Validation of the Children’s International Mucositis Evaluation Scale (ChIMES) in paediatric cancer and SCT

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Background: Objectives were to describe the reliability and validity of a new paediatric-specific mucositis scale, the Children’s International Mucositis Evaluation Scale (ChIMES).

Methods: In a multi-centre prospective study, children aged 0 to ≤18 years were eligible if they were receiving any of the following: myeloablative stem cell transplantation (SCT), ≥60 mg m⁻² course⁻¹ doxorubicin or ≥12 g m⁻² methotrexate. Multiple measures of mucositis were included along with ChIMES. Respondents were parent proxy report for children aged ≤12 years, and child self-report for children aged 12–18 years and 8 to ≤12 years. Mucositis diaries were completed at baseline and on Days 7–17 following chemotherapy/conditioning. On Day 14, the respondent reported presence of mucositis and change since the previous day.

Results: The 185 respondents included parents (N = 98), children aged 12–18 years (N = 66) and children aged 8 to < 12 years (N = 21). Test–retest reliability was excellent for ChIMES Total Score and ChIMES Percentage Score with r > 0.8 for all respondent types. Criteria for construct validation were met across all measures. ChIMES also demonstrated responsiveness with significant differences between baseline and Day 14.

Conclusion: ChIMES is a paediatric-specific measure of mucositis with favourable psychometric properties. It exhibits reliability, construct validity and responsiveness. ChIMES should be incorporated into clinical trials of mucositis prevention and treatment in paediatric cancer and SCT.

Oral mucositis is a common consequence of chemotherapy and stem cell transplantation (SCT) that reduces quality of life and results in morbidity in adults and children with cancer (Sonis, 2011). Prevention of mucositis is important to patients, parents and health-care providers (Ethier et al, 2012). In order to determine the most effective preventive and treatment strategies, outcome measures are needed to reliably quantify the degree of oral mucositis and its resultant morbidity. Considerable effort has resulted in many different mucositis scales being developed for adults, and currently, there are several options for the valid
measurement of mucositis among adults receiving chemotherapy and radiotherapy, including those undergoing SCT (Sonis et al., 2004). However, there remains a lack of validated instruments for assessing mucositis in children (Tomlinson et al., 2007).

Children have unique measurement issues compared with adults. Young children are more difficult to assess due to lack of co-operation with oral cavity examination. In addition, attribution of functional symptoms may not be linkable to an aetiology. For example, if a young child refuses to eat, it may be difficult to know whether this behaviour is related to mucositis, nausea or anorexia. The lack of a feasible, reliable and valid scoring system for mucositis in children has created an obstacle to intervention and epidemiological mucositis research in paediatric cancer (Qutob et al., 2013).

In order to address this gap in the literature, we formed a multi-disciplinary group with expertise in paediatric mucositis. Our previous work established the need for a new paediatric mucositis scale, generated items, drafted the scale and evaluated early psychometrics with a focus on understandability, content validity and acceptability from parents and children with cancer. Three iterations were required to arrive at the final version of the instrument that was then termed the Children’s International Mucositis Evaluation Scale (ChIMES; Tomlinson et al., 2007, 2008a, b, 2009a, b, 2010).

This manuscript describes the evaluation of reliability and validity of ChIMES in children with cancer or undergoing SCT. We hypothesised that ChIMES would be reliable (test–retest, inter-rater and internal consistency) and valid (convergent validity and responsiveness).

MATERIALS AND METHODS

We conducted a multi-centre prospective study of children at increased risk of developing oral mucositis in order to evaluate the psychometric properties of ChIMES. We measured oral mucositis at baseline (conducted between day — 2 to day 5 when mucositis was not expected) and then daily between Days 7 and 17 (when mucositis was expected) following start of chemotherapy or conditioning. Three groups of respondents were included: parent/guardian proxy respondents for children aged <12 years; child self-respondents aged 12–18 years; and child self-respondents aged 8 to <12 years.

The Research Ethics Board at The Hospital for Sick Children and all participating institutions approved the study. All participants provided written informed consent or assent as appropriate.

Subjects. Children aged 0 to ≤18 years were eligible if they were to receive any of the following treatments: myeloablative SCT, >60 mg m⁻² course⁻¹ doxorubicin or >12 g m⁻² methotrexate. We excluded respondents unable to read English and those who did not have the cognitive ability to complete instruments. Participants were recruited from: The Hospital for Sick Children, Toronto, ON, Canada; Children’s National Medical Center, Washington, DC, USA; and Lucile Salter Packard Children’s Hospital at Stanford, Palo Alto, CA, USA.

