Cystic Fibrosis in Adults: Short-Term and Long-Term Reproducibility of the Brody Score for Lung Morphology in Low-Dose MDCT Scans

Cystische Fibrose (CF) bei Erwachsenen: Kurz- und Langzeit-Reproduzierbarkeit des Brody-Scores bei Niedrig-Dosis-MDCT-Aufnahmen

Abstract

Purpose: The semi-quantitative Brody score measures the severity of cystic fibrosis (CF)-related lung disease. We investigated the short-term (28–60 days) and long-term (2–7 years) intra- and inter-observer reproducibility of the Brody score in low-dose multidetector row computed tomography examinations performed in inspiration (LDCTs) of adult CF patients.

Materials and Methods: Composite Brody scores and respective underlying bronchiectasis, mucus plugging, peribronchial thickening, parenchymal opacity, and hyperinflation subscores were evaluated twice (time interval, 1–84 months) by each of 3 independent radiologists (1–20 years of professional diagnostic radiology experience) in LDCTs (4–64 rows, 120 kVp, 10–15 mAs/slice, CTDIw approx. 1.0 mGy, effective dose approx. 0.5 mSv) of 15 adult patients with CF-related lung disease (8 female, 7 male, age, 18–50 years, mean, 33 years).

Results: The average reproducibility of the Brody score was within +/- 7% (range, 2–30%) between radiologists, and +/- 6% (3–12%) within radiologists (short-term, 28–60 days, 4%, 0–12%, long-term, 2–7 years, 12%, 1–36%). For the different subscores, the reproducibility was within +/- 25% (15–41%) between radiologists and +/- 23% (12–46%) within radiologists.

Conclusion: The Brody score shows high average inter-observer reproducibility in LDCTs of adult CF patients. The Brody score also demonstrates high average intra-observer reproducibility if subsequent assessments are made within 28–61 days. With time intervals of 2–7 years between subsequent evaluations, however, intra-observer reproducibility decreases. Respective subscores each demonstrate lower intra- and inter-observer reproducibility than does the composite Brody score.
Introduction

Cystic fibrosis (CF) is the most frequent life-shortening autosomal recessive disease among the Caucasian population in Europe, with an incidence about 1 in 2,500 live births and a carrier frequency of about 4–5% [1–3]. Mutations in a single large gene on chromosome 7 cause dysfunction of cAMP-dependent chloride channels in exocrine tissues, particularly in the lung and pancreas [3, 4]. Due to improved health care, life expectancy among CF patients has continuously increased, from about 25 years in 1985 to 37 years in 2008 [5].

There is no generally accepted gold standard for imaging CF-related lung disease in adult patients. However, some clinical institutions have established follow-up examinations at regular intervals that include computed tomography (CT) imaging [6]. In our institution, adult CF patients over the age of 18 years are regularly assessed by means of lung function tests once every 6 months and by low-dose-CT of the chest (LDCT) for lung morphology once every four years. Computed tomography (CT) has become the gold standard for imaging CF-related lung disease, with lung morphology appearing to be complementary to lung function tests (LFTs) in at least 50% of CF patients. Chest radiographs and LFTs alone fail to recognize structural lung alterations caused by CF [7]. MRI has not been widely applied to evaluate normal or mildly damaged lung structure in CF patients. Despite radiation exposure, the risk of CT-induced mortality is likely to be minimal in CF patients when compared to the benefit of a more accurate diagnosis [7]. The ACE systems of modern CT scanners significantly reduce dose exposure [8, 9]. Also, low-dose multidetector row CT of the lung (LDCT) shows high morphologic accuracy in non-malignant lung disease [10].

Different high-resolution (HR) CT-based scoring systems, with a range of morphologic parameters and different weighting of results, have been designed to support the evaluation of structural lung damage in CF patients. For some scoring systems, HRCT findings have been correlated with LFTs [11, 12]. A modification of the Bhalla score for structural lung alteration in CF has previously been applied in an adult population [2]. Brody and co-workers have developed a scoring system for HRCT scans obtained in both inspiration and expiration in children suffering from CF [13], which demonstrates both high reproducibility and accuracy [12, 14]. Since many CF patients live through adulthood, it appears necessary for optimal radiological patient management to extend CT scoring systems for children with CF to include adults with CF-associated lung disease [15] who underwent LDCT in inspiration.

