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Study of Nusinersen (BIIB058) in Participants With Spinal Muscular Atrophy

23/08/2025 08:08:25

Main Information	
Primary registry identifying number	Protocol number
LBCTR2021064778	232SM203
MOH registration number	
-	
Study registered at the country of origin	Study registered at the country of origin: Specify
Yes	
Type of registration	Type of registration: Justify
Prospective	N/A
Date of registration in national regulatory agency	
Primary sponsor	Primary sponsor: Country of origin
Biogen Idec Research Limited	United Kingdom
Date of registration in primary registry	Date of registration in national regulatory agency
18/09/2021	
Public title	Acronym
Study of Nusinersen (BIIB058) in Participants With Spinal Muscular Atrophy	
Scientific title	Acronym
Escalating Dose and Randomized, Controlled Study of Nusinersen (BIIB058) in Participants With Spinal Muscular Atrophy	
Brief summary of the study: English	
The primary objectives of this study are to examine the clinical efficacy of nusinersen administered intrathecally at higher doses to participants with SMA, as measured by the change in Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP INTEND) total score (Part B); to examine the safety and tolerability of nusinersen administered intrathecally at higher doses to participants with spinal muscular atrophy (SMA) (Parts A and C). The secondary objectives of this study are to examine the clinical efficacy of nusinersen administered intrathecally at higher doses to participants with SMA (Parts A, B, and C); to examine the effect of nusinersen administered intrathecally at higher doses to participants with SMA (Parts A and C); to examine the safety and tolerability of nusinersen administered intrathecally at higher doses to participants with SMA (Parts A and C); to examine the safety and tolerability of nusinersen administered intrathecally at higher doses to participants with SMA, to examine the effect of nusinersen administered intrathecally at higher doses to participants with SMA, to examine the effect of nusinersen administered intrathecally at higher doses to participants with SMA (Parts A and C); to examine the currently approved dose in participants with SMA (Part B).	

Brief summary of the study: Arabic

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المعطى داخل القراب بجر عات أعلى للمشاركين الذين nusinersen تتمثل الأهداف الأساسية لهذه الدراسة في فحص الفعالية السريرية لـ يعانون من ضمور العضلات الشوكي ، كما تم قياسه بالتغيير في اختبار الأطفال للاضطر ابات العصبية العضلية في مستشفى فيلادلفيا للأطفال المعطى داخل القراب بجر عات أعلى للمشاركين المصابين بضمور nusinersen (الجزء ب)، لفحص سلامة وتحمل (CHOP INTENC) المعطى داخل القراب بجر عات أعلى للمشاركين المصابين بمنمور nusinersen تتمثل الأهداف الثانويو في هذا المرابي العضلية في المعطى داخل القراب بجر عات أعلى للمشاركين المصابين منمور nusinersen الجزء ب)، فقحص سلامة وتحمل (SMA) العضليت

المُعطى داخل القراب بجر عات أعلى للمشاركين المصابين nusinersen تتمثل الأهداف الثانوية لهذه الدراسة في فحص الفعالية السريرية لـ المعطى داخل القراب بجر عات أعلى للمشاركين المصابين nusinersen ؛ لفحص تأثير (C و B و A الأجزاء) بضمور العضلات الشوكي المعطى داخل القراب بجر عات أعلى للمشاركين المصابين nusinersen ؛ لفحص سلامة وتحمل (C و A الجزأين) بضمور العضلات الشوكي المعطى داخل القراب بجر عات أعلى مقارنة بالجر عة المعتمدة حاليًا في المشاركين المشاركين مع مات على المصابين الشوكي المعطى داخل القراب بجر عات أعلى مقارنة بالجر عة المعتمدة حاليًا في المشاركين المشاركين مع الفراب المعضلات الشوكي المصابين بضمور العضلات الشوكي (الجزء ب) المشتركون في لبنان سيشاركون في الجزء ب

Health conditions/problem studied: Specify

Spinal Muscular Atrophy

Interventions: Specify

Drug: Nusinersen Administered as specified in the treatment arm Other Name: BIIB058

Key inclusion and exclusion criteria: Inclusion criteria

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Part A. B and C:

- Genetic documentation of 5q SMA (homozygous gene deletion, mutation, or compound heterozygote)

Part A:

- Onset of clinical signs and symptoms consistent with SMA at > 6 months (> 180 days) of age (i.e., later-onset SMA)