Procedures. Respondents were approached in the inpatient or clinic setting before the start of the chemotherapy cycle or SCT procedure. Measures of mucositis were ChIMES, World Health Organisation (WHO) mucositis scale, mucositis visual analogue scale (VAS), National Cancer Institute Common Terminology Criteria (NCI-CTC) v3.0 functional/symptomatic mucositis scale and the Oral Mucositis Daily Questionnaire (OMDQ). The recall period for all instruments was the same day except for the OMDQ, which was the past 24 h.

A study training manual, which detailed standard operating procedures, was circulated to all participating sites to maximise consistency in evaluations. Participants were provided descriptions and pictures of erythema and ulcers. Participants were encouraged to consult with their doctor or nurse if they were unsure about their presence. A standard light source was not used across sites. Timing and conduct of assessments were further standardised to occur before eating or drinking and at approximately the same time every day. The order of evaluations was also standardised. The baseline assessment occurred between 2 days before initiation of chemotherapy or SCT conditioning until 5 days following chemotherapy initiation as mucositis is unlikely to occur within this time frame. Follow-up assessments were then conducted once daily between Days 7 to 17 after starting chemotherapy or conditioning. Responses were recorded in a paper diary. The evaluation of test–retest reliability occurred on Days 13 and 14, when maximum mucositis was expected.

For children aged <8 years, the parent completed all measures and the child did not participate. On Day 14, the parent answered the following three additional questions: (a) reported whether their child was experiencing any mouth pain or sores that day (yes or no); (b) reported whether oral mucositis had changed since the previous day (Day 13) on a five-point scale consisting of much worse, somewhat worse, no change, somewhat better and much better; and (c) completed the Faces Pain Scale—Revised (Hicks et al., 2001) to explore how this measure is associated with the pain question of ChIMES.

For children aged ≥12 years, the child completed all instruments although he/she could request assistance from the parent/guardian if necessary. These older children also completed the three additional Day-14 questions as outlined above. The parent did not participate in daily assessments although on Day 14, the parent or guardian also completed ChIMES so that inter-rater reliability could be examined. For children aged 8 to <12 years, only the parent, only the child or both could participate. In the case of child self-report, the child completed the three additional Day-14 questions and the parent completed Day-14 ChIMES.

Outcome measures

ChIMES. ChIMES (parent version illustrated in Appendix 1) consists of seven elements: (1) Amount of mouth or throat pain (ChIMES1), (2) Effect of mouth or throat pain on swallowing (ChIMES2), (3) Effect of mouth or throat pain on eating (ChIMES3), (4) Effect of mouth or throat pain on drinking (ChIMES4), (5) Receipt of pain medication (ChIMES5), (6) Receipt of pain medication for mouth or throat pain (ChIMES6), and (7) Presence of ulcers (ChIMES7). Because in young children failure to eat or drink may not be attributable to mucositis vs other aetiologies such as nausea or anorexia, the instrument allows the respondent to choose ‘I can’t tell’ if the respondent is uncertain if the cause of functional impairment is due to oral mucositis. ChIMES1–4 each received a score of 0–5 where 5 is the worst degree of symptoms. ChIMES5 received a score of 1 if the child had received pain medications and ChIMES6 received a score of 1 if the child received pain medications because of mucositis. Otherwise, ChIMES5 and 6 received a score of 0. Finally, ChIMES7 received a score of 1 if oral ulcers were present and 0 if absent. Any question that was scored as missing or ‘I can’t tell’ was excluded from the total possible score. If all the questions were answered, the maximum score was 23. The ChIMES Total Score was the sum of all scores; ‘I can’t tell’ responses and missing responses both received a score of 0. The ChIMES Percentage Score was the ChIMES Total Score over the total maximum score taken into account ‘I can’t tell’ responses (by subtracting these items from the maximum score) multiplied by 100. In other words, the ChIMES Total Score does not take into account ‘I can’t tell’ or missing responses as they are given a score of 0 and the ChIMES Total Score keeps the weighting of all components constant.
In contrast, ChIMES Percentage Score does take into account ‘I can’t tell’ responses by changing the maximum score possible. Higher scores correspond to worse mucositis.

WHO. The WHO scale is based upon the ability to eat and drink combined with objective signs of mucositis, namely erythema and ulceration (World Health Organization, 1979). Visualisation of the oral cavity is critical for scoring, as the presence of oral ulcers delineates a WHO mucositis grade of $\geq 2$ vs $< 2$. It is one of the most commonly used outcome measures in clinical research (Sonis et al, 2004). WHO grade ranges from 0 to 4 where higher scores correspond to worse mucositis.

