International Consensus on Clinical Severity Scale Use in Evaluating Niemann-Pick Disease Type C in Paediatric and Adult Patients: Results From a Delphi Study

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Research

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Abstract

**Background:** Several scales have been developed in the past two decades to evaluate Niemann-Pick disease Type C (NPC) severity in clinical practice and trials. However, a lack of clarity concerning which scale to use in each setting is preventing the use of standardised assessments across the world, resulting in incomparable data sets and clinical trial outcome measures. This study aimed to establish agreed approaches for the use of NPC severity scales in clinical practice and research.

**Methods:** A Delphi method of consensus development was used, comprising three survey rounds. In Round 1, participants were asked nine multiple-choice and open-ended questions to gather opinions on the six severity scales and domains. In Rounds 2 and 3, questions aimed to gain consensus on the opinions revealed in Round 1 using a typical Likert scale.

**Results:** Nineteen experts, active in NPC paediatric and adult research and treatment, participated in this study. Of these, 16/19 completed Rounds 1 and 2 and 19/19 completed Round 3. Consensus (defined as ≥70% agreement or neutrality) was achieved for 66.7% of the multiple-choice questions in Round 2 and 83% of the multiple-choice questions in Round 3. Consensus was almost reached (68%) on the use of the 5-domain NPCCSS scale as the first choice in clinical practice. Consensus was reached (74%) for the 17-domain NPCCSS scale as the first choice in clinical trial settings, but the domains measured in the 5-domain scale should be prioritised as the primary endpoints. Experts called for educational and training materials on how to apply the NPCCSS (17- and 5-domains) for clinicians working in NPC.

**Conclusions:** In achieving a consensus on the use of the 17-domain NPCCSS scale as the first choice for assessing clinical severity of NPC in clinical trial settings, but prioritising the domains in the 5-domain NPCCSS scale for routine clinical practice, this study can help to inform future discussion around the use of the existing NPC clinical severity scales. For routine clinical practice, the study helps provide clarity on which scale is favoured by a significant proportion of a representative body of experts, in this case, the 5-domain NPCCSS scale.

Introduction

Niemann-Pick disease Type C (NPC) is a devastating, rare neurodegenerative disease characterised by a defect that severely impedes cellular lipid trafficking [1]. Inherited in an autosomal recessive manner, individuals with NPC have mutations in one of two genes, NPC1 or NPC2. Approximately 95% of affected individuals have mutations in NPC1 [1]. As a result, cholesterol and sphingolipids accumulate within the endosomal/lysosomal system, degrading the central nervous system (CNS) and causing a diverse number of neurological symptoms depending on the patient's age at onset. These symptoms may include cerebellar ataxia, dysarthria, dysphagia, cataplexy, seizures, dystonia, vertical gaze palsy, progressive dementia and death by 8–25 years of age [2].

The exact prevalence of NPC disease is difficult to calculate due to inadequate clinical awareness as well as the relative complexity of biochemical testing. However, it has been estimated to be 1 case per 100,000 live births [3]. The severe disabilities caused by NPC, particularly during the later stages of the disease, affect a patient's entire family and optimal disease management requires highly specialised healthcare within a multidisciplinary care setting.

Although NPC is not yet curable, knowledge on its pathogenesis has increased several-fold since the characterisation of the NPC1 and NPC2 genes. The focus of therapy remains symptom management, while advances are made in identifying effective disease-modifying treatments and investigational therapies.

The goal of the research into potential treatments for NPC is to develop drugs that are safe, effective and accessible to all members of the community. However, because NPC is an ultra-rare disease with considerable variability, designing and defining clinical trial inclusion criteria and endpoints can be challenging. Following a series of multidisciplinary discussions that culminated in an interactive workshop held at the Niemann Pick UK Annual Conference in 2019, it was agreed that there was a need to develop a consensus on the use of existing NPC clinical severity scales in routine clinical practice and clinical trials. By determining such consensus, assessments across the world could be standardised to establish comparable data sets and demonstrate treatment efficacy through meaningful outcome measures.

Several scales have been developed and published over the past two decades but, essentially, all are based on a four-domain scale initially developed by Iturriaga et al (2006) [4] (see Table 1). The present study aimed to establish consensus on the use of the clinical NPC severity scales listed in Table 1 in three different settings: routine clinical practice, clinical trial enrolment and clinical trial assessment. The list was agreed upon by the Core Working Group and informed by a comprehensive literature and clinical trial review. A Delphi method of consensus development was used to integrate anonymised perspectives from a group of international clinical experts with expertise in treating both paediatric and adult NPC patients and utilising scales to determine NPC severity. The Delphi method has proven to be a reliable measurement instrument to derive the opinion of a group of experts and evaluate the extent of agreement and to resolve any disagreement on a topic [5]. It has been widely used to establish a consensus across a range of subject areas. The study was coordinated as an iterative process of three surveys, with the questions in each round based on the previous round's results.
The objectives of this study were to build consensus among international experts in the field of NPC on: (i) the preferred clinical scale(s) for assessing NPC severity (ii) the most suitable NPC severity scale to be used within each of the following three settings: routine clinical practice, clinical trial enrolment and clinical trial assessment.