- Age 2 to \leq 15 years, inclusive, at the time of informed consent

Part B:

- Participants with SMA symptom onset < 6 months (< 180 days) of age (infantile onset) should have age > 1 week to < 7 months (< 210 days) at the time of informed consent

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- Participants with SMA symptom onset > 6 months (> 180 days) of age (later onset):

*Age 2 to < 10 years at the time of informed consent

*Can sit independently but has never had the ability to walk independently

*HFMSE score \geq 10 and \leq 54 at Screening

Part C:

- Participants ≥ 18 years of age at Screening must be ambulatory

- Currently on nusinersen treatment at the time of Screening, with the first dose being at least 1 year prior to Screening

Key inclusion and exclusion criteria: Gender	Key inclusion and exclusion criteria: Specify gender
Both	
Key inclusion and exclusion criteria: Age minimum	Key inclusion and exclusion criteria: Age maximum

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Key inclusion and exclusion criteria: Exclusion criteria

Part A, B and C:

- Presence of an untreated or inadequately treated active infection requiring systemic antiviral or antimicrobial therapy at any time during the Screening period

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- Presence of an implanted shunt for the drainage of cerebrospinal fluid (CSF) or of an implanted central nervous system (CNS) catheter - Hospitalization for surgery, pulmonary event, or nutritional support within 2 months prior to Screening or planned within 12 months after the participant's first dose.

Part A:

- Respiratory insufficiency, defined by the medical necessity for invasive or noninvasive ventilation for > 6 hours during a 24-hour period, at Screening

- Medical necessity for a gastric feeding tube

- Treatment with an investigational drug given for the treatment of SMA, biological agent, or device within 30 days or 5 half-lives of the agent, whichever is longer, prior to Screening or anytime during the study; any prior or current treatment with any survival motor neuron-2 (SMN2)-splicing modifier or gene therapy; or prior antisense oligonucleotide treatment, or cell transplantation.

Part B:

- Treatment with an investigational drug given for the treatment of SMA, biological agent, or device within 30 days or 5 half-lives of the agent, whichever is longer, prior to Screening or anytime during the study; any prior or current treatment with any SMN2-splicing modifier or gene therapy; or prior antisense oligonucleotide treatment, or cell transplantation

- Participants with SMA symptom onset > 6 months (> 180 days) of age (later onset):

*Respiratory insufficiency, defined by the medical necessity for invasive or noninvasive ventilation for > 6 hours during a 24-hour period, at Screening

*Medical necessity for a gastric feeding tube

*Participants with SMA symptom onset ≤ 6 months (≤ 180 days) of age (infantile-onset): Signs or symptoms of SMA present at birth or within



the first week after birth. Part C. - Concurrent or previous participation and/or administration of nusinersen in another clinical study NOTE: Other protocol defined Inclusion/Exclusion criteria may apply. Type of study Interventional Type of intervention Type of intervention: Specify type Pharmaceutical N/A **Trial scope** Trial scope: Specify scope Other Study design: Allocation Study design: Masking Randomized controlled trial Blinded (masking used) Study design: Control Study phase Active 2 to 3 Study design: Purpose Study design: Specify purpose N/A Treatment Study design: Assignment Study design: Specify assignment Other Sequential IMP has market authorization IMP has market authorization: Specify Yes, Worldwide US, European countries Name of IMP Year of authorization Month of authorization 2017 Nusinersen 5 Type of IMP Others Pharmaceutical class Antisense oligonucleotide inhibitor Therapeutic indication Spinal Muscular Atrophy Therapeutic benefit Nusinersen is an antisense oligonucleotide administered intrathecally via lumbar puncture (LP); it increases survival motor neuron (SMN) protein expression and significantly improves motor function in patients with spinal muscular atrophy (SMA). Study model Study model: Explain model N/A N/A Study model: Specify model N/A **Time perspective** Time perspective: Explain time perspective N/A

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Time perspective: Specify perspective	N/A
N/A	
Target follow-up duration	Target follow-up duration: Unit
Number of groups/cohorts	
Biospecimen retention	Biospecimen description
Samples without DNA	PK, Biomarker, and Immunogenicity samples will be retained for long-term storage (25 years). Safety samples and SMA genetic samples will be analyzed and destroyed after analysis.
Target sample size	Actual enrollment target size
2	
Date of first enrollment: Type	Date of first enrollment: Date
Anticipated	01/08/2021
Date of study closure: Type	Date of study closure: Date
Anticipated	30/09/2023
Recruitment status	Recruitment status: Specify
Pending	
Date of completion	
IPD sharing statement plan	IPD sharing statement description
Yes	In accordance with Biogen's Clinical Trial Transparency and Data Sharing Policy on http://clinicalresearch.biogen.com/
Additional data LIRI	
Admin comments	
Trial status	
Approved	



