The Chinese version of the Quality of Life in Childhood Epilepsy Questionnaire-16-C (QOLCE-16-C): translation, validity, and reliability

Ping Tang1, Qunfeng Lu1*, Yinghui Wu4*, Lin Wang4, Wenjuan Tang1, Yan Jiang1, Liling Yang4, Jianlin Ji4, Xiaomin Sun2, Jingmin Sun3 and Jie Yang2

Abstract

Background: Epilepsy is one of the most common chronic neurological diseases that adversely impact the quality of life of patients and their families. The “Quality of Life of Childhood Epilepsy Questionnaire” (hereinafter referred to as “QOLCE-16”) is a 16-item measure that was designed to assess health-related quality of life (HRQOL) among children with epilepsy. The purpose of the study was to translate and evaluate the psychometric properties of the QOLCE-16.

Methods: The 10 steps of Principles of Good Practices for translation and cultural adaptation of measures were adopted to translate the QOLCE-16 into Chinese. After that, item analysis, floor effect and ceiling effect, internal consistency, test–retest reliabilities, content validity and construct validity were conducted to test its applicability in children with epilepsy in China. A total of 435 native Chinese-speaking parents with children who had epilepsy from one children’s hospital were invited to take part in the study, including a cognitive interview sample of 5 and a validation sample of 430.

Results: A total of 414 objects were enrolled in our study for psychometric testing. The results of the item analysis revealed QOLCE-16-C to have good discrimination, the floor effect and ceiling effect were 0.2% and 1.0% respectively, and each item was significantly related to the total scale ($P < 0.001$). The Cronbach’s $\alpha$ value was 0.938 and the test–retest reliability was 0.724. For validity, results showed that the QOLCE-16-C had good content validity. Exploratory factor analysis indicated it was reasonable that the QOLCE-16-C consists of four dimensions after rotation. Confirmatory factor analysis demonstrated good construct validity ($\chi^2/df=1.698$, GFI = 0.913, CFI = 0.974, RMSEA = 0.058).

Conclusion: The Chinese version of QOLCE-16-C appears to be a culturally appropriate, valid and reliable tool to assess the health-related quality of life of children with epilepsy in China.

Keywords: Epilepsy, Pediatrics, Health related quality of life, Translation, Factor analysis, Reliability and validity

Plain English summary

Epilepsy is one of the most common chronic neurological diseases. Due to its high prevalence and serious adverse effects, epilepsy has become a major public health concern. Health related quality of life (HRQOL) is a multidimensional concept, used to obtain a wider range of treatment and recovery information for patients in clinical practice. As such, the ability to precisely measure HRQOL becomes critical for children living with...
epilepsy. To date, several inventories have been established to evaluate the HRQOL of children with epilepsy, among them, QOLCE-16 not only has adequate measurement characteristics, but also possesses a small number of items, which making it suitable for the busy clinical work in China. Despite this, there is no Chinese version of the scale in China at present. In this study, we have translated the QOLCE-16 from English into Chinese. Followed by this translation is an assessment of the psychometric properties of the Chinese version of the Quality of Life in Childhood Epilepsy Questionnaire-16-C (QOLCE-16-C). This study indicates that the Chinese version of QOLCE-16-C appears to be a culturally appropriate, valid and reliable tool to assess the health related quality of life of children with epilepsy in China.

Background
Epilepsy is one of the most common chronic neurological diseases, which is characterized by abnormal brain discharge accompanied with unpredictable and recurrent seizures [1]. Studies have shown that there are roughly 9.84 million people with epilepsy in mainland China, which most of these being pediatric patients [2]. Children with epilepsy are experiencing issues not only associated with neurological deficits and functional limitations, but are also going through profound psychiatric and psychosocial influences compared to patients with other chronic diseases [3–5]. Moreover, children with epilepsy are limited in social activities, face future driving restrictions, and discrimination with social stigma, which may lead to a decrease in self-esteem and increased negative emotions (e.g., depression and social isolation), all of which could pose obstacles for improving their quality of life [6]. High prevalence and serious adverse effects have made epilepsy as one of the major public health concerns [7].