**Methods**

**Study Design**

The Delphi technique is a reliable measurement instrument for developing novel concepts and setting the course of future-orientated research [6]. It assesses the opinion of a group of experts to gauge their levels of agreement and to resolve disagreement on an issue [5] and has been used successfully across a range of subject areas to gain a clinical consensus [7, 8, 9]. A Delphi study was carried out to gain a clinical consensus on six existing NPC clinical severity scales (see Table 1) that can be used within the following three settings: routine clinical practice, clinical trial inclusion criteria and clinical trial endpoints. These settings were identified via a literature review, conducted prior to the study, of how the six severity scales have been used in clinical practice and trials to date. Nineteen experts, active in NPC paediatric and adult research and treatment, participated in this study.

The Delphi technique is an iterative process that comprised three rounds. Ahead of the first round of this Delphi study, participants received two documents: 1) Literature review summary findings and 2) Clinical trials summary findings (see Appendices). Round 1 aimed to gather opinions on the use of the six severity scales and the key domains that should be measured in each clinical setting. Round 2 and 3 strived to gain consensus on these opinions. Ahead of Round 2, participants received the summary of the opinions revealed in Round 1, but the anonymity of each participant was ensured. This is an important consideration in Delphi studies to allow individuals to express their opinions freely and openly. However, the results of Round 2 were not shared ahead of Round 3 to avoid influencing the response.

**Round 1**

In Round 1, 16 specialists took part in a nine-question survey. Each of the nine questions constituted two parts: (a) a multiple-choice question and (b) a free-text question, that asked for reasoning, further insight or a recommendation based on their answer to part (a). The first round aimed to gather opinions on the six severity scales and domains that should be assessed in routine clinical practice, clinical trial inclusion criteria and clinical trial endpoints.

**Round 2**

In Round 2, 16 specialists, 11 of whom took part in Round 1, participated in an eleven-question survey. Participants were asked to independently rank nine statements using a 5-point Likert scale (‘strongly agree’, ‘agree’, ‘neither agree nor disagree’, ‘disagree’, ‘strongly disagree’). The final two questions of the survey were free-text questions about the NPC severity scales. Consensus was determined as agreement, or neutrality, by greater than or equal to 70% of the participants.

**Round 3**

In Round 3, 19 experts took part in a six-question survey, which used the same 5-point Likert scale as in Round 2. The aim of this final round was to gain consensus on what should be recommended based on opinions from Rounds 1 and 2. Consensus was defined in the same way as in Round 2.

Three survey rounds are considered optimal when trying to reach consensus [10]. They also allow the free-text question responses in Rounds 1 and 2 to be incorporated into Rounds 2 and 3, respectively. All surveys were administered using SurveyMonkey and survey links were distributed via email.

### Table 1

| Scale name                                                                 | List of domains measured                                                                 |
|---------------------------------------------------------------------------|------------------------------------------------------------------------------------------|
| 17-domain NPC Clinical Severity Score (NPCCSS) [18]                       | The NPCCSS measures 17-domains:                                                          |
| - Nine major domains: ambulation, cognition, eye movement, fine motor, hearing, memory, seizures, speech, swallowing. |                                                                                                |
| - Eight minor domains: auditory brainstem response, behaviour, gelastic cataplexy, hyperreflexia, incontinence, narcolepsy, psychiatric, respiratory problems. |                                                                                                |
| 5-domain NPCCSS [16]                                                      | Based on the 17-domain NPCCSS, the 5-domain NPCCSS measures ambulation, cognition, fine motor, speech and swallowing (five domains selected by NPC individuals, their caregivers and NPC experts as the most clinically relevant). |
| Disability Scale (NPC-specific) [4]                                      | It measures four domains: ambulation, manipulation, language and swallowing, with scores 1–4 or 5. |
| Disease-specific Disability Scale [19]                                   | Adaption of the scale developed by Iturriaga et al (2006) [4]. It measures four domains: ambulation, manipulation, language and swallowing, with weighted scores for each parameter on a scale from 0–1. |
| NPC-cdb Scale [20]                                                       | Unlike previous scales, the NPC-cdb scale represents the sum of all past and current symptoms present in a patient at any given time, with each symptom contributing a severity-weighted summand. |
| Functional Disability Scale [3]                                         | Modified from Pineda et al (2009) [19]. It measures seven domains: ambulation, manipulation, language, swallowing, eye movements, seizure and neurocognitive development (for patients under 12 years of age). |
Consensus was defined as greater than or equal to 70% of participants strongly agreeing/agreeing/neutrality on the Likert scale questions in Rounds 2 and 3. This level of agreement has been considered sufficient in several previous Delphi studies [11, 12].