Secondary Identifying Numbers			
Full name of issuing authority	Secondary identifying number		
clinicaltrials.gov	NCT04089566		
EudraCT Number	2019-002663-10		

Sources of Monetary or Material Support		
Name		
Biogen Idec Research Limited UK		

Secondary Sponsors	
Name	
N/A	

Contact for Public/Scientific Queries						
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Kamal Masri	Building S2B, Downtown Katameya, Road 90, 5th settlement, New Cairo, 11835, Cairo	Egypt	009618166 9400	kamal.masri2@ iqvia.com	IQVIA
Scientific	clinicaltrials@biogen.com -	Innovation house, 70 Norden Road, Maidenhead, Berkshire, SL6 4AY	United Kingdom	0044 1628 501000	cta.submissions @biogen.com	Biogen Idec

Centers/Hospitals Involved in the Study				
Center/Hospital name	Name of principles investigator	Principles investigator speciality	Ethical approval	
Saint Georges University Medical Center	Dr. Hicham Mansour	Pediatric Neurology	Approved	

Ethics Review				
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
Saint George Hospital University Medical Center	18/01/2021	Dr. Michel Daher	N/A	+9611441000



Countries of Recruitment
Name
United Kingdom
United States of America
France
Canada
Germany
Estonia
Italy
Republic of Korea
Poland
Greece
Hungary
Ireland
Latvia
Spain
Russian Federation
Taiwan
Lebanon

Health Conditions or Problems Studied				
Condition	Code	Keyword		
Spinal Muscular Atrophy	Spinal muscular atrophy, unspecified (G12.9)	Muscular Atrophy Muscular Atrophy, Spinal Atrophy Pathological Conditions, Anatomical Neuromuscular Manifestations Neurologic Manifestations Nervous System Diseases Spinal Cord Diseases Central Nervous System Diseases Motor Neuron Disease Neurodegenerative Diseases Neuromuscular Diseases		





Interventions			
Intervention	Description	Keyword	
Drug	Nusinersen	Nusinersen	

Primary Outcomes		
Name	Time Points	Measure
Part B Infantile-onset SMA	Baseline up to Day 183	The CHOP INTEND test is designed to evaluate the motor skills of infants with significant motor weakness. It includes 16 items (capturing neck, trunk, and proximal and distal limb strength) structured to move from easiest to hardest with the grading including gravity eliminated (lower scores) to antigravity movements (higher scores). All item scores range from 0-4.
Part A and C: Number of Participants with Adverse Events (AEs) and Serious Adverse Events (SAEs)	Screening up to Day 389	An AE is any untoward medical occurrence in a participant administered a pharmaceutical product and that does not necessarily have a causal relationship with this treatment. An AE can therefore be any unfavorable and unintended sign (including an abnormal assessment such as an abnormal laboratory value), symptom, or disease temporally associated with the use of a medicinal (investigational) product, whether or not related to the medicinal (investigational) product. An SAE is any untoward medical occurrence that at any dose results in death, in the view of the Investigator, places the participant at immediate risk of death, requires inpatient hospitalization or prolongation of existing hospitalization, results in persistent or significant disability/incapacity, results in a birth defect.
Part A and C: Number of Participants with Clinically Significant Shifts from Baseline in Clinical Laboratory Parameters	Screening up to Day 302	Clinically Significant Shifts from Baseline in Clinical Laboratory Parameters
Part A and C: Number of Participants with Clinically Significant Shifts from Baseline in Electrocardiograms (ECGs)	Screening up to Day 302	Clinically Significant Shifts from Baseline in Electrocardiograms (ECGs)
Part A and C: Number of Participants with Clinically Significant Shifts from Baseline in Vital Signs	Screening up to Day 302	Clinically Significant Shifts from Baseline in Vital Signs
Part A and C: Change from Baseline in Body Length/Height	Baseline up to Day 302	Change from Baseline in Body Length/Height
Part C Infantile-onset SMA: Change from Baseline in Head Circumference	Baseline up to Day 302	Change from Baseline in Head Circumference
Part C Infantile-onset SMA: Change from Baseline in Chest Circumference	Baseline up to Day 302	Change from Baseline in Chest Circumference
Part C Infantile-onset SMA: Change from Baseline in Arm Circumference	Baseline up to Day 302	Change from Baseline in Arm Circumference
Part A and C Later-onset SMA: Change from Baseline in Ulnar Length	Baseline up to Day 302	Change from Baseline in Ulnar Length
Part A and C: Ratio of Weight for Age	Baseline up to Day 302	Ratio of Weight for Age
Part A and C: Ratio of Weight for Length	Baseline up to Day 302	Ratio of Weight for Length
Part C: Ratio of Head-to-chest Circumference	Baseline up to Day 302	Ratio of Head-to-chest Circumference
Part A and C: Change from Baseline in Activated Partial Thromboplastin Time (aPTT)	Baseline up to Day 269	Change from Baseline in Activated Partial Thromboplastin Time (aPTT)