VAS. We used a horizontal 10-cm VAS anchored at 0 (no mouth or throat pain) and 10 (most severe mouth or throat pain) and asked the respondent to indicate that day’s level of pain. We have previously used this outcome measure to validate the Oral Mucositis Assessment Scale (OMAS) and as an outcome measure in a randomised trial (Sung et al, 2007a, b).

NCI-CTC. The NCI-CTC is the standard adverse event reporting system used by the National Cancer Institute and it is widely used for grading oral mucositis (National Cancer Institute, 2003). The NCI-CTC v3.0 mucositis functional/symptomatic scale was used. NCI-CTC grade ranges from 0 to 5 where higher scores correspond to worse mucositis.

OMDQ. The OMDQ was developed through multiple focus groups and one-on-one interviews with cancer patients (Bellm et al, 2002; Stiff et al, 2006). It consists of seven questions that relate to: (1) Amount of mouth and throat pain (OMDQ1), (2) Effect of pain on sleeping (OMDQ2), (3) Effect on swallowing (OMDQ3), (4) Effect on drinking (OMDQ4), (5) Effect on eating (OMDQ5), (6) Effect on talking (OMDQ6) and (7) Amount of diarrhoea (OMDQ7). The OMDQ has been validated in paediatric cancer by parent proxy report and child self-report for all items other than for the diarrhoea item. Thus, OMDQ7 was not included in this study. Each component of the OMDQ was scored separately as an aggregate score has not been validated. For each component, the score ranges from 0 to 4 where higher scores correspond to worse mucositis.

**Statistical considerations.** All analyses were stratified by respondent type. ChIMES evaluations focused on ChIMES Total Score and ChIMES Percentage Score, but we also illustrated the properties of individual items. To evaluate the test–retest reliability of ChIMES, we calculated the Spearman’s correlation coefficient between Days 13 and 14 for those who reported no change in mucositis between these days and for all respondents. We hypothesised an $r \geq 0.7$. To evaluate the inter-rater reliability of ChIMES, we calculated the Spearman’s correlation coefficient between parents and children aged 8 to <12 and 12–18 years on Day 14 and anticipated an $r \geq 0.5$. We evaluated internal consistency by Cronbach’s alpha and anticipated an alpha $\geq 0.7$ (Streiner and Norman, 1995).

To evaluate convergent validity of ChIMES, we hypothesised that ChIMES scores would be positively correlated with WHO, VAS, NCI-CTC and OMDQ. For this analysis, we described the Spearman’s correlation coefficients using all evaluations but in order to account for the same child providing multiple measures at baseline and on Days 7–17, we obtained the $P$ values using a repeated-measures linear model with Proc Mixed in SAS (Cary, NC, USA). We anticipated a Spearman’s correlation of $\geq 0.35$ based on our previous studies. To evaluate the responsiveness of ChIMES, we compared the ChIMES scores obtained at baseline with Day-14 evaluations in children who had oral mucositis on Day 14 and in all children. These two scores were compared using the Wilcoxon signed-rank test. An exploratory aim was to compare the Faces Pain Scale-Revised and the ChIMES pain question (ChIMES1) on Day 14 to address the question of whether the smiley faces scale used in ChIMES may be confounding pain affect and intensity. These scores were evaluated using the Spearman’s correlation coefficient.

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**Figure 1. Flow diagram of participants stratified by respondent type.**

| Total approached: 222 | Total refused: 30 |
|-----------------------|-------------------|
| Parent proxy report=114 | Parent proxy report=13 |
| Child (12–18 years) self-report=82 | Child (12–18 years) self-report=14 |
| Child (8 to <12 years) self-report=26 | Child (8 to <12 years) self-report=3 |

| Total participants: 192 | Total not evaluable: 7 |
|------------------------|-----------------------|
| Parent proxy report=101 | Parent proxy report=3 |
| Child (12–18 years) self-report=68 | Child (12–18 years) self-report=2 |
| Child (8 to <12 years) self-report=23 | Child (8 to <12 years) self-report=2 |

| Total evaluable participants: 185 | Diary lost: 6 |
|-------------------------------|----------------|
| Parent proxy report=98 | Parent proxy report=3 |
| Child (12–18 years) self-report=66 | Child (12–18 years) self-report=2 |
| Children (8 to <12 years) self-report=21 | Child (8 to <12 years) self-report=1 |

| Withdrew: 1 | |
|-------------|---|
| Parent proxy report=3 | |
| Child (12–18 years) self-report=2 | |
| Child (8 to <12 years) self-report=1 | |
The sample size was based on evaluating the test–retest reliability of ChIMES. Assuming that the \( r \) under the null hypothesis was 0.4 and under the alternate hypothesis was 0.7, an \( \alpha \) of 0.05 and a \( \beta \) of 0.20, we planned to recruit at least 90 parent respondents to ensure that we had 45 who reported no change in oral mucositis between Days 13 and 14.

