Development of the PREMature Infant Index (PREMII™), a clinician-reported outcome measure assessing functional status of extremely preterm infants

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ABSTRACT

Background: Comprehensive measures to evaluate the effectiveness of medical interventions in extremely preterm infants are lacking. Although length of stay is used as an indicator of overall health among preterm infants in clinical studies, it is confounded by nonmedical factors (e.g. parental readiness and availability of home nursing support).

Objectives: To develop the PREMature Infant Index (PREMII™), an electronic content-valid clinician-reported outcome measure for assessing functional status of extremely preterm infants (<28 weeks gestational age) serially over time in the neonatal intensive care unit. We report the development stages of the PREMII, including suggestions for scoring.

Methods: We developed the PREMII according to US Food and Drug Administration regulatory standards. Development included five stages: (1) literature review, (2) clinical expert interviews, (3) Delphi panel survey, (4) development of items/levels, and (5) cognitive interviews/usability testing. Scoring approaches were explored via an online clinician survey.

Results: Key factors reflective of functional status were identified by physicians and nurses during development of the PREMII, as were levels within each factor to assess functional status. The resulting PREMII evaluates eight infant health factors: respiratory support, oxygen administration, apnea, bradycardia, desaturation, thermoregulation, feeding, and weight gain, each scored with three to six gradations. Factor levels are standardized on a 0–100 scale; resultant scores are 0–100. No usability issues were identified. The online clinician survey identified optimal scoring methods to capture functional status at a given time point.

Conclusions: Our findings support the content validity and usability of the PREMII as a multi-function outcome measure to assess functional status over time in extremely preterm infants. Psychometric validation is ongoing.

Introduction

Survival of infants born extremely preterm, defined as birth at <28 weeks gestational age (GA) by the World Health Organization, and used interchangeably with extremely low gestational age newborn (ELGAN), has improved over time [1,2]. The majority of extremely preterm infants require intensive care in the neonatal period [3], and survivors remain at risk of short- and long-term morbidities, such as intraventricular hemorrhage, necrotizing enterocolitis ( NEC), chronic lung disease, and neurodevelopmental impairment [4–7].

A challenge for this patient population is the lack of outcome measures to evaluate treatment effects in clinical studies, and clinical assessment tools that monitor how the neonates grow and mature over time. While length of stay (LOS) is often used as an outcome measure in clinical studies, LOS can be influenced by nonmedical factors such as parental...
readiness and availability of home nursing support [8], and institutional variations in organization of care [9], thus limiting the appropriateness of LOS as a measure of infant health and development and as an endpoint in clinical trials. Existing neonatal illness measures, developed primarily to predict mortality and morbidity, combine neonatal data shortly after admission to the Neonatal Intensive Care Unit (NICU) and not over time. For example, the Score for Neonatal Acute Physiology (SNAP) [10] and SNAP Perinatal Extension version II (SNAPPE-II) collect infant data within 24 and 12 h of admission, respectively [11], while the Clinical Risk Index for Babies (CRIB) [12] and CRIB II collect data within 12 and 1 h of admission, respectively, to evaluate risk for mortality [13].

The aim of this study was to develop a comprehensive content-valid clinician-reported outcome (ClinRO) measure, the PREMature Infant Index (PREMII™), to assess the functional status of extremely preterm infants (<28 weeks GA) over time in the NICU, for use in a phase 2 clinical trial. In the current article, we report on the development of the PREMII.

Materials and methods

Study design

Development of the PREMII followed US Food and Drug Administration (FDA) regulatory guidance for patient-reported outcome instruments [14] – standards that apply to other clinical outcome assessment tools, including ClinROs. The PREMII development process (phase 1) consisted of five stages: (1) targeted literature review, (2) clinical expert interviews, (3) Delphi panel survey, (4) development of PREMII items and levels, and (5) cognitive interviews and usability testing of the electronic version. These stages were designed to provide evidence of content validity (i.e. relevance, clarity, and comprehensiveness) of the PREMII to measure accurately the clinical condition, specifically functional status as it changes over time, of the target population (i.e. extremely preterm infants). Additionally, an online clinician survey was conducted to explore potential approaches to scoring the PREMII.