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Part A and C: Change from Baseline in Prothrombin Time (PT)	Baseline up to Day 269	Change from Baseline in Prothrombin Time (PT)
Part A and C: Change from Baseline in International Normalized Ratio (INR)	Baseline up to Day 269	Change from Baseline in International Normalized Ratio (INR)
Part A and C: Change in Urine Total Protein	Baseline up to Day 302	Change in Urine Total Protein
Part A and C: Change from Baseline in Neurological Examination Outcomes	Baseline up to Day 302	Change from Baseline in Neurological Examination Outcomes
Part A and C: Percentage of Participants with a Postbaseline Platelet Count Below the Lower Limit of Normal on at least 2 Consecutive Measurements	Baseline up to Day 302	Percentage of Participants with a Postbaseline Platelet Count Below the Lower Limit of Normal on at least 2 Consecutive Measurements
Part A and C: Percentage of Participants with a Postbaseline Corrected QT Interval Using Fridericia's Formula (QTcF) of > 500 millisecond (msec) and an Increase from Baseline to Any Postbaseline Timepoint in QTcF of > 60 msec	Baseline up to Day 302	Percentage of Participants with a Postbaseline Corrected QT Interval Using Fridericia's Formula (QTcF) of > 500 millisecond (msec) and an Increase from Baseline to Any Postbaseline Timepoint in QTcF of > 60 msec

Key Secondary Outcomes		
Name	Time Points	Measure
Part B Infantile-onset SMA: Percentage of Hammersmith Infant Neurological Examination (HINE) Section 2 Motor Milestone Responders	Day 302	Section 2 of the HINE is used to assess motor milestones of the participants It is composed of 8 motor milestone categories: voluntary grasp, ability to kick in supine position, head control, rolling, sitting, crawling, standing, and walking.
Part B Infantile-onset SMA: Change from Baseline in HINE Section 2 Motor Milestones Total Score	From baseline to Day 302	Section 2 of the HINE is used to assess motor milestones of the participants. It is composed of 8 motor milestone categories: voluntary grasp, ability to kick in supine position, head control, rolling, sitting, crawling, standing, and walking.
Part B Infantile-onset SMA: Time to Permanent Ventilation	Screening up to Day 302	Permanent ventilation is defined as tracheostomy or ≥ 16 hours of ventilation/day continuously for > 21 days in the absence of an acute reversible event.
Part B Infantile-onset SMA: Time to Death (Overall Survival)	Screening up to Day 399	Overall survival
Part A and B Later-onset SMA: Change from Baseline in Hammersmith Functional Motor Scale - Expanded (HFMSE) Score	Baseline up to Day 302	HFMSE scale is a tool to assess motor function in children with SMA. The original 20 item Hammersmith Functional Motor Scale was expanded to include 13 additional adapted items from the Gross Motor Function Measure to improve sensitivity for the higher functioning ambulant population.
Part A and B Later-onset SMA: Change from Baseline in Revised Upper Limb Module (RULM) Score	Baseline up to Day 302	The RULM is developed to assess upper limb functional abilities participants with SMA. This test consists of upper limb performance items that are reflective of activities of daily living.
Part A and B Later-onset SMA: Total Number of New WHO Motor Milestones	Baseline up to Day 302	Total Number of New WHO Motor Milestones
Part A and B Later-onset SMA: Change from Baseline in Assessment of Caregiver Experience with Neuromuscular Disease (ACEND)	Baseline up to Day 302	ACEND is designed to quantify the caregiver impact experienced by parents/caregivers of children affected with severe neuromuscular diseases. It includes domains assessing physical impact (including feeding/grooming/dressing, sitting/play, transfers, and mobility) and general caregiver impact (including time, emotion, and finance).
Part A and B Later-onset SMA: Change from Baseline in Pediatric Quality of Life Inventory™ (PedsQL)	Baseline up to Day 302	PedsQL is used to measure healthrelated quality of life (HRQOL) in children and adolescents. The PedsQL Measurement 4.0 Generic Core Scales include assessment of physical functioning, emotional functioning, social functioning, and school functioning and the PedsQL 3.0 Neuromuscular Module measures HRQOL.