**RESULTS**

Between 6 July 2010 and 29 April 2013, 222 potentially eligible respondents were evaluated. Figure 1 outlines the flow of participants; 30 refused and 7 were not evaluable, thus leaving 185 respondents in the final analysis. Of these, 98 were parent/guardian proxy respondents for children aged <12 years, 66 were child self-respondents aged 12–18 years and 21 were child self-respondents aged 8 to <12 years. Of the 34 children aged 8 to <12 years, 14 children agreed to self-report mucositis scores alongside their parents, 13 children refused and only their parents participated and 7 children participated alone without their parents. Table 1 illustrates the demographics of the study cohort stratified by respondent type. Approximately 40–45% of respondents had previous experience with mucositis.

**DISCUSSION**

We have demonstrated that ChIMES, a new paediatric-specific measure of oral mucositis, is reliable, valid and responsive to change in children and adolescents with cancer and undergoing SCT. In order to decide whether to incorporate an instrument as an outcome measure in clinical trials, assessment of all of these properties is important. Our data also suggest that ChIMES may be

**Table 1.** Demographics of the study cohort stratified by respondent type

| Characteristic | Parent proxy report for children aged <12 years (N = 98) | Child self-report aged 12–18 years (N = 66) | Child self-report aged 8 to <12 years (N = 21) |
|---------------|-------------------------------------------------|---------------------------------|---------------------------------|
| **Parent**    |                                                 |                                 |                                 |
| Male (%)      | 17 (17.5)                                       | —                               | 5 (38.5)                        |
| Median age (IQR) in years | 36.8 (32.8, 41.1) | —                               | 41.6 (39.5, 45.1) |
| At least college education (%) | 75 (76.5) | —                               | 10 (47.6)                       |
| **Child**     |                                                 |                                 |                                 |
| Male (%)      | 57 (58.2)                                       | 40 (60.6)                       | 15 (71.4)                       |
| Median age (IQR) in years | 5.3 (3.0, 8.5) | 14.7 (13.5, 16.6) | 9.8 (9.0, 11.1) |
| **Diagnosis (%)** |                                             |                                 |                                 |
| Leukemia/lymphoma | 34 (34.7)                                  | 28 (42.2)                       | 8 (38.1)                        |
| Solid tumour | 31 (31.6)                                      | 30 (45.5)                       | 8 (38.1)                        |
| Brain tumour | 15 (15.3)                                      | 4 (6.1)                         | 3 (14.3)                        |
| Metabolic | 4 (4.08)                                       | —                               | —                               |
| Other | 14 (14.3)                                      | 4 (6.1)                         | 2 (9.5)                         |
| **Treatment at enrolment (%)** |                                             |                                 |                                 |
| Chemotherapy | 16 (16.3)                                      | 35 (53.0)                       | 7 (33.3)                        |
| Stem cell transplantation | 82 (83.7)                                  | 31 (47.0)                       | 14 (66.7)                       |
| TBI containing conditioning | 16/62 (19.5)                          | 7/31 (22.6)                     | 6 (42.9)                        |
| Previous history of mucositis (%) | 43 (43.9) | 27 (40.9)                       | 9 (42.9)                        |

Abbreviations: IQR = interquartile range, TBI = total body irradiation.

**Table 2.** Test–retest reliability of ChIMES total and percentage scores measured on Days 13 and 14*

| Respondent type | ChIMES total score \( r \) (P-value) | ChIMES percentage score \( r \) (P-value) |
|-----------------|--------------------------------------|------------------------------------------|
| **Parent proxy report** |                                       |                                          |
| Report no change, n = 53 | 0.967 (<0.0001) | 0.968 (<0.0001) |
| All, n = 98 | 0.941 (<0.0001) | 0.942 (<0.0001) |
| **Child self-report aged 12–18 years** |                                       |                                          |
| Report no change, n = 33 | 0.894 (<0.0001) | 0.888 (<0.0001) |
| All, n = 66 | 0.854 (<0.0001) | 0.852 (<0.0001) |
| **Child self-report aged 8 to <12 years** |                                       |                                          |
| Report no change, n = 10 | 1.000 (<0.0001) | 1.000 (<0.0001) |
| All, n = 21 | 0.902 (<0.0001) | 0.902 (<0.0001) |

Abbreviation: ChIMES = Children’s International Mucositis Evaluation Scale.