**Survey Development**

The initial survey development involved the definition of a research question and development of the questions to be used in Round 1, based on the study team's expertise and a review of the literature. This initial development was carried out by the Core Working Group. To meet the study objectives, the survey was split into three sections. The first round included questions to establish opinions on the most useful NPC severity scales and domains measured in each clinical setting and the second and third round aimed to gain consensus on the opinions gathered in Round 1.

**Expert Panel Recruitment**

In Delphi studies, a minimum of 12 participants is considered sufficient for achieving a consensus, as larger sample sizes can deliver diminishing returns concerning the validity of the findings [13, 14, 15]. Twenty international specialists from Europe, the United States and South America were invited to complete the Delphi study, of which 19 agreed to participate. The professional community in NPC is very small, given the rarity of the disease, so the authors of the existing clinical severity scales that are still practising as NPC clinicians were also invited to take part. The participants were identified by Dr Will Evans, Chair of NPUK, and ratified by the Core Working Group as the key specialists in NPC around the world and invited via email to participate in this Delphi study.

**Results**

**Participants**

Each survey round of this Delphi study comprised a representative panel of clinical experts (the Expert Panel) treating both paediatric and adult NPC patients, from seven different countries: United States of America (n = 6), United Kingdom (n = 5), Germany (n = 3), Spain (n = 2), Brazil (n = 1), France (n = 1) and Australia (n = 1). More than half (58%) of the study participants included in the study were paediatric specialists, which was expected given that the condition is more prevalent in babies and young children.

**Round 1**

In Round 1, consensus was reached amongst the 16 international experts on the five most important domains to be measured to assess NPC clinical severity in the context of all three clinical settings (routine clinical practice, trial enrolment and clinical trial outcome measures). These included: ambulation, cognition, fine motor, speech and swallowing. Although these are the five domains captured in the 5-domain NPCCSS scale, the group was far from unanimous in the ambition to use a single scale across each of the clinical settings. Nonetheless, the 5-domain and 17-domain NPCCSS were among the highest-ranked for preferred use within all three settings, with an average consensus of 43.75% and 31.25% respectively. The most divisive question of the survey was regarding the adoption of a single severity scale in all scenarios, with some responses supportive of the consistency and optimisation of a scale on a global scale while others suggested that a single scale would be too reductive. Based on Round 1 results, detailed in Table 2, the second round focused on questions that asked participants to rate statements according to a typical Likert scale.
Table 2
Responses to statement included in Round 1

| 1a. Which of the following NPC Severity Scales is the most useful as a practical measure of disease severity in normal clinical practice? | 17-domain NPCCSS[18] | 5-domain NPCCSS[16] | Disability scale[4] | Disease-specific disability scale[19] | NPC-cdb scale[20] | Functional disability scale[3] | None | Other |
|---|---|---|---|---|---|---|---|---|
| | 18.75% | 43.75% | 12.5% | 18.75% | 0% | 18.75% | 0% | 6.25% |

1b. Please explain the reason for your answer

Summary of key insights:

- The 5-domain NPCCSS and Disease-specific disability scale were highlighted by multiple respondents as being simple, quick to administer and complete in any clinical environment in a routine clinical exam, with no additional work, tools or expertise required.
- The increased validity of the 17- and 5-domain NPCCSS scales, given their recent use by multiple groups for large cohorts of NPC patients and in clinical trials, was cited in further support of their use.
- While the time-effectiveness and accuracy of the 5-domain NPCCSS scale in clinical practice were acknowledged, its limitations were also flagged in terms of evaluation of certain subsets of patients, e.g. those with mainly psychiatric involvement or experiencing seizures.
- The granularity of scores and the comprehensiveness of the 17-domain NPCCSS scale was appreciated by multiple respondents.
- Notably, the accuracy of the description of eye movement impairment was questioned across all of the scales.
- The challenges of capturing progression in late-onset patients with more slowly-progressing disease when using these scales was also raised, with the suggestion for greater granularity of scoring across domains.