Considering epilepsy’s chronic nature and significant negative effects on multiple facts of children's lives, traditional treatments for seizure control no longer meet the current focus of long-term care for epilepsy. Thus, reducing seizure frequency and improving the quality of life should be regarded as the primary goals of antiepileptic treatments [8]. Health related quality of life (HRQOL) is a multidimensional concept, which is used to obtain a wider range of treatment and recovery information of patients in clinical practice [9–11]. Measurement of HRQOL in children with epilepsy helps to accurately reflect their physical, emotional, and psychosocial functions [12]. Hence, the ability to precisely measure HRQOL becomes critical for children living with epilepsy.

To date, several inventories have been established to evaluate the HRQOL of children with epilepsy, such as the Quality of Life in Epilepsy for Adolescents (QOLIE-AD-48) [13], the Quality of Life in Childhood Epilepsy Questionnaire (QOLCE-55) [14], and the Health-Related Quality of Life Measure for Children with Epilepsy (CHEQOL-25) [15]. However, the relatively lengthy items of these tools have cost reporters large amounts of time and patience to complete. Additionally, management burdens have been given to medical staff, which may limit its practical use in busy clinical research settings, especially China.

In order to minimalize the burden of respondents, Goodwin and other scholars recently developed a 16-item version measure (QOLCE-16) from the QOLCE-55 that allows for the capturing of HRQOL while maintaining the strong properties of the original tool based on the item response theory methods [8, 16]. The reliability and sensitivity of the QOLCE-16 was tested across age, sex, and time, which shows this short version is both time-saving and appropriate as measure for clinic [8].

Although QOLCE-16 is an excellent and practical inventory for China’s busy clinical and scientific environment, its use in China is currently blank due to the absence of a Chinese version. The purpose of this study was to first translate the QOLCE-16 from English into Chinese, followed by an assessment of the psychometric properties of the Chinese version of the Quality of Life in Childhood Epilepsy Questionnaire-16-C (QOLCE-16-C). Expectations for the Chinese version of the QOLCE-16-C were the following: (1) appropriate cultural equality; (2) high internal consistence; (3) good test–retest reliability; (4) strong structural validity.

Methods
This study consists of two phases: (1) translation of the QOLCE-16 to Chinese, followed by a cross-cultural adaptation among the parents with children with epilepsy; (2) testing of the psychometric properties of the QOLCE-16-C, including reliability and validity assessments.

Phase 1: translation and cross-cultural adaptation procedures
The goal of phase 1 was to produce a semantically equivalent, grammatically fluent, culturally compatible, and readily comprehensible QOLCE-16-C. Due to the lack of gold translation standards [17], we adopted the 10 steps of Principles of Good Practices for Translation and Cultural Adaptation of measures established by the International Society for Pharmacoeconomics and Outcomes Research (ISPOR) [18], which has commonly been used in large international researches, including the Patient-reported outcome measurement information system (PROMIS) [19], the World Health Organization...
Quality of Life (WHOQOL) [20], the Neuro-quality of life (Neuro-QoL) [21], and others [22, 23]. Figure 1 shows the specific translation steps.

**The forward and backward translation**

After receiving permission for translating from the development group of QOLCE-16 (Professor Kathy N. Speechley) through E-mails, the forward translation team (1 pediatric neurology nurse, 2 graduate students) translated the instrument independently. All three of them were bilingual with experience living in English-speaking countries for at least one year. The forward translated team that translated the English version were then reconciled into one through discussion among the three translators as well as the leader of the project. After that, another group of three bilingual translators (1 child neurologist, 1 child health doctor, 1 English-speaking translator) who were blind to the study objects and original QOLCE-16 conducted the back-translation. After discussion, an integrated and back-translated English version was obtained.

**Harmonization**

A 5-member harmonization team (1 linguistic specialist, 2 medical school teachers with experience in scale translation research, 2 pediatric nursing doctoral students) conducted an evaluation and comparison between the original scale, forward version and back-translated version to produce an agreement on a draft version.

**Cross-cultural adaption**

To evaluate the accuracy of the draft version with the original version, the cognitive interviewing technique was used in 5 parents whose child had epilepsy. Prior to the cognitive interview, an interview outline was developed and evaluated by experts to ensure its rationality. According to the interview outline, parents were asked about their understanding of the whole scale and the expression of specific items. A 10-point scale was used to measure how well parents understood the items. A rating of “1” meant that the item was difficult to answer, while a rating of “10” meant that the item was easy to answer. General probing and paraphrasing techniques were used for some of the items, which consist of asking respondents to explain the questions, requesting them to define meanings of words used in the questions, explaining their responses and identifying areas of the questionnaire that pose difficulty in understanding, interpretation or completion [24, 25].