*Table represents Spearman’s correlation coefficients with \( P \) values in parentheses for ChIMES total and percentage scores measured on 2 consecutive days (Days 13 and 14). Results are presented for those who indicated no change in mucositis between Days 13 and 14 and all respondents.
Validation of ChIMES

Table 3. Construct validation of ChIMES total and percentage scores*

| Respondent type | Parent proxy report for children aged < 12 years (N = 98) | Child self-report aged 12-18 years (N = 66) | Child self-report aged 8 to < 12 years (N = 21) |
|-----------------|-------------------------------------------------------|------------------------------------------|-----------------------------------------------|
| Other measures of mucositis | ChIMES total score | ChIMES percentage score | ChIMES total score | ChIMES percentage score | ChIMES total score | ChIMES percentage score |
| WHO Mucositis | 0.847 (<0.0001) | 0.846 (<0.0001) | 0.782 (<0.0001) | 0.785 (<0.0001) | 0.830 (<0.0001) | 0.827 (<0.0001) |
| VAS Mucositis | 0.854 (<0.0001) | 0.857 (<0.0001) | 0.808 (<0.0001) | 0.809 (<0.0001) | 0.727 (<0.0001) | 0.731 (<0.0001) |
| CTC Mucositis | 0.862 (<0.0001) | 0.863 (<0.0001) | 0.779 (<0.0001) | 0.781 (<0.0001) | 0.795 (<0.0001) | 0.795 (<0.0001) |
| OMDQ1* | 0.903 (<0.0001) | 0.906 (<0.0001) | 0.851 (<0.0001) | 0.852 (<0.0001) | 0.813 (<0.0001) | 0.822 (<0.0001) |
| OMDQ2 | 0.723 (<0.0001) | 0.706 (<0.0001) | 0.585 (<0.0001) | 0.587 (<0.0001) | 0.551 (<0.0001) | 0.549 (<0.0001) |
| OMDQ3 | 0.900 (<0.0001) | 0.911 (<0.0001) | 0.882 (<0.0001) | 0.882 (<0.0001) | 0.922 (<0.0001) | 0.917 (<0.0001) |
| OMDQ4 | 0.896 (<0.0001) | 0.908 (<0.0001) | 0.884 (<0.0001) | 0.886 (<0.0001) | 0.928 (<0.0001) | 0.926 (<0.0001) |
| OMDQ5 | 0.908 (<0.0001) | 0.917 (<0.0001) | 0.904 (<0.0001) | 0.905 (<0.0001) | 0.900 (<0.0001) | 0.906 (<0.0001) |
| OMDQ6 | 0.864 (<0.0001) | 0.876 (<0.0001) | 0.724 (<0.0001) | 0.726 (<0.0001) | 0.783 (<0.0001) | 0.780 (<0.0001) |

Abbreviations: ChIMES = Children’s International Mucositis Evaluation Scale; CTC = National Cancer Institute’s Common Terminology Criteria v3.0; WHO = World Health Organisation mucositis scale; VAS = pain visual analogue scale; OMDQ = Oral Mucositis Daily Questionnaire.
*Table represents Spearman’s correlation coefficients with P values derived from a generalised linear mixed model with repeated measures in parentheses.
*OMDQ items were as follows: (1) Amount of mouth and throat pain (OMDQ1), (2) Effect of pain on sleeping (OMDQ2), (3) Effect on swallowing (OMDQ3), (4) Effect on drinking (OMDQ4), (5) Effect on eating (OMDQ5), and (6) Effect on talking (OMDQ6).

Table 4. Responsiveness of ChIMES total and percentage scores between baseline and day 14*

| Respondent type | ChIMES total scores | ChIMES percentage scores |
|-----------------|---------------------|----------------------------|
| Mean baseline total score ± s.d. | Mean day 14 total score ± s.d. | Mean difference (95% CI) P-value | Mean baseline percentage score ± s.d. | Mean day 14 percentage score ± s.d. | Mean difference (95% CI) P-value |
| Parent proxy report | | | | | | |
| Report D14 mucositis (n = 51) | 1.5 ± 3.6 | 13.0 ± 7.5 | 11.4 (−1.0, 22.0) <0.0001 | 6.4 ± 15.8 | 58.7 ± 32.6 | 51.0 (−4.3, 95.7) <0.0001 |
| All patients (n = 98) | 1.5 ± 3.6 | 8.0 ± 8.5 | 6.6 (−3.0, 22.0) <0.0001 | 6.4 ± 15.8 | 36.0 ± 37.5 | 29.5 (−13.0, 95.7) <0.0001 |
| Child self-report aged 12-18 years | | | | | | |
| Report D14 mucositis (n = 26) | 0.9 ± 2.2 | 10.6 ± 7.2 | 9.8 (1.0, 21.0) <0.0001 | 3.7 ± 9.5 | 46.5 ± 31.3 | 43.3 (4.3, 91.3) <0.0001 |
| All patients (n = 66) | 0.9 ± 2.2 | 5.1 ± 7.1 | 4.3 (−1.0, 19.0) <0.0001 | 3.7 ± 9.5 | 22.3 ± 31.0 | 18.9 (−4.3, 82.6) <0.0001 |
| Child self-report aged 8 to < 12 years | | | | | | |
| Report D14 mucositis (n = 6) | 0.9 ± 1.5 | 12.6 ± 8.9 | 12.3 (0.0, 23.0) 0.063 | 3.7 ± 6.5 | 54.7 ± 38.7 | 53.6 (0.0, 100.0) 0.063 |
| All patients (n = 21) | 0.9 ± 1.5 | 5.1 ± 8.1 | 3.7 (−5.0, 23.0) 0.333 | 3.7 ± 6.5 | 22.0 ± 35.4 | 16.1 (−21.7, 100.0) 0.363 |