2a. In the context of routine clinical practice, if you had to limit measurements to only 5 of these domains, which would you select?

| Ambulation | Cognition | Eye movement | Fine motor | Hearing | Memory | Seizures | Speech | Swallowing | Other |
|---|---|---|---|---|---|---|---|---|---|
| 100% | 100% | 12.5% | 93.75% | 6.25% | 12.5% | 12.5% | 81.25% | 87.5% | 25% |

2b. Please let us know which are the minimum number of domains you feel would be sufficient to reflect disease burden and progression in everyday practice and why

| 1–4 domains | 5-domains | 7–9 domains |
|---|---|---|
| 18.75% | 43.75% | 12.5% |

3a. Which of the following NPC Severity Scales is the most useful as a measure of disease severity for enrolment, in the context of a research study or clinical trial?

| 17-domain NPCCSS[18] | 5-domain NPCCSS[16] | Disability scale[4] | Disease-specific disability scale[19] | NPC-cdb scale[20] | Functional disability scale[3] | None | Other |
|---|---|---|---|---|---|---|---|
| | 43.75% | 37.5% | 0% | 12.5% | 12.5% | 6.25% | 6.25% | 6.25% |
## 1a. Which of the following NPCS Severity Scales is the most useful as a practical measure of disease severity in normal clinical practice?

| Scale | 17-domain NPCCSS [18] | 5-domain NPCCSS [16] | Disability scale [4] | Disease-specific disability scale [19] | NPC-cdb scale [20] | Functional disability scale [3] | None | Other |
|-------|------------------------|----------------------|----------------------|----------------------------------------|--------------------|---------------------------------|------|-------|
| Score | 18.75%                 | 12.5%                | 18.75%               | 0%                                      | 18.75%             | 0%                              | 0%   | 6.25% |

### Summary of key insights:
- The 17-domain NPCSS scale was most popular among respondents in the context of clinical trial enrolment; it was seen as the most refined scale with the broadest coverage of the disease and the largest score range in each domain (5 instead of 4 or less). However, it was noted that the scale could be improved with respect to the linearity of the rating in some domains.

- Granularity was seen as critical to measuring change and baseline assessment within clinical trials; it should be as comprehensive as possible while remaining quantifiable.
- As more data becomes available, e.g. genomic data, there may be a need to reconsider which parameters are most important and whether preferred scales need to be amended accordingly.
- Simplicity was seen as valuable for multi-centre trials. The simplicity of the 5-domain NPCSS scale, as well as its proven correlation with the 17-domain scale, may drive the preference for use in some trials.
- Additionally, the question of which parameters can be expected to change in a clinical trial should be considered, as they determine both the endpoint and inclusion criteria and the identification of patients who can demonstrate measurable progression. Given the heterogeneity of the condition, general scores may not be suitable for every trial.

## 4a. In the context of trial enrolment, if you had to limit measurements to only 5 of these domains, which would you select?

| Domain   | Ambulation | Cognition | Eye movement | Fine motor | Hearing | Memory | Seizures | Speech | Swallowing | Other |
|----------|------------|-----------|--------------|------------|---------|--------|----------|--------|------------|-------|
| Score    | 100%       | 93.75%    | 25%          | 93.25%     | 0%      | 18.75% | 18.75%   | 87.5%  | 87.5%      | 6.25% |

## 4b. Please let us know which are the minimum number of domains you feel would be sufficient to reflect disease burden and progression in a clinical trial setting and why

| Number of Domains | 1–4 domains | 5-domains | 7–9 domains |
|------------------|-------------|-----------|-------------|
| Score            | 12.5%       | 43.75%    | 12.5%       |

## 5a. Do you think that the ASIS score (Annual Severity Incremental Score), is a suitable measure to capture the rate of disease progression for trial enrolment and/or clinical trial outcome measures?

| Suitability       | Yes, it is suitable for trial enrolment | Yes, it is suitable for clinical trial outcome measures | No, it is not suitable for either |
|-------------------|----------------------------------------|-------------------------------------------------------|---------------------------------|
| Score             | 68.75%                                 | 62.5%                                                 | 12.5%                           |

### Summary of key insights:
- Multiple respondents highlighted that as ASIS is a general scale and should only be a secondary outcome measure. It is not as sensitive as other scales, particularly over a potentially short period of a clinical trial.

- Broadly, its value for both prospective and retrospective measures was recognised by the majority of respondents, particularly in regard to quantifying progression in a respective age group over multiple years of treatment.
1a. Which of the following NPC Severity Scales is the most useful as a practical measure of disease severity in normal clinical practice?

| Scale | 17-domain NPCCSS [18] | 5-domain NPCCSS [16] | Disability scale [4] | Disease-specific disability scale [19] | NPC-cdb scale [20] | Functional disability scale [3] | None | Other |
|-------|-----------------------|----------------------|---------------------|-------------------------------------|-------------------|----------------------------------|------|-------|
| %      | 18.75%                | 43.75%               | 12.5%               | 10.75%                              | 0%                | 18.75%                           | 0%   | 6.25% |

6b. Please explain the reason for your answer.