**Expert review**

The Delphi Method was used during this procedure [26]. 9 experts (1 child neurologist, 1 child neurosurgeon, 1 child health doctor, 1 child rehabilitation doctor, 1 pediatrician in public health, 2 professors of nursing, 1 head nurse of neurosurgery, and 1 head nurse of neurology) were invited to make comparison among the above original scale, forward version, back-translated version, and cognitive interview integrated version through E-mail, and to form the final integrated version, the QOLCE-16-C.
Phase 2: QOLCE-16-C psychometric testing procedure

Study settings and subjects

Settings  Parents with children who had epilepsy from Shanghai Children’s hospital were invited to take part in the study. A convenience sampling method was adopted to recruit the eligible participants in the Neurology department’s inpatient and outpatient clinics.

Acceptance and row standard  The inclusion criteria were children who were (1) aged 4–18 years, (2) with a diagnosis of epilepsy, (3) with parents able to speak and read Chinese, and (4) willing to participate in this study and have their parent’s permission. Parents who have cognitive or mental impairment and refuse to participate in the investigation were excluded.

Informed consent  When children and their parents came to the clinic, the researchers explained the purpose of this study to the participants and asked them if they would be willing to participate in the survey. After obtaining the written informed consent of the participants, data were aggregated through self-filled questionnaires which were filled out in the doctor’s office.

Sample size  It is generally considered that two independent sample sizes of 200 were deemed accepted for exploratory factor analysis (EFA) and confirmatory factor analysis (CFA) [27]. Taking into account the possibility of missing data, a final total of 430 questionnaires were distributed from June to November 2020, of which 414 were effectively returned, with an effective response rate of 96.3% (16 scales were missing data). Of these, a sample of 203 respondents was recruited to conduct the EFA, and 211 were recruited for CFA. The data collection was divided into two steps: the data for EFA was firstly collected, and then to collected data for CFA. To assess the test–retest reliability, 40 parents were randomly selected to answer the QOLCE-16-C again 2–3 weeks after their first completement. The test–retest data was collected in two ways: paper questionnaires and electronic questionnaires filled out on the WeChat platform. Finally, 32 valid retest questionnaires were recovered.

Instruments

General demographic scale  The general demographic scale was used to obtain demographic data, including information about children with epilepsy and family information. Data related to children with epilepsy included gender, age, grade of study and information related to epilepsy disease. Family data included parents’ age, education background, family monthly income, payment methods for medical expenses.

Chinese version of the QOLCE-16  This is a short-form version of the QOLCE-55 [14] and provides an overall assessment of parent-reported HRQOL of children with epilepsy aged 4–18 years across four domains: cognitive (4 items), emotional (4 items), social (4 items), and physical functioning (4 items). The internal consistency reliability of the original QOLCE-16 was excellently high (α = 0.90) [16]. A 5-point Likert scale was used to calculate the scores for each domain and the scores for the total scale: 1 = very often, 2 = fairly often, 3 = sometimes, 4 = almost never, 5 = never [16]. The reverse code included the following items: item d of the emotional domain, and items a, b, c of the physical functioning domain. These items were then transformed as follows: response option 1 transformed to response option 5, response option 2 transformed to response option 4, response option 4 transformed to response option 2, and response option 5 transformed to response option 1. Linearly transform items from the pre-coded numeric values of items to a 0–100 point scale, with higher converted scores reflecting better quality of life. Responses were transformed such that: response option 1 = 0, response option 2 = 25, response option 3 = 50, response option 4 = 75, response option 5 = 100, presenting total scores take values of the unweighted mean of the four subscales from 0 (low HRQOL) to 100 (high HRQOL) [8, 14]. If more than one subscale is missing, the total score should be set to missing. It is worth noting that the scale also included the sixth option: 6 = not applicable, however this option was not incorporated into the scoring. If the answer to an item is “not applicable”, the item will be considered as missing data. The translation procedure of the Chinese version of the QOLCE-16 has been reported in phase 1.

Data analysis

IBM SPSS Statistics 23.0 and SPSS Amos24.0 (Armonk, NY, US) were used for analyzing the demographic characteristics and psychometric properties. Descriptive statistics were accepted to illustrate the participants.