Abbreviations: ChIMES = Children’s International Mucositis Evaluation Scale; CI = confidence interval.
**Table represents difference between day 14 and baseline with P-values from Wilcoxon signed-rank test.

used confidently for child self-report in those aged ≥ 12 years and likely for those as young as 8 years of age.

There are several components of ChIMES that are particularly relevant to children. First, ChIMES focuses on functional elements, as these were considered more clinically important rather than simply the presence of ulcers (Tomlinson et al, 2009b). Second, the assessment of ulcers is limited to a yes/no question rather than a detailed assessment as is conducted for the OMAS. With the OMAS (Sonis et al, 2001), nine sites of the oral cavity are evaluated for erythema and ulceration. Although we found the OMAS to be valid in paediatric cancer patients receiving chemotherapy (Sung et al, 2007b), the correlation coefficients between OMAS with WHO and VAS were much lower (0.56 and 0.37, respectively) compared with ChIMES. We also found OMAS difficult for some
children given the time required for oral cavity evaluation. These issues suggest that ChIMES is preferable to OMAS for children, although OMAS is likely to be an excellent outcome measure for adult cancer patients. Third, ChIMES allows ‘I can’t tell’ responses. With WHO, the inability to attribute symptoms to aetiology and the inability to visualise the oral cavity result in a missing score. Given the importance of avoiding missing scores in clinical trials, these issues suggest that ChIMES may be a better measure for paediatric clinical trials compared with WHO (Little et al., 2012).

We calculated and evaluated two ChIMES outcomes, the ChIMES Total Score and the ChIMES Percentage Score. Our results suggest that either may be used in clinical trials. However, the incorporation of ‘I can’t tell’ responses may make the ChIMES Percentage Score preferable.

We had a limited number of children 8 to <12 years of age who self-reported mucositis scores, and there are several aspects of our trial that merit specific consideration. First, some children reported mucositis alongside their parents; we do not know whether the simultaneous completion of diaries by parents and children may have biased the child responses favourably. Second, we invited 34 children aged 8 to <12 years to our study and only 21 agreed to self-report mucositis scores. Thus, our results may not be generalisable to all 8- to <12-year olds. Third, the evaluation of responsiveness focused on those with mucositis on Day 14, and only six respondents met this criterion within the 8- to <12-year self-report respondent group. Consequently, the failure to show a statistically significant difference for the analysis is more likely related to inadequate power rather than lack of responsiveness in this age range. Further, the magnitude of the differences between baseline and Day 14 is similar to that seen in parent and child respondents aged 12-18 years.

The major strengths of our study include the multi-centre design, incorporation of multiple measures of mucositis and evaluation of responsiveness in addition to reliability and validity. However, our results must be interpreted in light of its limitations. As previously noted, the 8- to <12-year-old group is likely to be self-selected as a more compliant and motivated group. Second, we only included English speakers in this study. Another limitation is that we have little insight into whether parents were able to accurately assess subjective symptoms in the youngest children using the six-point scale or whether a dichotomous or simpler scale would have been preferable. Finally, another limitation is that we do not have clinically defined thresholds for ChIMES scores that delineate the presence or absence of mucositis or degree of severity (such as mild, moderate and severe).

Future work should focus on determining whether children aged <8 years can self-report mucositis and parent/guardian evaluation of mucositis in the youngest children. Future work should also focus on identifying clinically meaningful thresholds for ChIMES scores to categorise the presence of mucositis and degree of severity. Additionally, ChIMES should be translated into other languages and evaluated in other cultural contexts. In conclusion, ChIMES is a paediatric-specific measure of mucositis with favourable psychometric properties. It exhibits test–retest reliability, inter-rater reliability, internal consistency, construct validity and responsiveness. ChIMES should be incorporated into mucositis prevention and treatment clinical trials in paediatric cancer and SCT.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

REFERENCES

Bell LA, Cunningham G, Durnell L, Eilers J, Epstein JB, Fleming T, Fuchs HJ, Haskins MN, Horowitz MM, Martin PJ, McGuire DB, Mulane K, Oster G (2002) Defining clinically meaningful outcomes in the evaluation of new treatments for oral mucositis: oral mucositis patient advisor board. Cancer Invest 20(5–6): 793–800.