Since the reproducibility of scoring results in the repetitive evaluation of the same chest CT images is a key feature of clinical reliability of any particular CT scoring system, we conducted an intra- and interobserver reproducibility study of the Brody scoring system in a selected set of LDCTs of the chest performed in adult CF patients. We determined the short-term reproducibility (28–60 days) of the Brody score among second-year radiology residents, to understand how firmly its different features would be integrated in new clinical knowledge. We also assessed the long-term (at least two years) reproducibility of the Brody score in an attending radiologist with 10 years of post-fellowship chest radiology experience, since time intervals between follow-up CT scans of the chest in CF patients are often in the range of several years [2, 6]. It has previously been established that the short-term reproducibility of the Brody score is high among experienced chest radiologists [12, 14].

We hypothesized that the intra-observer reproducibility of the Brody score and its different subscores in adult patients with CF who undergo LDCT of the chest in inspiration would be within +/-10% of the average score, both with a short time interval of 28–60 days for second-year radiology residents and with a long time interval of at least 2 years for an attending radiologist. Similarly, we hypothesized that the inter-observer reproducibility of the Brody score and its different subscores in adult patients with CF who undergo LDCT of the chest in inspiration would be within +/-10% of the average score between second-year radiology residents and an attending radiologist.

Materials and Methods

Patients

This retrospective analysis was based on LDCT examinations of the chest performed between February 2006 and October 2010, in 15 consecutive adult CF patients (male, n = 7; female n = 8; age range, 18–50 years, average, 33 years), each of whom had only had one LDCT examination at our institution. The clinical indication for LDCT was to assess the current lung morphology status. Institutional policy for adult CF patients over the age of 18 years includes the assessment of lung morphology status by means of LDCT once every four years. Additional MDCT scans are acquired when there is clinical suspicion or evidence of aggravation of CF-related lung disease.

Ethical Issues

Institutional ethics committee approval was obtained for the retrospective analysis of data previously obtained for individual clinical treatment of CF patients. Data included in this study were evaluated and presented in accordance with the World Medical Association (WMA) Declaration of Helsinki Ethical Principles for Medical Research Involving Human Subjects, as last amended by the 59th WMA General Assembly, Seoul, Korea, October 2008.

LDCT Protocol

LDCT of the chest was invariably performed in inspiration, at a tube charge of 120 KVP, with the lowest technically achievable tube current that would still generate images of diagnostic value and left no option of further dose reduction by means of dose modulation programs.

Through January 2007, examinations were conducted on a 4-row MDCT scanner (Mx 8000, Philips Medical Systems, Hamburg,
Results

The respective median time intervals and ranges between the first and second LDCT readings were 4.7 years (2.0 – 7.0) for R1, 42 days (32 – 60) for R2, and 37 days (28 – 47) for R3.

Brody Score (Total Score)

Among the 15 patients, the average Brody scores between the three independent readers ranged from 20.0 to 132.7 points (median, 58.4 points, mean, 68.0 points, possible range, 0 – 207.0 points). The individual Brody scores ranged from 10.0 – 146.5 points (median, 63.0 points, mean, 70.7 points) for R1, 25.0 – 133.5 points (median, 57.9 points, mean, 66.4 points) for R2, and 22.0 – 134.5 points (median, 59.9 points, mean, 66.9 points) for R3.

The average deviation of all individual Brody scores from the mean value of all readers was 4.9 points (7 %), with a range of 1.5 – 10.3 points (2 – 30 %). The mean deviation of Brody scores from individual respective average values was 8.7 points (12 %, range, 0.5 – 26.0 points, 1 – 36 %) for R1, 3.1 points (5 %, range, 0 – 6.3 points, 0 – 12 %) for R2, and 2.2 points (3 %, range, 0 – 7.0 points, 0 – 9 %) for R3. Findings implied an average intra-individual reproducibility of Brody scoring results within about 7 % of the individual mean score, with greater deviations in the long-term re-assessment by the most experienced reader. There was no evidence of skewing of the average deviation toward lower or higher average values of the Brody score (Fig. 1a – 3).