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Part B: Number of Participants with AEs and SAEs	Screening up to Day 399	An AE is any untoward medical occurrence in a participant administered a pharmaceutical product and that does not necessarily have a causal relationship with this treatment. An AE can therefore be any unfavorable and unintended sign (including an abnormal assessment such as an abnormal laboratory value), symptom, or disease temporally associated with the use of a medicinal (investigational) product, whether or not related to the medicinal (investigational) product. An SAE is any untoward medical occurrence that at any dose results in death, in the view of the Investigator, places the participant at immediate risk of death, requires inpatient hospitalization, results in persistent or significant disability/incapacity, results in a birth defect
Part B: Number of Participants with Clinically Significant Shifts from Baseline in Clinical Laboratory Parameters	Screening up to Day 302	Clinically Significant Shifts from Baseline in Clinical Laboratory Parameters
Part B: Number of Participants with Clinically Significant Shifts from Baseline in ECGs	Day 1 up to Day 302	Clinically Significant Shifts from Baseline in ECGs
Part B: Number of Participants with Clinically Significant Shifts from Baseline in Vital Signs	Screening up to Day 302	Clinically Significant Shifts from Baseline in Vital Signs
Part B: Change from Baseline in Body Length/Height	Baseline up to Day 302	Change from Baseline in Body Length/Height
Part B Infantile-onset SMA: Change from Baseline in Head Circumference	Baseline up to Day 302	Change from Baseline in Head Circumference
Part B Infantile-onset SMA: Change from Baseline in Chest Circumference	Baseline up to Day 302	Change from Baseline in Chest Circumference
Part B Infantile-onset SMA: Change from Baseline in Arm Circumference	Baseline up to Day 302	Change from Baseline in Arm Circumference
Part B Later-onset SMA: Change from Baseline in Ulnar Length	Baseline up to Day 302	Change from Baseline in Ulnar Length
Part B: Ratio of Weight for Age	Baseline up to Day 302	Ratio of Weight for Age
Part B: Ratio of Weight for Length	Baseline up to Day 302	Ratio of Weight for Length
Part B: Ratio of Head-to-chest Circumference	Baseline up to Day 302	Ratio of Head-to-chest Circumference
Part B: Change from Baseline in aPTT	Baseline up to Day 279	Change from Baseline in aPTT
Part B: Change from Baseline in PT	Baseline up to Day 279	Change from Baseline in PT
Part B: Change from Baseline in INR	Baseline up to Day 279	Change from Baseline in INR
Part B: Change in Urine Total Protein	Baseline up to Day 302	Change in Urine Total Protein
Part B: Change from Baseline in Neurological Examination Outcomes	Baseline up to Day 302	Change from Baseline in Neurological Examination Outcomes
Part B: Percentage of Participants with a Postbaseline Platelet Count Below the Lower Limit of Normal on at least 2 Consecutive Measurements	Baseline up to Day 302	Percentage of Participants with a Postbaseline Platelet Count Below the Lower Limit of Normal on at least 2 Consecutive Measurements
Part B: Percentage of Participants with a Postbaseline QTcF of > 500 msec and an Increase from Baseline to Any Postbaseline Timepoint in QTcF of > 60 msec	Baseline up to Day 302	Percentage of Participants with a Postbaseline QTcF of > 500 msec and an Increase from Baseline to Any Postbaseline Timepoint in QTcF of > 60 msec
Part A, B and C: Number of Hospitalizations	Day 1 to Day 279	Number of Hospitalizations
Part A, B and C: Duration of Hospitalizations	Day 1 to Day 279	Duration of Hospitalizations