Ethier MC, Regier DA, Tomlinson D, Judd P, Doyle J, Gassas A, Naevi A, Sung L (2012) Perspectives toward oral mucositis prevention from parents and health care professionals in pediatric cancer. Support Care Cancer 20(8): 1771–1777.

Hicks CL, von Baeyer CL, Spafford PA, van Korlaar I, Goodenough B (2001) The Faces Pain Scale-Revised: toward a common metric in pediatric pain measurement. Pain 93(2): 173–183.

Little RJ, Cohen ML, Dickerson K, Emerson SS, Farrar JT, Neaton JD, Shih W, Siegel JP, Stern H (2012) The design and conduct of clinical trials to limit missing data. Stat Med 31(28): 3433–3443.

National Cancer Institute (2003) Common Terminology Criteria for Adverse Events v3.0 (CTCAE) Cancer Therapy Evaluation Program, March 31, 2003 http://ctep.cancer.gov, Publish Date: August 9, 2006.

Qutob AF, Gue S, Revesz T, Logan RM, Keefe D (2013) Prevention of oral mucositis in children receiving cancer therapy: a systematic review and evidence-based analysis. Oral Oncol 49(2): 102–107.

Sonis ST (2011) Oral mucositis. Anticancer Drug Rev 22(7): 607–612.

Sonis ST, Elting LS, Keefe D, Peterson DE, Schubert M, Hauer-Jensen M, Bekele BN, Raber-Durlacher J, Donnelly JP, Rubinstein EB (2004) Perspectives on cancer therapy-induced mucosal injury: pathogenesis, measurement, epidemiology, and consequences for patients. Cancer 100(9 Suppl): 1955–2025.

Sonis ST, Oster G, Fuchs H, Bellm L, Bradford WZ, Edelsberg J, Hayden V, Eilers J, Epstein JB, Levesque FG, Miller C, Peterson DE, Schubert MM, Spijkervet FK, Horowitz M (2001) Oral mucositis and the clinical and economic outcomes of hemopoietic stem-cell transplantation. J Clin Oncol 19(8): 2201–2205.

Stiff PJ, Erder H, Bensinger WI, Emmanouilides C, Gentile T, Isitt J, Lu ZI, Spieler R (2006) Reliability and validity of a patient self-administered daily questionnaire to assess impact of oral mucositis (OM) on pain and daily functioning in patients undergoing autologous hemopoietic stem cell transplantation (HSCT). Bone Marrow Transplant 37(8): 393–401.

Streiner DL, Norman GR (1995) Health Measurement Scales: A Practical Guide to their Development and Use. 2nd edn. Oxford University Press Oxford, UK.

Sung L, Tomlinson GA, Greenberg ML, Koren G, Judd P, Ota S, Feldman BM (2007a) Serial controlled N-of-1 trials of topical vitamin E as prophylaxis for mucositis in children treated with hematopoietic cell transplantation (HSCT). Bone Marrow Transplant 37(8): 393–401.

Sung L, Tomlinson GA, Greenberg ML, Koren G, Judd P, Ota S, Feldman BM (2007b) Validation of the oral mucositis assessment scale in pediatric cancer. Pediatr Blood Cancer 49(2): 149–153.

Tomlinson D, Gibson F, Treister N, Baggett C, Judd P, Hendershot E, Maloney AM, Doyle J, Feldman B, Kwong K, Sung L (2009a) Understandability, content validity, and overall acceptability of the Children’s International Mucositis Evaluation Scale (ChIMES): child and parent reporting. J Pediatr Hematol Oncol 31(6): 416–423.

Tomlinson D, Gibson F, Treister N, Baggett C, Judd P, Hendershot E, Maloney AM, Doyle J, Feldman B, Kwong K, Sung L (2009b) Designing an oral mucositis assessment instrument for use in children: generating items using a nominal group technique. Support Care Cancer 17(5): 555–562.
Tomlinson D, Gibson F, Treister N, Hendershot E, Maloney A, Judd P, Doyle J, Baggott C, Feldman B, Sung L (2008a) Challenges of mucositis assessment in children: expert opinion. *Eur J Oncol Nurs* 12(5): 469–475.
Tomlinson D, Judd P, Hendershot E, Maloney AM, Sung L (2007) Measurement of oral mucositis in children: a review of the literature. *Support Care Cancer* 15(11): 1251–1258.
Tomlinson D, Judd P, Hendershot E, Maloney AM, Sung L (2008b) Establishing literature-based items for an oral mucositis assessment tool in children. *J Pediatr Oncol Nurs* 25(3): 139–147.