Item analysis  In item analysis, the normality of data distribution was visually inspected. Floor and ceiling effects were respectively defined as the number of participants with the highest or lowest scores on a scale. A percentage of 15% or higher indicated poor performance in the questionnaire [28]. Independent-samples for the T test were chosen to calculate the critical ratio (CR) for every item, where a value above 3 was considered statistically significant [29]. After that, the correlation coefficient between each item and the total scale scores was checked, with a value greater than 0.4 considered to have a good correlation between the two [29].

Item analysis  In item analysis, the normality of data distribution was visually inspected. Floor and ceiling effects were respectively defined as the number of participants with the highest or lowest scores on a scale. A percentage of 15% or higher indicated poor performance in the questionnaire [28]. Independent-samples for the T test were chosen to calculate the critical ratio (CR) for every item, where a value above 3 was considered statistically significant [29]. After that, the correlation coefficient between each item and the total scale scores was checked, with a value greater than 0.4 considered to have a good correlation between the two [29].
Reliability  Reliability was tested by internal consistency and test–retest reliability. Cronbach’s α coefficient was used to assess the internal consistency where a value of α above 0.7 is considered as acceptable [30]. Interclass correlation coefficients (ICC) was used for test–retest reliability, in general, values between 0.70/0.75 and 0.9 and greater than 0.90 are indicative of good, and excellent reliability, respectively [31].

Content validity  Content validity index (CVI) was used to evaluate the content validity of the scale, including the scale level CVI (S-CVI) and the item level CVI (I-CVI). 9 experts (see above section on expert review process) were asked to score the whole scale and each item on clarity and relevancy on a 4-point Likert scale (0 = undesirable, 1 = somewhat desirable, 2 = desirable, 3 = fully desirable). The CVI was then calculated by the number of the evaluators gave fully desirable and desirable divided by the total number of questions. The cut-off of the S-CVI should be 0.9 or higher, and the I-CVI should be 0.78 or higher [32].

Construct validity  Although the QOLCE-16 has been proven to have good construct validity by its original author, EFA and CFA were conducted considering that the translated QOLCE-16-C was applied for the first time in Chinese children with epilepsy. First, EFA was performed to obtain the factor structure by using principal components analysis with varimax rotation [33]. Before the EFA, Kaiser–Meyer–Olkin’s (KMO) measure and Bartlett’s sphericity were chosen for verification, where a KMO index greater than 0.50 was considered eligible to conduct EFA [34]. Criteria used to determine the number of effective factors were: (1) eigenvalues greater than 1.0, (2) the percentage of total explained variance accounted for, and (3) items with loading greater than 40% in absolute value [33]. Second, CFA was conducted to verify the factor structure determined in the exploratory study. The maximum likelihood estimation method was used to estimate the parameters of the model. Items were first loaded onto 4 factors and then integrated into one factor. The model fit indices are deemed acceptable by $\chi^2/df < 3$, comparative fit index (CFI) > 0.90, goodness of fit index (GFI) > 0.90 and root-mean-square error of approximation (RMSEA) < 0.08 [35]. After that, the convergent validity of each dimension was tested by the values of the average variance extracted (AVE), where the value of AVE higher than 0.5 was considered acceptable, and discriminant validity was determined by comparing the square root value of AVE of each dimension with the correlation coefficient of a particular dimension with those of other dimensions [36].

Results  Phase 1: translation and cross-cultural adaptation

The forward and backward translation  The results of the three forward translators were able to reached a general consensus for translation, except for the phrase “health and life quality.” Additionally, there were objections to the translation of “directions,” as it contains both the direction and indication two kinds of meanings. However, since this item belonged to the cognitive dimension, it was more appropriate to translate it as an indication, as cognitive impairment is common in children with epilepsy. During the backward translation, we observed that most of the items and instructions were almost coincided with the original scale, possibly owing to majority being daily expressions with few abstract concepts. The integrated forward and backward version of the QOLCE-16 were delivered for further harmonization and review.