Bronchiectasis Subscore

The average deviation of individual bronchiectasis sub scores (possible range, 0 – 72.0 points) from the mean value of all readers was 4.2 points (16 %), with a range of 1.2 – 11.7 (8 – 43 %). The mean deviation of bronchiectasis sub scores from individual respective average values was 7.9 points (25 %, range, 0.5 – 17.0 points, 2 – 67 %) for R1, 3.7 points (16 %, range, 0.5 – 21.0 points, 1 – 57 %) for R2, and 1.2 points (5 %, range, 0 – 4.0 points, 0 – 20 %) for R3, implying an average intra-individual reproducibility of bronchiectasis sub scores results within about 15 % of the individual mean score, with greater deviations for long-term re-assessment by the attending radiologist (Fig. 1b, 3).

Mucus Plugging Subscore

The average deviation of all individual mucus plugging sub scores (possible range, 0 – 36.0 points) from the mean value of all readers was 1.9 points (19 %), with a range of 0.3 – 4.2 points (2 – 133 %). The mean deviation of mucus plugging sub scores from individual respective average values was 2.1 points (18 %, range, 0 – 6.0 points, 0 – 67 %) for R1, 1.4 points (16 %, range, 0 – 5.0 points, 0 – 67 %) for R2, and 1.1 points (11 %, range, 0 – 6.0 points, 0 – 100 %) for R3, implying an average intra-individual reproducibility of mucus plugging sub score results within about 12 % of the individual mean score, with similar deviations for short-term and long-term re-assessment (Fig. 1c, 3).

Peribronchial Thickening Subscore

The average deviation of all individual peribronchial thickening sub scores (possible range, 0 – 54.0 points) from the mean value of all readers was 2.7 points (15 %), with a range of 0.3 – 7.0 points (3 – 38 %). The mean deviation of peribronchial thickening sub scores from individual respective average values was 4.7 points (22 %, range, 0 – 12.0 points, 0 – 67 %) for R1, 2.4 points (14 %, range, 0 – 10.3 points, 0 – 36 %) for R2, and 0.67 points (4 %, range,
0 – 2.8 points, 0 – 13 %) for R3, implying an average intra-individual reproducibility of mucus plugging subscore results within about 13 % of the individual mean score, with greater deviations for long-term re-assessment by the attending radiologist (Fig. 1 d, 3).

**Parenchymal Opacity Subscore**
The average deviation of all individual parenchymal opacity subscores (possible range, 0 – 54.0 points) from the mean value of all readers was 1.8 points (41 %), with a range of 0 – 5.2 points (0 – 111 %). The mean deviation of parenchymal opacity subscores from individual respective average values was 2.6 points (86 %, range, 0 – 13.0 points, 0 – 200 %) for R1, 1.5 points (35 %, range, 0 – 5.0 points, 0 – 200 %) for R2, and 0.9 points (17 %, range, 0 – 7.0 points, 0 – 93 %) for R3, implying an average intra-individual reproducibility of parenchymal opacity subscore results within about 46 % of the individual mean score, with similar deviations for short-term and long-term re-assessment (Fig. 1e).
Hyperinflation Subscore

The average deviation of all individual hyperinflation subscores (possible range, 0 – 27.0 points) from the mean value of all readers was 3.1 points (37 %), with a range of 0.9 – 7.4 points (4 – 71 %). The mean deviation of hyperinflation subscores from individual respective average values was 2.7 points (62 %, range, 0 – 8.0 points, 0 – 200 %) for R1, 1.2 points (11 %, range, 0 – 5.5 points, 0 to 71 %) for R2, and 1.4 points (14 %, range, 0 – 10.5 points, 0 – 67 %) for R3, implying an average intra-individual reproducibility of hyperinflation score results within about 29 % of the individual mean score (Fig. 1f), with greater deviations for long-term re-assessment by the attending radiologist (Fig. 1f).

Discussion

In a retrospective analysis, we applied the Brody score to assess lung morphology in adult CF patients, based on LDCT scans of the lung performed in inspiration, and tested for the reproducibility of scoring results. The Brody score is a weighted composite...
score that was developed to describe and quantify lung morphology in children suffering from CF based on incremental HRCT scans of the lung obtained in both inspiration and expiration. The Brody score sums up its five different subscores for bronchiectasis, mucus plugging, peribronchial thickening, parenchymal opacity, and hyperinflation [13]. Reproducibility was tested among two second-year radiology residents, with short-term intervals of 28–60 days between subsequent assessments of the same LDCT scan, and an attending radiologist with 10 years of post-fellowship clinical chest radiology experience who re-assessed all LDCT scans in this study after long-term intervals of 2–7 years.