Concept of interest

The concept of interest that the PREMII is designed to measure is functional status. Functional status is defined as an indicator of neonates’ overall health and development encompassing physical, physiological, and clinical status – specifically, what an infant can do and what support the infant requires, on a day-to-day basis, as a reflection of their overall health and development, which can be also considered as maturation over time. Functional status can be assessed with respect to eight key functional areas included in the PREMII (feeding, weight gain, thermoregulation, respiratory support, apnea, bradycardia, desaturation [ABD] events, and oxygen administration). The PREMII can measure functional status as it changes over time with the baby’s development.

The original target concept for the study was discharge readiness. However, evidence gathered from the literature review and clinical expert interviews highlighted challenges to standardizing assessment of physical readiness for discharge. These included variability in standards of neonatal care, home medical support, and proximity and availability of outpatient support. Therefore, the target concept evolved to functional status, which is independent of the health care system or home situation.

Stage 1: Targeted literature review

A targeted literature review was undertaken to identify relevant concepts for inclusion in the PREMII. We searched Embase, MEDLINE, and PubMed for English-language articles published from 2001 to 2015. The search strategy used search terms relevant to factors, attributes, and measures related to physical discharge readiness and LOS for extremely preterm infants (Supplementary Tables 1–2).

Stage 2: Clinical expert interviews

Telephone semistructured qualitative interviews were conducted. Criteria for inclusion included specialized training in neonatology, with ≥10 years of experience caring for preterm infants (Table 1). The interviews were designed to obtain feedback from clinicians on the physical factors infants need to achieve to be considered ready for NICU discharge, as identified by the literature review. See Supplementary Table S3 for an overview of the interview questions.

Stage 3: Delphi panel survey

The Delphi method is a structured communication technique that involves participants (in this case, a panel of experts) who answer a questionnaire in an iterative manner after being provided with an anonymized summary of group responses [15]. Participants were asked to rate the relative importance of factors,
Table 1. Participant inclusion criteria for the PREMII development stages.

| PREMII development stage | Participant inclusion criteria |
|--------------------------|-------------------------------|
| Clinical expert interviews | - General medical license or registration, plus a specialty license or registration in neonatology, as applicable in country of origin  
- Practicing neonatologist with ≥10 years of experience in the care of preterm infants  
- Coauthored hospital management guidelines on the care of preterm infants or a neonatology-related textbook  
- Oral and written fluency in English  
- Availability for a 1-h interview and periodic consulting and/or review of short documents via email or telephone call throughout the duration of the study (~10 months) |
| Delphi panel survey | - General medical license or registration, plus a specialty license or registration in neonatology, as applicable in country of origin  
- Practicing neonatologist with ≥5 years of experience in the care of preterm infants  
- Coauthored peer-reviewed publications, hospital management guidelines on the care of preterm infants, or neonatology-related textbook; was a speaker at conferences or neonatology clinical meetings; or acted as a principal investigator/subprincipal investigator in any past or present neonatology-related trials  
- Oral and written fluency in English  
- Availability to complete up to three brief (10- to 15-min) online surveys |
| Cognitive interviews and usability testing | - Practicing neonatologist with ≥5 years of experience in the care of preterm infants, or neonatal nurse with ≥5 years of experience working in the NICU  
- Oral and written fluency in English  
- Coauthored peer-reviewed publications, hospital management guidelines on the care of preterm infants, or neonatology-related textbook; was a speaker at conferences or neonatology clinical meetings; or acted as a principal investigator/subprincipal investigator in any past or present neonatology-related trials  
- Agreement to complete a 35- to 40-min online survey in English or native language of country of origin |
| Online survey | - General medical license or registration  
- Specialist training in pediatrics or neonatology  
- Practicing neonatologist or pediatrician, with responsibilities that include the care of preterm infants  
- ≥5 years of experience in the care of preterm infants  
- Agreement to complete a 35- to 40-min online survey in English or native language of country of origin |

