The Square, Bristol, BS1 6PN, United Kingdom; +44 207 104 8118; oxforda. rec@hra.nhs.uk), ref: 24/SC/0403

Study design

Interventional double blind randomized parallel group placebo controlled trial

Primary study design

Interventional

Study type(s)

Safety, Efficacy

Health condition(s) or problem(s) studied

Duchenne muscular dystrophy (DMD)

Interventions

Part A

Experimental Arm: ENTR-601-44.

- Participants will receive a fixed number of doses at one of three dose levels. One dose will be given every six weeks.
- Drug: ENTR-601-44: Given by IV infusion as specified under Participant Group/Arm.

Placebo Comparator Arm: ENTR-601-44 matching placebo

- Participants will receive a fixed number of placebo doses matched to ENTR-601-44 doses. One dose will be given every six weeks.
- Drug: ENTR-601-44 Matching Placebo: Given by IV infusion as specified under Participant Group/Arm.

Intervention Type

Drug

Phase

Phase I/II

Drug/device/biological/vaccine name(s)

ENTR-601-44

Primary outcome(s)

Safety and tolerability of ENTR-601-44 measured using incidence and severity of treatmentemergent adverse events (TEAEs); changes in vital sign measurements, clinical laboratory results, electrocardiogram (ECG) parameters, physical examination findings from baseline through End of Study visit

Key secondary outcome(s))

- 1. Plasma, muscle, and urine concentration of ENTR-601-44 and its final metabolite at timepoints as specified in the study protocol
- 2. Change from baseline in dystrophin by Western blot from muscle biopsy at End of Study
- 3. Change from baseline in dystrophin expression and localization from muscle biopsy at End of Study
- 4. Percent change from baseline in exon 44 skipping measured in muscle biopsy at End of Study
- 5. Anti-drug antibody (ADA) and anti-dystrophin antibody in serum at Baseline, End of Study, and additional timepoints as specified in the study protocol

Completion date

Eligibility

Key inclusion criteria

Principal inclusion criteria

- 1. Genetic diagnosis of DMD and confirmed pathologic variant in the dystrophin gene amenable to exon 44 skipping as reviewed by a central genetic counselor.
- 2. Assigned male at birth with clinical signs compatible with Duchenne muscular dystrophy as determined by the investigator.
- 3. Part A: 4-20 years of age, inclusive.
- 4. Ambulatory Status Part A: ambulatory with a Performance of the Upper Limb v2.0 (PUL 2.0) Entry as per protocol at Screening
- 5. Adequate muscle for obtaining tissue biopsy as assessed by the investigator.
- 6. Other protocol-defined criteria apply.

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Mixed

Lower age limit

4 years

Upper age limit

20 years

Sex

Male

Key exclusion criteria

Principal exclusion criteria

1. Any significant concomitant medical condition that might interfere with the ability to comply with protocol

requirements.

- 2. Has an acute illness within 4 weeks prior to the first dose of study drug which may interfere with study measurements or jeopardize participant's safety.
- 3. Use of the following medications:
- 3.1. Prior treatment with any exon skipping therapy at any time
- 3.2. Prior treatment with any gene therapy at any time
- 3.3. Use of anti-coagulants, anti-thrombotics, or anti-platelet agents from at least 30 days prior to the start of the screening period until the end of the study
- 3.4. Use of an immunosuppressant for a non-DMD condition from 30 days prior to screening until the end of the study
- 3.5. Has taken or is currently taking a histone deacetylase (HDAC) inhibitor, including (but not limited to) givinostat from at least 30 days prior to the start of the screening period until the

end of the study

- 4. Laboratory abnormalities.
- 5. Daytime ventilator dependence or any use of invasive mechanical ventilation via tracheostomy.
- 6. Has an abnormal electrocardiogram (ECG) reading assessed as clinically significant by the investigator, and/or a QT interval with Fridericia correction method (QTcF) >450 msec at Screening or prior to the first dose of study drug on Day 1.
- 7. Received any experimental or investigational drug, etc. within 3 months prior to first dose or within 5 half-lives (whichever is longer).
- 8. Other protocol-defined criteria apply.

Date of first enrolment 03/07/2025

Date of final enrolment 18/08/2028

Locations

Countries of recruitment United Kingdom

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Italy

Belgium

Spain

Study participating centre UZ Leuven

Belgium

Study participating centre University Hospital Gent Belgium

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Study participating centre Centre Hospitalier Régional de la Citadelle Belgium

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Study participating centre IRCCS Ospedale San Raffaele Italy

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Study participating centre Fondazione Serena Onlus - Centro Clinico NeMO Milano Italy

Study participating centre

Fondazione Policlinico Universitario A. Gemelli IRCCS - Universita Cattolica del Sacro Cuore Italy

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Study participating centre Ospedale Pediatrico Bambino Gesu Italy

Study participating centre Hospital Sant Joan de DeuSpain

Study participating centre Hospital Universitario Vall d'HebronSpain

Study participating centre
Great Ormond Street Hospital for Children
Great Ormond Street
London
United Kingdom
WC1N 3JH

Study participating centre Freeman Hospital

Freeman Road High Heaton Newcastle upon Tyne United Kingdom NE7 7DN

Study participating centre Alder Hey Children's NHS Foundation Trust

Alder Hey Hospital
Eaton Road
West Derby
Liverpool
United Kingdom
L12 2AP

Study participating centre Leeds General Infirmary

Great George Street Leeds United Kingdom LS1 3EX

Study participating centre Royal Manchester Childrens Hospital

Hospital Road Pendlebury Swinton Manchester United Kingdom M27 4HA

Study participating centre

Oxford University Hospitals NHS Foundation Trust

Dep of Paediatrics Level 2 Children's Hospital, John Radcliffe, Headley Way, Headington Oxford United Kingdom OX3 9DU

Sponsor information

Organisation

Entrada Therapeutics, Inc.

Funder(s)

Funder type

Industry

Funder Name

Entrada Therapeutics, Inc.

Results and Publications

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are not expected to be made available due to due to the data's high commercial sensitivity.

IPD sharing plan summary

Not expected to be made available, Data sharing statement to be made available at a later date

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes
Study website	Study website	11/11/2025	11/11/2025	No	Yes