Harmonization  The 5-member harmonization team generally agreed with the forward and backward translation, but put some adjustments to better order items in the same dimension. For example, in the original scale, “Had difficulty following complex instructions” came before “health and life quality.” Additionally, there were objections to the translation of “directions,” as it contains both the direction and indication two kinds of meanings. However, since this item belonged to the cognitive dimension, it was more appropriate to translate it as an indication, as cognitive impairment is common in children with epilepsy. During the back translation, we observed that most of the items and instructions were almost coincided with the original scale, possibly owing to majority being daily expressions with few abstract concepts. The integrated forward and backward version of the QOLCE-16 were delivered for further harmonization and review.

Cross-cultural adaptation  Five native Chinese parents (one farther, four mothers) whose child had epilepsy were interviewed, ranging in age from 31 to 40 years, with educational levels covering junior high school education to doctoral degrees. Generally, participants reported that all the items and instructions were appropriate without any uncomfortableness. Two parents had differing opinions on one translation respectively: “instructions” was understood as “imperative commands” by one mother, whereas “supervision” was understood as “overprotective” by another mother. Considering that the two mothers were of a low educational level, we believe that it may be an isolated phenomenon. As such, we elected not to revise it and instead the results over to experts for review.
**Expert review**

A total of three rounds of expert reviews were conducted, and all nine experts gave feedback actively. The semantic meaning and content of the scale were examined according to the local Chinese culture. Beyond that, the uncertainties encountered in the translation and cultural adaption process were discussed in detail. The phrase “health and well-being” was eventually translated into “quality of life” in Chinese, as it was consistent with the content measured by the scale. The word “supervision” in the item “Need more supervision than other children in his/her age” was determined to be translated as “caring” for parents can better understand it. In addition, experts determined the expression of the answers to the scale, such as “All of the time” and “Very often” were translate as “always” in Chinese. The Chinese version of QOLCE-16 was obtained in Appendix 1.

**Phase 2: psychometric properties of the QOLCE-16-C**

**Demographic characteristics of the subjects**

From the total number of 430 parents, 414 participants from Shanghai Children’s hospital were recruited, excluded those for not filling out sufficient data. Among these 414 epilepsy children, 203 (49%) were boys, with the average age of 9.73 years old and a standard deviation of 3.31 years old, 170 (41.1%) were partial seizures, 173 (41.8%) were generalized seizures, 298 (63.0%) of them took only one or zero antiepileptic drug. Among these 414 respondents, 298 (72.0%) were mothers, 13 (3.1%) were fathers, 103 (24.8%) were grandparents, with age ranged from 27 to 80 (44.2 ± 12.52) years old, 246 (59.42%) of them had a bachelor degrees or above. Of those participants, 203 of them are for the EFA and 211 are for the CFA. The detailed demographic characteristics of the total subjects are presented in Table 1.

| Variable                                      | N (%)                            |
|-----------------------------------------------|----------------------------------|
| Information about epilepsy children           |                                  |
| Gender                                        |                                  |
| Male                                          | 203 (49.03%)                     |
| Female                                        | 211 (50.97%)                     |
| Mean age (SD)                                 | 9.73 (3.31)                      |
| Grade level                                   |                                  |
| Haven’t started school                        | 4 (1.0%)                         |
| Kindergarten                                  | 62 (14.98%)                      |
| Primary school                                | 235 (56.76%)                     |
| Secondary school                              | 91 (21.98%)                      |
| High school                                   | 14 (3.38%)                       |
| Special school                                | 8 (1.93%)                        |
| Epilepsy type                                 |                                  |
| Partial                                       | 170 (41.06%)                     |
| Generalized                                   | 173 (41.79%)                     |
| Don’t know                                    | 71 (17.15%)                      |
| Mean duration of epilepsy (SD)                | 3.95 (3.19)                      |
| Types of AEDs                                  |                                  |
| 0–1                                           | 298 (71.98%)                     |
| 2–3                                           | 113 (27.29%)                     |
| > 3                                           | 3 (0.72%)                        |
| Information about the respondents and families |                                  |
| Relationship to child                         |                                  |
| Farther                                       | 13 (3.14%)                       |
| Mother                                        | 298 (71.98%)                     |
| Grandparents                                  | 103 (24.88%)                     |
| Mean age (years)                              | 44.24 (12.52)                    |
| Education level                               |                                  |
| Primary school                                | 4 (0.97%)                        |
| Secondary school                              | 56 (13.53%)                      |
| High school/Technical school                  | 108 (26.09%)                     |
| Undergraduate/Junior College                  | 153 (36.96%)                     |
| Master or above                               | 93 (22.46%)                      |
| Average monthly household income              |                                  |
| 2000–5000                                     | 16 (3.86%)                       |
| 5000–8000                                     | 67 (16.18%)                      |
| 8000–10,000                                   | 120 (28.99%)                     |
| > 10,000                                      | 63 (50.97%)                      |