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Part A, B and C: Clinical Global Impression of Change (CGIC)	Day 302	The CGIC scale is a 7-point scale that requires the clinician to assess how much the participant's illness has changed relative to a baseline state at the beginning of the intervention, where 1=very much improved, 2=much improved, 3=minimally improved, 4=no change, 5=minimally worse, 6=much worse, 7=very much worse. Higher rating will indicate worsening of the condition.
Part A, B and C: Number of Participants with Serious Respiratory Events	Screening up to Day 399	Number of Participants with Serious Respiratory Events
Part B Infantile-onset SMA: Percentage of Time on Ventilation	Screening up to Day 302	Percentage of Time on Ventilation
Parts A, B and C: Ventilator Use	Screening up to Day 302	Ventilator Use
Part B Infantile-onset SMA: Change from Baseline in the Parent Assessment of Swallowing Ability (PASA) Scale]	Baseline up to Day 302	PASA questionnaire is developed to assess the signs and symptoms of dysphagia. It includes 33 items across 4 domains. General feeding, drinking liquids and eating solid foods will be assessed with 5 levels of response (Never, Rarely, Sometimes, Often, and Always), and 2 items will be assessed with 'Yes'/'No'. The assessment of swallowing concerns has 4 levels of response: Strongly Agree, Agree, Disagree, and Strongly Disagree.
Part C: Change from Baseline in HFMSE Score	Baseline up to Day 302	HFMSE scale is a tool to assess motor function in children with SMA. The original 20 item Hammersmith Functional Motor Scale was expanded to include 13 additional adapted items from the Gross Motor Function Measure to improve sensitivity for the higher functioning ambulant population
Part C: Change from Baseline in HFMSE Score	Baseline up to Day 302	HFMSE scale is a tool to assess motor function in children with SMA. The original 20 item Hammersmith Functional Motor Scale was expanded to include 13 additional adapted items from the Gross Motor Function Measure to improve sensitivity for the higher functioning ambulant population.
Part C: Change from Baseline in RULM Score	Baseline up to Day 302	The RULM is developed to assess upper limb functional abilities participants with SMA. This test consists of upper limb performance items that are reflective of activities of daily living.
Part C: Total Number of New WHO Motor Milestones	Baseline up to Day 302	Total Number of New WHO Motor Milestones
Part C: Change from Baseline in ACEND	Baseline up to Day 302	ACEND is designed to quantify the caregiver impact experienced by parents/caregivers of children affected with severe neuromuscular diseases. It includes domains assessing physical impact (including feeding/grooming/dressing, sitting/play, transfers, and mobility) and general caregiver impact (including time, emotion, and finance).
Part C: Change from Baseline in PedsQL™	Baseline up to Day 302	PedsQL is used to measure healthrelated quality of life (HRQOL) in children and adolescents. The PedsQL Measurement 4.0 Generic Core Scales include assessment of physical functioning, emotional functioning, social functioning, and school functioning and the PedsQL 3.0 Neuromuscular Module measures HRQOL.
Part C: Change from Baseline in CHOP INTEND Total Score	Baseline up to Day 302	The CHOP INTEND test is designed to evaluate the motor skills of infants with significant motor weakness. It includes 16 items (capturing neck, trunk, and proximal and distal limb strength) structured to move from easiest to hardest with the grading including gravity eliminated (lower scores) to antigravity movements (higher scores). All item scores range from 0-4.
Part C: Change from Baseline in HINE Section 2 Motor Milestones Total Score	Baseline up to Day 302	Section 2 of the HINE is used to assess motor milestones of the participants. It is composed of 8 motor milestone categories: voluntary grasp, ability to kick in supine position, head control, rolling, sitting, crawling, standing, and walking.





Parts A and B Later-onset SMA: Change from Baseline in the PASA Scale	Baseline up to Day 302	PASA questionnaire is developed to assess the signs and symptoms of dysphagia. It includes 33 items across 4 domains. General feeding, drinking liquids and eating solid foods will be assessed with 5 levels of response (Never, Rarely, Sometimes, Often, and Always), and 2 items will be assessed with 'Yes'/'No'. The assessment of swallowing concerns has 4 levels of response: Strongly Agree, Agree, Disagree, and Strongly Disagree.
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Trial Results	
Summary results	
Study results globally	
Date of posting of results summaries	Date of first journal publication of results
Results URL link	
Baseline characteristics	
Participant flow	
Adverse events	
Outcome measures	
URL to protocol files	