Summary of key insights:

- To market an expensive drug, a sponsor will need to demonstrate a positive impact on the dynamics of a composite clinical progression score.
- The challenge of conducting an outcome trial of sufficient duration (probably > 24 mo) to see a robust statistically significant clinical effect in any of the scales with a reasonable number of participants was raised by more than one respondent.
- A severity score was seen as a suitable outcome measure if the data are collected properly and in a rigorous and consistent manner across sites and with proper (and fairly simple) training. Otherwise data are less reliable and more objective measures are needed, such as MRI, BAEPs, oxysterols, and videos with blind raters, of walking and the 9HPT, as suggested by other respondents.
- To support reproducibility and reliability across trial sites, limiting the severity score to the 5 major domains was seen as sensible. These need to be guided with precise assessments (named tests) and be age/cognition dependent.
- The 5-domain scale addresses the five most important domains, based on clinician and family opinion, and does not include items that can vary due to other treatments and thus act as confounders.

7a. Which do you think are the key domains to capture as a clinical trial outcome measure? Please select all that apply.

| Domain               | %    |
|----------------------|------|
| Ambulation           | 93.75% |
| Cognition            | 75%  |
| Eye movement         | 25%  |
| Fine motor           | 100% |
| Hearing              | 12.5% |
| Memory               | 25%  |
| Seizures             | 25%  |
| Speech               | 87.5% |
| Swallowing           | 81.25% |
| Other                | 12.5% |

7b. Please provide any further insights.

Summary of key insights:

- The top 5-domains chosen by the group were seen as the most relevant to describe neurological disease progression. However, it was suggested the impact of seizures needs to be accounted for, as well as the quality of life of the patient and their caregivers.
- Sophisticated computer assessment to measure speech in trials was suggested for consideration.
- Until an effective disease modifying therapy becomes available, deciding what to measure in clinical trials remains a challenge. The solution proposed was to start by measuring everything and adapting endpoints dependent on the findings, particularly with different age groups involved.

8a. Do you think that the adoption of a single severity scale in all scenarios is optimal, even if this means losing some refinement? If ‘yes’, which of the following NPC Severity Scales would you recommend?

| Scale | 17-domain NPCCSS [18] | 5-domain NPCCSS [16] | Disability scale [4] | Disease-specific disability scale [19] | NPC-cdb scale [20] | Functional disability scale [3] | None | Other |
|-------|-----------------------|----------------------|---------------------|-------------------------------------|-------------------|----------------------------------|------|-------|
| %      | 12.5%                 | 50%                  | 0%                  | 6.25%                              | 0%                | 12.5%                            | 25%  | 0%    |

8b. Please provide any further insights.

Summary of key insights:

- This was the most divisive question for the group with many calling for greater consistency and optimisation of a single multi-domain score system.
1a. Which of the following NPC Severity Scales is the most useful as a practical measure of disease severity in normal clinical practice?

- 17-domain NPCSS [18]
- 5-domain NPCSS [16]
- Disability scale [4]
- Disease specific disability scale [19]
- NPC-cdb scale [20]
- Functional disability scale [3]
- None
- Other

- In the absence of a proven composite score that can work in all settings, the use of different scales in clinical trials should be at the liberty of each investigator/sponsor.
- Neither clinical research nor clinical practice should be compromised by a one size fits all approach. This would be regression to the least common denominator.
- Alternatively, it may be appropriate to consider that if a scale cannot be implemented in routine clinical practice, it is not justifiable to use in a trial.
- It is critically important to try to standardize scoring and implementation to make datasets comparable.
- The 5-domain NPCCSS scale would be best suited to all three settings.

9a. Are a limited number of domains sufficient to meet needs in all scenarios? If 'yes', tick the limited number of domains you believe would be sufficient.

- Ambulation
- Cognition
- Eye movement
- Fine motor
- Hearing
- Memory
- Seizures
- Speech
- Swallowing
- Other

- 50%
- 43.75%
- 12.5%
- 50%
- 0%
- 0%
- 6.25%
- 37.5%
- 43.75%
- 6.25%

9b. Please provide any further insights

- Many respondents did not agree with the question that a limited number of domains could be sufficient to meet the needs all scenarios.
- It was suggested that there should be a focus on domains where change can be expected with therapy and a domain where changes can be quantified. Measuring everything at baseline within clinical trials would show where changes occur.
- The five identified domains are almost always all involved as the disease progresses. Only a small percentage of patients experience hearing loss and seizures, memory is a part of general cognition and too hard to separate from this domain, and eye movement change are difficult to measure.
- In very young children, an additional developmental scale (e.g. Bayleys, Kaufman, etc.) should be used and, in adults, a dementia scale should be used.
- One respondent suggested that it remains unclear if breaking down scores into domains is particularly helpful, while the dynamics of additive sum scores (across domains or without domains) is what matters for outcome trials.
- An additional suggestion included videoing of the walk-test and 9HPT as functionally most relevant; with analysis performed by blinded raters on a 0 +/- 3 scale.