NICU: neonatal intensive care unit; PREMII: PREMature Infant Index.

identified through the literature review and clinical expert interviews, for the assessment of functional status on a scale of 0 (not at all important) to 5 (extremely important). Additionally, participants were asked to provide feedback on the definitions of the levels for each factor, as well as other important aspects related to the factors and level definitions. The levels for each factor were intended to reflect a scale of functional status from very poor to very good. The purpose was to build consensus on the most important factors for evaluation of a preterm infant’s functional status for inclusion in the PREMII, and to determine the importance of factors.

**Stage 4: Development of PREMII items and levels**

This stage refers to the drafting of the instrument, namely, the formulation of instructions, items or questions capturing each of the identified factors relevant in assessing infant functional status, and response options.

**Stage 5: Cognitive interviews and usability testing of the electronic version**

Note: cognitive interviews and the online clinician survey occurred in parallel.

Semistructured telephone interviews were conducted in two rounds. The purpose of the cognitive interviews was to assess the clarity of the instructions, items, and levels, as well as ease of completion of the instrument. Additionally, the interviews were designed to elicit any potential logistical difficulties with completing the instrument (e.g. due to nursing shift patterns, and differences in geographical or institutional NICU practices). Usability testing of the electronic version was undertaken via interviews to assess the ease of completion on an electronic device (e.g. a tablet device).

**Online clinician survey**

The online survey was developed to explore the most appropriate scoring method to capture accurately a preterm infant’s functional status at a given time point during their NICU stay.

The online survey included questions designed to explore the following: the best approach to calculate daily factor scores, the relative importance of each factor in rating an infant’s overall functional status, and the best approach to calculate a weekly summary score. The questions were based on sample infant profiles that were presented to respondents.

**Daily factor scores**

Participants were presented with example individual factor ratings for each shift over a 24-h period and asked for their opinion on the optimal method to calculate a daily factor score from the shift ratings from the following options: the “most frequent” score across shift scores provided over the 24-h evaluation period, the “numerical average” score across shift scores provided over the 24-h evaluation period, the “worst” (or “best”, as applicable) shift score during that period, the “most recent” shift score during that
period, or “other” (with a request to provide details). Respondents were not asked for a preferred method for calculating a daily weight factor score, as weight is not measured repeatedly across shifts.

Relative importance in rating overall functional status
Participants were asked to rate the relative importance (on a scale of 1 [most important] to 8 [least important]) of each factor in rating an infant’s functional status; respondents were allowed to equally rate multiple factors. Respondents were presented with eight clinical examples of infants and their overall functional status scores over a 7-day period. The overall functional status scores were summarized as the infant’s most frequent, worst (or best), average, and today’s score, as well as the trend over the last 3 days ratings recorded over the 7-day evaluation period.

Weekly summary score
Respondents were asked to rate the weekly summary functional status of the infant (very poor, poor, moderate, good, very good). Additionally, they were asked to rate the importance of each rating approach.

The survey was developed in English and then translated into the following languages: Spanish (Spain, Latin America), French (France), German (Germany), Italian (Italy), Portuguese (Brazil), and Japanese (Japan). Translations met the requirements of the ISO 17100 standard.

Data analysis
Data are reported as descriptive statistics (n and percentage, mean, median). For the clinical expert interviews and cognitive interviews, data were analyzed using qualitative methods. For the online clinician survey, a linear regression analysis was performed to compare weekly summary PREMII scores (“most frequent”, “worst”, “average”, “today”, “trend [past three days]”) with the actual weekly scores provided by the respondents (“weekly summary functional status”) for the online infant profiles.