**Item analysis**

A total of 414 samples were included for item analysis, and there were no missing data for each item. The QOLCE-16-C showed a 0.2% floor effect and 1.0% ceiling effect, which means of no floor or ceiling effect in the Chinese version of QOLCE-16. The item analysis results showed that the CR coefficients of all items were higher than 3 ($P<0.001$). The scores of all items are significantly related to the total scores (Table 2).

**Reliability**

The results shown that the Cronbach’s $\alpha$ coefficients for the total and four dimensions of the QOLCE-16-C were 0.938, 0.945, 0.814, 0.922, and 0.891 respectively. There were 32 people completed the retest survey. The ICC for the total and each dimension were 0.724, 0.716, 0.615, 0.650, and 0.672 respectively (Table 3).

**Validity**

**Content validity** The positive coefficients of the 9 experts were 100%. The S-CVI and the I-CVI was 1, except for item 5, which was 0.89, indicating that the importance of
all items in the QOLCE-16-C has reached an expert consensus.

Construct validity The exploratory factor analysis shown that the KMO value was 0.928, Bartlett’s test \(\chi^2 = 5514.698, P < 0.01\), which was suitable for exploratory factor analysis. Principal component analysis method was used to extract 4 common factors. The characteristic values after the axis were 3.573, 3.145, 3.131, 2.660, all greater than 1. Its cumulative variance contribution rate was 78.183%, which meets the requirement that the cumulative variance contribution rate to be at least 40% [33]. The factor loading of each item was between 0.621 and 0.877. Table 4 shows more details.

The CFA was performed to test the fitness of the total scale. The construct of the total scale was acceptable for \(\chi^2/df = 1.698\), GFI = 0.913, CFI = 0.974, RMSEA = 0.058 (Table 5). The value of AVE of each dimension, the square root of every AVE value, and correlation coefficients between dimensions are shown in the Table 6. It was revealed that the square root of AVE was significantly higher than the dimensions of other correlated coefficients, thus suggesting acceptable discrimination validity.

Discussion
To the best of our knowledge, this study presents the first attempt to translate the QOLCE-16 into Chinese, tests its validity and reliability among Chinese children with epilepsy.

The steps of cross-cultural translation are rigorous and have firm basis. The 10-step translation method of the guidelines for the Principles of Good Practices for translation and cultural adaptation was used to translate QOLCE-16, which provides effective guidance for cross-cultural translation. Few items were adjusted during the process based on the results of parents’ cognitive interviews and the opinions given by expert consultants, such as translating “supervision” to “caring” for better understanding. In addition, the order in which items were placed had been adjusted according...
to the cognitive habits of Chinese. Consequently, the translated scale can convey the meaning smoothly in the semantics and is also acceptable in the culture.

The QOLCE-16-C had no floor or ceiling effects, with items being well identified and strongly correlated to the total scale. The results of item analysis illustrated that the CR values of each item were greater than 3, which meets the requirement of CR values being greater than 3 and statistically significant [29]. Moreover, there was a strong correlation between the scores of each item and the total scores, indicating that all items were valid.

Internal consistency and test–retest reliability of the QOLCE-16-C are acceptable. The Cronbach’s α coefficients for each dimension was higher than 0.7, and the cognitive and social dimensions were higher, 0.945 and 0.922 respectively, which was consistent with the results of Goodwin (value of 0.92 and 0.85) [16]. The test–retest reliability was completed among 32 parents, and the results shown that the ICC between the two tests ranged between 0.615 and 0.716. The lower correlation may be related to the length of the interval in our study (2–3 weeks). In addition, the way of those retest data been collected may also have contributed to the low correlation, as more than half of the retest data was collected through WeChat electronic questionnaires.

Content validity and construct validity are reasonable. The S-CVI in QOLCE-16-C reached 1, and the I-CVI was 0.89–1, indicating good expert content validity. In structural validity, EFA produced four common factors, which were highly consistent with the results of the original scale. The results of CFA were acceptable for $\chi^2/df = 1.698$, $GFI = 0.913$, $CFI = 0.974$, RMSEA = 0.058. In addition, the QOLCE-16-C also showed good discriminative validity. In future studies, the measurement equivalence of the QOLCE-16-C should be further verified by variety of groups, such as differing in age, gender or time [8].