Summary of key insights:

- Many respondents did not agree with the question that a limited number of domains could be sufficient to meet the needs all scenarios.
- In very young children, an additional developmental scale (e.g. Bayleys, Kaufman, etc.) should be used and, in adults, a dementia scale should be used.
- One respondent suggested that it remains unclear if breaking down scores into domains is particularly helpful, while the dynamics of additive sum scores (across domains or without domains) is what matters for outcome trials.
- An additional suggestion included videoing of the walk-test and 9HPT as functionally most relevant; with analysis performed by blinded raters on a 0 +/- 3 scale.

Round 2

In Round 2, a consensus was achieved on six of the nine statements (see Table 3). However, two statements missed reaching a consensus by 1% (69% consensus respectively). These two statements that failed to reach consensus related to whether it was essential to measure all 17-domains during a clinical trial and whether the 5-domain scale satisfies the requirements for use in all clinical settings. The final statement on which consensus was not reached related to the feasibility and need to develop a novel NPC clinical severity scale that satisfies requirements for use in all clinical settings. The panel of experts agreed that it was 'desirable' (81%) and 'achievable' (75%) to determine a single, standardised NPC clinical severity scale for routine clinical practice and clinical research on a global scale within the scope of the existing scales. Further, 100% of respondents agreed that a clinical paper recommending which NPC clinical severity scale should be used in each clinical setting would be valuable to the international clinical and patient community. Consensus was also reached on the statement that the domains measured in the 5-domain scale provided an accurate clinical understanding of NPC severity in clinical practice and trials (87%) and, if there was only one international scale recommended for use evaluating the disease, it would be the 5-domain NPCCSS (81%).
| Question | Round 2 | Agree/neural | Disagree |
|----------|---------|--------------|----------|
| 1. A single, standardised NPC clinical severity scale that can be used in routine clinical practice as well as clinical research on a global scale is desirable | 81% | 19% |
| 2. A single, standardised NPC clinical severity scale that can be used in routine clinical practice as well as clinical research on a global scale is achievable within the scope of existing scales | 75% | 25% |
| 3. A clinical consensus paper recommending which NPC clinical severity scale to use per different clinical setting (comprising routine practice and trial research) would be valuable to the international clinical and patient community | 100% | 0% |
| 4. Assessment across the following 5-domains, provides an accurate clinical understanding of NPC severity: Ambulation, Cognition, Fine motor, Speech, Swallowing | 87% | 13% |
| 5. If only one existing NPC severity scale was to be used for the evaluation of disease in normal clinical practice internationally, I would recommend the 5-domain NPCCSS scale | 81% | 19% |
| 6. It is essential to measure all 17-domains in the NPCCSS during a clinical trial to capture all potential treatment benefits for people living with NPC | 69% | 31% |
| 7. It is sufficient to measure the 5-domains in the 2018 NPCCSS during a clinical trial to capture relevant potential treatment benefits for people living with NPC | 75% | 25% |
| 8. I believe the 5-domain NPCCSS scale satisfies requirements for use in all clinical settings, to standardise assessments on a global scale | 69% | 31% |
| 9. I believe it is feasible and there is a need to develop a new NPC clinical severity scale that satisfies requirements for use in all clinical settings, to standardise assessments on a global scale | 45% | 56% |
| 10. If a new universal NPC clinical severity scale were to be developed, the most important way that it would differ from existing scales would be... |  |  |

**Summary of key insights:**

- To balance breadth with brevity and usability
- To focus on domains where change can be expected with disease progression or therapy
- To evaluate cognition at different ages
- To include quality of life measures
- To determine the impact of epilepsy
- To incorporate video of the performance of patients during the 9HPT and 8-min walk test
- To include age/subtypes-dependant items (e.g. epilepsy and cataplexy in late infantile-juvenile, psychiatry in adolescent-adult...)
- Based on the largest possible source data from natural history cohorts as well as clinical trials and take into account that NPC manifests and progresses differently across age groups and patient populations
- Used across regions, languages and cultures
11. What would be your recommendations to implement a more uniform approach to the use of NPC clinical severity scales?

Summary of key insights:
• To publish a systematic review of the current scales and consensus
• To publish an expert consensus on which scale is preferred for clinical routine practice and which for trials
• To develop detailed SOPs and training on the use of severity scales
• To select a simple scale that can be used in different setting and is sensitive enough to capture the impact of the disease in the NPC patient
• To add QoL measures to 5-domain NPCCSS
• To gain insights from the community on what matters to patients and carers
• To provide patients with score sheets, a booklet or app, to complete regularly and which they present to their doctors at every appointment
• To include clinical scale biochemical markers and neuroimaging
• To evolve clinical scales with available data and distinct uses (e.g. in a specific NPC sub-population, or to track changes in a specific subject), particularly as personalised medicine is a goal of this decade
• To capture real-world results of scales systematically (e.g. INPDR) so that pre/post treatment effect are comparable

The key themes of the responses about a new, universal NPC clinical severity scale (Question 10) included: a need to incorporate quality of life measures, age/subtype dependant items (such as epilepsy and cataplexy in late infantile-juvenile) and a video of patient performance during a 9-Hole Peg Test (9HPT) and 8-minute walk test. When asked for recommendations to implement a more uniform approach to the use of NPC severity scales, participants suggested a published systematic review of the current scales, a published expert consensus, the inclusion of biochemical markers and neuroimaging, and to provide more agency to each patient (such as an app to fill in regularly) to help the doctors achieve personalised treatment. The key insights from the open-ended questions in Round 2 are summarised in Table 3.