World Health Organization (1979) *Handbook for Reporting Results of Cancer Treatment*. WHO: Geneva, Switzerland.

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**APPENDIX 1**

**CHILDREN’S INTERNATIONAL MUCOSITIS EVALUATION SCALE (ChIMES)**

**PAIN**

1. Which of these faces best describes how much pain your child feels in their mouth or throat today? Circle one.

   0
   1
   2
   3
   4
   5
   0
   1
   2
   3
   4
   5

**FUNCTION**

2. Which of these faces shows how hard it is for your child to SWALLOW their saliva/spit today because of mouth or throat pain? Circle one.

   0
   1
   2
   3
   4
   5
   0
   1
   2
   3
   4
   5

3. Which of these faces shows how hard it is for your child to EAT today because of mouth or throat pain? Circle one.

   0
   1
   2
   3
   4
   5
   0
   1
   2
   3
   4
   5

4. Which of these faces shows how hard it is for your child to DRINK today because of mouth or throat pain? Circle one.

   0
   1
   2
   3
   4
   5
   0
   1
   2
   3
   4
   5

**PAIN MEDICATION**

5. Has your child taken medicine for any kind of pain today?

   □ Yes  □ No

   If yes, did your child need the medicine because they had mouth or throat pain?

   □ Yes  □ No

**APPEARANCE**

6. Please look in your child’s mouth. Can you see any mouth sores (ulcers)?

   □ Yes  □ No  □ Can’t tell

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**Figure A1.** Child International Mucositis Evaluation Scale (ChIMES).
## APPENDIX 2

### Table A1. Test-retest reliability of each ChIMES item measured on Days 13 and 14*

| Respondent type                  | ChIMES1 \( r \) (P-value) | ChIMES2 \( r \) (P-value) | ChIMES3 \( r \) (P-value) | ChIMES4 \( r \) (P-value) | ChIMES5 \( r \) (P-value) | ChIMES6 \( r \) (P-value) | ChIMES7 \( r \) (P-value) |
|----------------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
| **Parent proxy report**          |                             |                             |                             |                             |                             |                             |                             |
| Report no change, \( n = 53 \)   | 0.966 (<0.0001)             | 0.954 (<0.0001)             | 0.961 (<0.0001)             | 0.972 (<0.0001)             | 0.967 (<0.0001)             | 0.958 (<0.0001)             | 0.935 (<0.0001)             |
| All, \( n = 98 \)                | 0.921 (<0.0001)             | 0.931 (<0.0001)             | 0.925 (<0.0001)             | 0.928 (<0.0001)             | 0.921 (<0.0001)             | 0.898 (<0.0001)             | 0.862 (<0.0001)             |
| **Child self-report aged 12–18 years** |                             |                             |                             |                             |                             |                             |                             |
| Report no change, \( n = 35 \)   | 1.000 (<0.0001)             | 0.999 (<0.0001)             | 0.999 (<0.0001)             | 0.995 (<0.0001)             | 0.832 (<0.0001)             | 0.921 (<0.0001)             | 0.844 (<0.0001)             |
| All, \( n = 66 \)                | 0.877 (<0.0001)             | 0.776 (<0.0001)             | 0.854 (<0.0001)             | 0.819 (<0.0001)             | 0.727 (<0.0001)             | 0.780 (<0.0001)             | 0.782 (<0.0001)             |
| **Child self-report aged 8 to <12 years** |                             |                             |                             |                             |                             |                             |                             |
| Report no change, \( n = 11 \)   | 1.000 (<0.0001)             | 0.995 (<0.0001)             | 1.000 (<0.0001)             | 1.000 (<0.0001)             | 0.849 (0.0001)              | 0.826 (0.0003)              | 0.810 (0.003)               |
| All, \( n = 21 \)                | 0.909 (<0.0001)             | 0.908 (<0.0001)             | 0.923 (<0.0001)             | 0.871 (<0.0001)             | 0.906 (<0.0001)             | 0.868 (<0.0001)             | 0.750 (0.0003)              |

*Table represents Spearman’s correlation coefficients with \( P \) values in parentheses for each ChIMES item when measured on 2 consecutive days (Days 13 and 14). Results are presented for those who indicated no change in mucositis between Days 13 and 14 and for all respondents.