Results
Stage 1: Targeted literature review
In total, 998 unique abstracts were identified, of which 48 duplicates were excluded. An additional 918 publications were excluded based on predefined exclusion criteria (Figure 1). A total of 32 full-text articles were assessed for eligibility, of which nine were excluded for lack of relevance. The remaining 23 articles were
included in the analysis: 19 related to discharge readiness or LOS (original target concept) [9,16–33] (Table 2), three discussed instruments for assessing infant mortality/morbidity risk [11,17,34] (Supplementary Table 4; one of these reported findings relevant to both LOS and instruments) [17], and two reported national guidelines on the care of preterm/high-risk infants [8,35] (Supplementary Table 5). No measures specifically assessing physical readiness for discharge were identified. From the included literature, over one-half of the articles noted the infant’s cardiorespiratory stability and weight or ability to gain weight as key factors in determining discharge readiness or LOS (Table 2).

### Stage 2: Clinical expert interviews

Four expert neonatologists (RMW [USA], MAT [United Kingdom], IH-P [Sweden], JH [USA]) participated (Supplementary Table 6). The findings were similar to those identified in the literature, namely, oral feeding ability, consistent weight gain, physical/physiological stability, respiratory stability (e.g. absence of apnea), and thermostability (capacity to maintain normal temperature; Table 3). Additionally, two clinical experts noted retinopathy of prematurity (one each in relation to discharge readiness and LOS).

### Stage 3: Delphi panel survey

In total, 17 neonatologists participated in the Delphi panel survey (Supplementary Table 6). In order of importance, participants endorsed respiratory status, ABD events, feeding ability, oxygen supplementation,
Table 4. Participant feedback from rounds 1 and 2 of the cognitive interviews, and the subsequent revisions made to the PREMII following consultation with the clinical experts.

| Item                  | Participant Feedback | Revisions to the PREMII                                                                 |
|-----------------------|----------------------|----------------------------------------------------------------------------------------|
| Respiratory support   |                      | • Suggest clarifying “no supplemental oxygen” in last level                           |
|                       | Round 2              | • “Intratracheal” not a familiar term                                                   |
|                       |                      | • “Negative pressure support” isn’t commonly used                                        |
|                       |                      | • “Only” doesn’t fit with instructions                                                  |
| Oxygen administration  | Round 1              | • Levels are clearer if explicit ranges are reported                                    |
|                       |                      | • Distinction between > 50% and < 50% is important to capture                          |
|                       | Round 2              | • Uncertainty on how to rate the item for infants on low-flow nasal cannula             |
| Apnea                 | Round 1              | • ABD (as one item) should be separated                                                |
|                       | Round 2              | • Important to clarify if infant needed intervention or not                            |
| Bradycardia           | Round 1              | • ABD (as one item) should be separated                                                |
|                       | Round 2              | • Definition may change based on gestational age of the infant                         |
|                       |                      | • “Clinically relevant” may cause confusion                                            |
| Desaturation          | Round 1              | • ABD (as one item) should be separated                                                |
|                       | Round 2              | • Important to clarify if event requires intervention or is self-resolving             |
|                       |                      | • Definition should include “≤”                                                       |
| Feeding               | Round 1              | • Need to define “oral feeds” and add/clarify regarding enteral feeding                |
|                       |                      | • Suggest including reference to feeds via catheter                                     |
|                       | Round 2              | • “Catheter” may cause confusion                                                      |
|                       |                      | • Need to clarify “no feeds occurred”                                                  |
| Weight gain           | Round 1              | • Infants may not be weighed every day                                                 |
|                       | Round 2              | —                                                                                      |
| Temperature           | Round 1              | • “Bundled” not a clear or familiar term                                               |
|                       |                      | • Need to better highlight differences between levels                                   |
|                       | Round 2              | • Some words redundant                                                                |

ABD: apnea, bradycardia, desaturation; IV: intravenous; PREMII: PREMature Infant Index.
thermoregulation, and weight gain (Figure 2). Retinopathy of prematurity was originally included but subsequently removed, as it was not considered to fall under the definition of functional status.