Round 3

In Round 3, consensus was reached on five out of the six statements (see Table 4). Despite consensus (81%) achieved during Round 2 that the 5-domain NPCCSS scale was the preferred scale for routine clinical practice and trials, the suggested recommendation in Round 3 that this be positioned as the first-choice scale in routine clinical practice, did not quite reach consensus (68%). However, the panel of 19 experts agreed that the 17-domain NPCCSS scale should be recommended as the first choice to assess the severity of NPC in clinical trial settings, but the domains listed in the 5-domain scale should be prioritised as the primary endpoints (74%). Furthermore, 74% of respondents agreed that there is no need for a new universal scale for all settings to be developed. However, resources or training on how to apply the NPCCSS (17- and 5-domains) should be developed and provided to clinicians working in NPC (89%). Further, 84% agreed that the consensus paper should be reviewed every five years to ensure that recommendations remain accurate.

Table 4

Responses to statement included in Round 3

| Question                                                                 | Round 3 |
|--------------------------------------------------------------------------|---------|
| 1. The 5-domain NPCCSS scale is the first choice for assessing clinical severity of NPC in routine clinical practice | Agree/neutral 68% Disagree 32% |
| 2. The 17-domain NPCCSS scale is the first choice for assessing clinical severity of NPC in clinical trial settings, prioritising the domains in the 5-domain scale (e.g. as primary endpoints) | Agree/neutral 74% Disagree 26% |
| 3. There is no need for a new universal scale for all settings to be developed | Agree/neutral 74% Disagree 26% |
| 4. Resources/training on how to apply the NPCCSS (17- and 5-domains) should be developed and provided to clinicians working in NPC | Agree/neutral 89% Disagree 11% |
| 5. The consensus paper is reviewed periodically to ensure that its recommendations remain accurate | Agree/neutral 100% Disagree 0% |
| 6. The timescale for periodic review of the consensus paper should be every 5 years | Agree/neutral 84% Disagree 16% |

Discussion

This Delphi study achieved consensus during Round 2 that the domains measured in the 5-domain NPCCSS scale provided an accurate clinical understanding of NPC severity. If there was only one international scale recommended for use in routine clinical practice, the respondents would recommend use of the 5-
domain NPCCSS scale. Although this statement achieved consensus in Round 2, amongst a panel of 16 NPC specialists who completed the first two rounds, it did not quite reach consensus in Round 3 from a panel of 19 experts.

In Round 1, respondents highlighted the 5-domain NPCCSS scale as simple, accurate and quick to administer and complete in a routine clinical examination and that its simplicity was valuable for multi-centre trials to support reproducibility and reliability across sites. Further, it was noted that the domains measured in the 5-domain scale are present in nearly all cases of NPC as the disease develops, unlike hearing loss and seizures, which are typically present in only a small percentage of patients. Respondents also noted that the domains measured in the 17-domain scale posed several challenges. For example, as a domain, memory is difficult to separate from the cognition domain and that measuring changes in the eye movement domain can be problematic.

However, the 5-domain scale was seen as insufficient for evaluation of specific subsets of patients, such as those with mainly psychiatric involvement or experiencing seizures. Moreover, answers in Round 1 stressed the importance of the granularity of scores and the comprehensiveness provided by the 17-domain NPCCSS scale, in capturing the progression of late-onset patients with a slowly progressing disease, as well as for measuring change and baseline assessment in clinical trials. This likely led to the 74% consensus in Question 2 of Round 3 that the 17-domain NPCCSS should be the first-choice severity scale in clinical trial settings.

Given these insights, the Core Working Group recommends that the 17-domain NPCCSS is used as the preferred scale to assess NPC severity across clinical trial enrolment and trial outcome measures. However, the domains listed in the 5-domain scale (ambulation, cognition, fine motor, speech and swallowing) should take precedence as primary endpoints as they are the most relevant to describe neurological disease progression and quality of life [16]. As supported by the experts in Round 1, use of the 5-domain NPCCSS is recommended in multi-centre trials to support reproducibility and reliability of results across multiple trial sites. Lastly, the Core Working Group recommends that the 5-domain NPCCSS scale is used within routine clinical practice to assess the clinical severity of NPC patients. These recommendations provide greater global consistency and optimisation of both the 17- and 5-domain NPCCSS scales, whilst not becoming too reductive, which was noted as important by respondents in Round 1.