Finally, we have to reiterate the purpose of the development of QOLCE-16 proposed by the original author Goodwin: to simplify QOLCE-55 and to provide an easy-to-use measurement tool for clinic. The QOLCE-55 is a simplified version of the original 76-item scale and provides an overall assessment of the HRQOL of 4–18 years old children with epilepsy reported by parents, which covering the four genres of HRQOL: cognitive (22 items), emotional (17 items), social (7 items), and physical (9 items) [14]. The scoring method of QOLCE-55 is consistent with QOLCE-16, and it also has excellent internal consistency reliability ($\alpha = 0.96$). Therefore, QOLCE-55 is a preferred measure in cases where specific domain information or a more comprehensive HRQOL assessment is required. In contrast, QOLCE-16 is more suitable for focusing on aggregate summary scores [16].

There are several limitations of this study need to take further consideration. (1) Convenience sampling survey conducted in only one children’s hospital, which may have some impacts on the scale’s generalizability. (2) The sample size of the 5 cognitive interviewees we included were relatively small, compared to the recommended acceptable range in the literature being 10–15 subjects [37]. This may result in an unreasonable translation and expression of items. (3) Convergent and divergent validity were performed on 4 dimensions, but did not on the total QOLCE-16-C, which would have required a comparison of the QOLCE-16-C scores with other validated measures. Future studies need to be carried out in different groups (e.g., age, sex, and time) in order to further confirm its applicability in Chinese children with epilepsy.

### Conclusions

In conclusion, this study conducted a translation of QOLCE-16 in China, along with its reliability and validity tests in Chinese children with epilepsy. The contents and items of the QOLCE-16-C are simple and easy to understand. This simplicity was largely due to the small items’ effectiveness in reducing time and effort in two key areas—answering and reporting. After verification, the QOLCE-16-C has been proved to have good reliability and validity, and is suitable for the evaluation of health-related quality of life of children with epilepsy in China. Future studies need to be carried out in different groups to make further confirmation on its applicability in Chinese children with epilepsy.

### Table 5

| $\chi^2$ | df | $P$ | $\chi^2/df$ | CFI | GFI | RMSEA |
|---------|----|-----|------------|-----|-----|-------|
| 166.378 | 98 | 0.000 | 1.698 | 0.974 | 0.913 | 0.058 |

### Table 6

| AVE | Cognitive | Emotional | Social | Physical |
|-----|-----------|-----------|--------|----------|
| Cognitive | 0.825 | 0.908 | 0.677 | 0.839 |
| Emotional | 0.458 | 0.503 | 0.654 | 0.745 | 0.845 |

The bold parts are the square root of AVE.
Appendix 1: Chinese version of QOLCE-16

问卷说明

尊敬的家长:

您好！以下问题有关您孩子的总体健康情况，请勾选出合适的答案。某些问题看起来可能比较相似，但其实各不相同。还有些问题可能您的孩子并没有相关表现，请尽量回答每一题。因为对我们而言，了解您的孩子什么时候还不存在这些问题非常重要。答案没有对错之分。如果您不确定如何回答某个问题，请尽量给出您自己认为最贴切的回答。

第一部分：您的孩子认知功能

以下问题涉及孩子在专注力、记忆力和语言能力方面存在的一些问题。

与同龄孩子相比，您的孩子在**过去4周内**出现以下情况的频率程度：

|  | 总是 | 经常 | 有时 | 几乎不 | 从不 | 不适用 |
|---|---|---|---|---|---|---|
| a. 在理解指令时存在困难？ |  |  |  |  |  |  |
| b. 在遵循简单指令时存在困难？ |  |  |  |  |  |  |
| c. 在遵循复杂指令时存在困难？ |  |  |  |  |  |  |
| d. 难以记住别人告诉过他/她的事情？ |  |  |  |  |  |  |