Feedback from the Delphi survey highlighted perceived differences in the relative importance of each ABD event in evaluating functional status, and underlined the need to separate ABD events into individual factors due to potential different underlying physiologic causes of events.

**Stage 4: Development of the PREMII items and levels**

The draft PREMII was developed based on the factors identified in the previous development stages, with further rounds of review by the four clinical experts to refine levels within each factor. Items included in the first version of the PREMII included weight gain, feeding ability, temperature, respiratory support, a single ABD item, and extent of oxygen supplementation.

**Stage 5: Cognitive interviews and usability testing of the electronic version**

The first round of interviews was completed by 23 physicians and nurses; the second round was completed by nine nurses (Supplementary Table 6). Each of the PREMII items’ levels underwent revisions based on findings from the interviews (Table 4). No issues relating to usability of the electronic version of the instrument were identified among the five nurses who participated in usability interviews.

**Online clinician survey**

The online survey was completed by 201 pediatricians and neonatologists (Supplementary Table 6). The “numerical average” score across the 24-h evaluation period was the most frequently reported preferred method for calculating daily factor scores for each of the seven applicable factors (respiratory support, oxygen administration, apnea, bradycardia, desaturation, thermoregulation, and feeding; weight gain was excluded from this analysis because weight is not measured repeatedly across nursing shifts; Supplementary Figure 1). In calculating a weekly summary score, the “trend” score over the past 3 days and “today’s” score were most commonly reported to be most important in determining an infant’s overall functional status (53.0 and 34.7%, respectively), based on the previous 7-day period using hypothetical infant profiles. With regard to relative importance, on a scale of 1–8 (most to least important), respiratory support, apnea, and bradycardia were considered the most important of the eight factors (weight included in the assessment) in rating an infant’s functional status (Supplementary Figure 2). However, there was variability among physicians in terms of relative importance of the factors.

**Finalization of instrument**

The resulting PREMII comprises eight items capturing each of the identified relevant factors (respiratory support, oxygen administration, apnea, bradycardia, desaturation, thermoregulation, feeding, and weight gain), each scored on three to six levels, representing a scale of functional status ranging from very poor to very good (Appendix). The assessment is intended to be repeated over the course of a study to capture change. The intended frequency of administration of the PREMII during a Takeda-sponsored clinical trial is described here. The PREMII assessment will start $\geq 48$ h after birth on the day the infant reaches the next postmenstrual age (PMA) week. For example, if the infant is born at 23 weeks $+ 4$ days, PREMII assessment will begin at 24 weeks PMA, but if an infant is born at 23 weeks $+ 5$ days, PREMII assessment will begin the following PMA week at 25 weeks PMA. In the clinical trial, the PREMII will be administered weekly until 32 weeks PMA and then daily until discharge or 40 weeks PMA, whichever is the earliest. The nurse primarily responsible for the infants’ care will score the PREMII on a tablet device near the end of each nursing shift. The PREMII captures a 24-h period and the number of PREMII assessments carried out during this time will depend on the duration of nursing shifts (e.g. 8 or 12 h). Formal training will be provided for PREMII users before using the tool.

**Discussion**

We developed the PREMII, a ClinRO with evidence of content validity, designed to measure treatment benefit in clinical trials by assessing the functional status of extremely preterm infants in the NICU. To our knowledge, the PREMII is the first comprehensive multifunction outcome measure developed to capture and measure health and development repeatedly in extremely preterm infants over time from birth until discharge from the NICU. While illness severity scores are available for the purpose of predicting mortality and morbidity [10–13],
they primarily collect infant data within 24 h of admission to the NICU, and are not designed to assess the process of development and maturation over time. LOS is considered an important outcome measure in clinical studies; however, using LOS to assess treatment effect in neonatal studies can be challenging on account of factors not directly related to infant health that may influence time to discharge, such as parental readiness and organizational factors [8,9]. The PREMII includes eight infant health factors (respiratory support, oxygen administration, apnea, bradycardia, desaturation, thermoregulation, feeding, and weight gain), which will enable the assessment of functional status as an outcome measure in neonatal studies, thus providing a comprehensive approach to comparing groups of infants, for example, when examining the effects of treatments.