The Core Working Group also recommends that resources or training on the NPCCSS scales (17- and 5-domains) should be developed and provided to clinicians working with NPC patients to optimise the standardisation of their application. Further, it is advised that this consensus paper should be reviewed every five years to ensure that the recommendations remain accurate.

This Delphi study gathered consensus on the use of six existing NPC clinical severity scales, the findings for which have enabled the research team to deduce several significant recommendations and areas for further development. Drawing on an international panel of NPC clinicians, who treat both paediatric and adult NPC patients, views were gathered from a select, yet representative panel of experienced experts in the field. However, the rarity of NPC disease means that there is a limited global community of NPC specialists. As a result, the size and composition of the expert panel may reduce the generalisability of the results. Nonetheless, the final sample size (16 participants in Round 1 and 2 and 19 participants in Round 3) was greater than the lower limit threshold of 12 [17]. Given the global scale upon which this field operates, the Delphi consensus method, which can be conducted quickly and online, was an appropriate tool for collecting responses. In addition to identifying the areas of consensus, the study highlighted areas where there is less certainty in the field, such as balancing the need for greater consistency of a single, global multi-domain scale with the concern of becoming too reductive.

While a strength of the study was its ability to access an international network of specialists in the field of NPC research and treatment, some of the participants included in the study were those who developed the clinical severity scales under evaluation. The strong opinions from these participants may therefore have introduced some response bias. Further, it is acknowledged that the concept of ‘consensus’ is fairly fluid. While we have consensus, there are still experts among the group who strongly disagree with the recommendations and hold these views firmly. Given the small size of the expert community, research is unlikely to ever reach consensus across all statements. However, the fact that 19 out of 20 invited participants took part in the Delphi study highlights both the perceived importance of this piece of work to the NPC community, and the influential role that patient groups can have in bringing together stakeholders for such projects. According to guidance from the National Institute for Health Research (NIHR) Health Technology, the Delphi technique typically results in a 20% dropout rate over the three rounds of consensus development. In this study, there was an absence of dropouts in any of the three rounds, therefore substantiating the validity of our recommendations.

A key limitation of this study is that it does not offer definitive guidance, as consensus in Round 2 on the 5-domain NPCCSS as the preferred scale for routine clinical practice did not reach final consensus in Round 3. This may be a result of nuances in question phrasing, but the insights obtained were adequate to make several reliable recommendations. As a result, this consensus might facilitate a platform to enable standardisation of data capture and agreement on use for outcome measures.

We believe this study can help to inform and position future discussion around the use of the existing NPC clinical severity scales in clinical practice and trials. As more data, including genomic data, for NPC become available, the findings will become even more important and there may be a need to reconsider which parameters are most important and whether the preferred scales should be amended accordingly. Similarly, outcomes of ongoing trials of disease-modifying therapies for NPC will drive the need to identify the most appropriate clinical severity scale for determining drug efficacy.

Conclusion

Within this Delphi study, experts confirmed that there was no need for a new universal scale for all settings to be developed. However, they highlighted a need to strike a balance between greater optimisation of a global, single multi-domain scale and it becoming too reductive when choosing between the six existing scales. Although consensus was achieved in Round 2 on the 5-domain NPCCSS as the preferred scale for routine clinical practice, this did not achieve a final consensus in Round 3. Given the small size of the expert community, research is unlikely to ever reach consensus across all statements. However, several meaningful recommendations could be drawn from the study. In line with the consensus achieved in Round 3, this study recommends the use of the 17-
domain NPCCSS scale across clinical trial settings, but the five domains measured in the 5-domain scale should be prioritised as primary endpoints. Further, this study recommends the use of the 5-domain NPCCSS scale in routine clinical practice. The findings also indicate a need to develop educational and training materials on how to apply the NPCCSS (17- and 5-domains) for clinicians working in NPC.

**Abbreviations**

9HPT: 9-Hole Peg Test  
NIHR: National Institute for Health Research  
NPC: Niemann-Pick disease Type C  
NPCCSS: NPC Clinical Severity Score  
CNS: Central Nervous System

**Declarations**

**Ethics approval and consent to participate:**
Not applicable

**Consent for publication:**
Not applicable

**Availability of data and materials:**
The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

**Competing interests:**

MP has stock in IntraBio, has consulted for Orphazyme (with monies directed to the Mayo Clinic) and has received research support from Amicus, Glycomine, Idorsia, Orphazyme and Shire-Takeda (with funds to the Mayo Clinic). FP is cofounder and consultant to IntraBio, has stock in IntraBio, and has consulted for Actelion and Orphazyme. CG is an employee of Orphazyme A/S, which is conducting clinical research in NPC.

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**Authors' contributions:**
All authors were involved in the design and analysis of the Delphi study. All authors were also contributors in writing, reading and approving the final manuscript.

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