第二部分：您孩子的情感功能

下列条目描述了您孩子可能存在的总体感受。

您认为您的孩子，在**过去4周内**有多少时间存在下列现象：

|  | 总是 | 经常 | 有时 | 偶尔 | 从不 | 不适用 |
|---|---|---|---|---|---|---|
| a. 感觉没有人理解他/她？ |  |  |  |  |  |  |
| b. 感觉情绪低落或沮丧？ |  |  |  |  |  |  |
| c. 感觉挫折？ |  |  |  |  |  |  |
| d. 感觉自信？ |  |  |  |  |  |  |
第三部分：您孩子的社交功能

以下是一些涉及孩子社会交往和社会活动的描述。

a. 在过去4周内，与同龄孩子相比，您的孩子由于癫痫或与癫痫相关的问题，他/她的社交活动受到了限制吗？程度如何？

| □ | □ | □ | □ | □ | □ |
|---|---|---|---|---|---|
| 是的，严重 | 是的，有 | 是的，轻度 | 是的，但 | 不，没有受限 | 不适用 |
| 重受限 | 些受限 | 微受限 | 少受限 | 受限 |

在过去4周内，由于癫痫您孩子出现以下状况的频数程度：

b. 影响了他/她在学校或工作中的社交互动？

c. 导致他/她被孤立？

d. 导致他/很难维持友谊？

第四部分：您孩子的身体机能

以下问题有关您孩子可能进行的体育活动。

在过去的4周的日常活动中，您的孩子在多大频率上：

| □ | □ | □ | □ | □ | □ |
|---|---|---|---|---|---|
| 总是 | 经常 | 有时 | 几乎不 | 从不 | 不适用 |
| a. 像同龄孩子一样在户外自由玩耍？ | | | | | |
| b. 像同龄孩子一样从事体育活动？ | | | | | |
| c. 像同龄孩子一样在室内自由玩耍？ | | | | | |
| d. 比其他同龄孩子需要更多的看护？ | | | | | |

Abbreviations
HRQOL: Health related quality of life; QOLIE-AD-48: Quality of Life in Epilepsy for Adolescents; QOLCE-55: Quality of Life in Childhood Epilepsy Questionnaire; CHEQOL-25: Quality of Life Measure for Children with Epilepsy; QOLCE-16: Quality of Life Childhood Epilepsy Questionnaire; QOLCE-16-C: Chinese version of the Quality of Life in Childhood Epilepsy Questionnaire; ISPOR: International Society for Pharmacoeconomics and Outcomes Research; PROMIS: Patient-reported outcome measurement information system; WHOQOL: World Health Organization Quality of Life; Neuro-QoL: Neuro-quality of life; CR: Critical ratio; ICC: Interclass correlation coefficients; CVI: Content validity index; S-CVI: Scale level of content validity index; I-CVI: Item level of content validity index; EFA: Exploratory factor analysis; CFA: Confirmatory factor analysis; KMO: Kaiser–Meyer–Olkin; CFI: Comparative fit index; GFI: Goodness of fit index; RMSEA: Root-mean-square error of approximation; AVE: Average variance extracted.
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Authors' contributions
Conceived, designed and directed the study: PT, QL, YW, YJ, LY and JJ. Performed the experiments: PT, QL, L.Y., J.J., X.S., J.S. and J.Y. Analyzed the data: PT, QL, YW and LW. Wrote the article: PT, QL, LW, YJ, LY, JY and JJ. Critical revision of the manuscript: PT, QL, YW, LW, XS and JS. All authors read and approved the final manuscript.

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Availability of data and materials
The datasets generated and/or analyzed during the current study are not publicly available to preserve the privacy of the participants but are available from the corresponding author upon reasonable request.

Declarations
Ethical approval and consent to participate
This study protocol was approved by the Ethics Committee of the Shanghai Children's hospital, and informed consent was obtained before the study began, the ethical approval number is 2020R053-E01. Informed consent and assent were obtained.

Consent for publication
Not application.

Competing interests
The authors declare that they have no competing interests.

Author details
1 Department of Nursing, Shanghai Children's Hospital, Shanghai Jiao Tong University, Shanghai 200062, People's Republic of China. 2 Department of Neurology, Shanghai Children's Hospital, Shanghai Jiao Tong University, Shanghai 200062, People's Republic of China. 3 Department of Outpatient, Shanghai Children's Hospital, Shanghai Jiao Tong University, Shanghai 200062, People's Republic of China. 4 School of Nursing, Shanghai Jiao Tong University, Shanghai 200025, People's Republic of China.

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