The development stages demonstrated that the PREMII adequately measures functional status in extremely preterm infants and therefore has good content validity, which is in accordance with US FDA regulatory standards for developing patient-reported outcome instruments [14]. Development of the PREMII was guided by neonatologists and NICU nurses, who provided their opinions based on clinical experience. Through the Delphi approach, expert neonatologists reached consensus agreement on the factors for inclusion in the PREMII, and the importance of factors. An example of this was the consensus that respiratory status and the level of support required would adequately measure the severity of lung disease. Participants represented countries across a number of global regions, including North America, Europe, Latin America, and Asia-Pacific. This approach highlighted cultural differences in clinical practice across regions and aided the development of the PREMII to maximize applicability. Although designed for clinical trials, the PREMII could be used as a key performance indicator in NICUs, for benchmarking between sites/hospitals, or to adjust for illness severity as extremely preterm infants approach term equivalent age. The tool may even provide a structured approach to informing discharge readiness by providing the relevant data to inform discharge decision making. It should be noted, however, that the PREMII is not specifically intended to predict discharge readiness or LOS, but rather to assess functional status over time. Furthermore, although the PREMII was developed specifically for the population of extremely preterm infants (<28 weeks GA), it could be applied to infants born at other GA during their growth and development in the NICU as the factors for assessment will remain consistent.

There are limitations of the PREMII that should be considered. One is that local policies regarding neonatal care may differ (e.g. oxygen saturation limits), as well as definitions of what constitutes an event (e.g. apnea or bradycardia). The difficulty of controlling for differing standards of care and the potential for variability of practice across sites remain a challenge in clinical research. We standardized the factors and level ranges captured by PREMII items to the greatest extent by gaining consensus input from expert clinicians based on global considerations. Additionally, instructions and training are included in the PREMII instrument to minimize variation. A further consideration is the element of subjectivity in the clinician responses (e.g. “worst experience”). The development steps were designed to ensure appropriate and clear response options, to measure the abilities to respond using the response options, and consistency of interpretation across respondents. PREMII items and levels were developed with extensive clinical expert input and we expect a high degree of consistency in item interpretation; there remains, however, the possibility that interpretation may vary among clinicians. We acknowledge that some factors (e.g. feeding and weight gain) can be affected by various comorbidities, such as NEC; this will be further explored in a separate study (outlined below).

A separate real-world, prospective, psychometric validation study is underway to evaluate the psychometric properties of the PREMII for clinical application. Specifically, we will evaluate inter- and intrarater reliability, construct validity, criterion (i.e. predictive) validity, sensitivity to change, and responder definition. Comorbidities, especially those that impact nutrition such as NEC, will be captured in the study, and outcomes will be categorized. Additionally, the psychometric validation study will further explore the scoring of the PREMII and evaluate the optimal frequency of administration of PREMII in real-world clinical practice. The PREMII is designed for use from shortly after birth through discharge from the NICU; longer term validation (e.g. at 2 years of age) is challenging owing to variation in clinical practice and patient attrition over time.

In conclusion, the PREMII represents a ClinRO measure with well-supported content validity and usability to assess the functional status of extremely preterm infants serially over time in the NICU. It is hoped this unique tool will be suitable for use in neonatal clinical studies.

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Statement of ethics
The authors have no ethical conflicts to disclose. Ethical approval was not required by the institutional review board because the study did not involve direct patient involvement or personal health information.

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Data availability statement
All relevant data are within the paper and its supplementary information files.

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