The respective mean, median, and range values for the Brody score were similar between readers. The range covered both very minor and severe lung involvement with CF-related disease, leaving out only the most severe part of the possible range. The findings imply that, overall, the different radiological features underlying the Brody scoring system are easy to understand, recognize, and weigh in their respective severity even with relatively little chest radiology experience. These findings expand previous experience which was largely based on expert readings of chest images of CF patients [12–14, 17].

For the Brody score as the weighted composite score sum of its different subscores, the study hypothesis of inter- and intra-observer deviations of 10% or less from the respective averages was met by the average of our findings, and, particularly, by the second-year radiologists who re-assessed LDCTs within 28–60 days. The latter finding corroborates previous reports of high short-term reproducibility of different HRCT scoring systems for CF patients of different ages among expert readers [12–14, 17].

The findings appear to expand previous knowledge to include a more general perception of the Brody score as a tool that reproducibly assesses lung morphology changes not only among pediatric patients in the hand of expert readers [12–14], but also among adult CF patients in the hands of radiology residents, as in this study. However, high reproducibility only seems to hold for short-term re-assessments. In fact, our finding of decreased reproducibility of Brody score results in the hands of an experienced chest radiologist after time intervals of 2–7 years, when compared with short-term reproducibility among radiology residents, implies that either memory effects play a role in the short-term reproduction of scoring results or visual perception may change over time. From a clinical point of view, it appears to be better to review the previous chest CT along with the current one and perhaps newly apply the Brody scoring system to both than to score only the current scan and rely on the old Brody scoring results of the previous one.

For the individual subscores, the study hypothesis of an inter- and intra-observer deviation of 10% or less from the respective averages was not met. Rather, individual sub-scoring results deviated by as much as 200 percent from the average. This finding was more pronounced among subscores with smaller score point ranges, i.e., mucus plugging and hyperinflation, and among subscores that appear to be more difficult to assess by visual means, i.e., parenchymal opacity and hyperinflation. Among the former, the lower point range increases both the susceptibility to small changes and the relative impact of deviations between subsequent assessments. Among the latter, visual impressions of lung parenchymal density and its distribution could be a source of uncertainty. When two neighboring areas of lung parenchyma show with different densities, the first could be hyperinflated while the second is normal, or the first could be normal while the second demonstrates ground glass opacity of some origin. Possible solutions would include the addition of an expiratory CT scan of the chest to more easily recognize areas of hyperinflation by their relative lack of volume change when compared with normal lung, or the addition of CT density measurements in areas of uncertainty. While the former would expose the patient to additional radiation, potentially at uncertain gain in the individual case, the latter would require lists of CT density value ranges that would be accepted as normal, too high, or too low for CF patients.

Limitations of our retrospective reproducibility study include the lack of experienced readers reassessing CT examinations after a short time interval, the restriction of experience at long time intervals to one experienced reader only, and the relatively low number of patients included. Since previous studies were based on the short-term reproducibility of CT scoring results in CF patients among experienced readers, it did not appear to add crucial knowledge to the field to obtain those results in addition. Among readers with brief postgraduate medical experience, it would not have been possible by design to perform a reproducibility study with intervals of several years between individual readings. Therefore, long-term reproducibility results could only have been obtained for readers with more extensive post-graduate chest radiology experience. In our department, such long-term experience with radiological examinations in CF patients was restricted to only one attending radiologist.

In conclusion, the Brody score, as a weighted composite score that describes and quantifies the respective presence, location, and extent of CF-related changes in lung morphology as detected by means of CT, appears to be reproducible within about 10% of an average value between different radiologists, both with a short time interval of 28 to 60 days, and with a long time interval of 2 to 7 years. However, its individual subscores for bronchiectasis, mucus plugging, peribronchial thickening, parenchymal opacity, and hyperinflation are less reproducible, with values exceeding 10%. Overall, the long-term reproducibility was not as good as the short-term reproducibility. For clinical practice, it appears advisable to review both the previous chest CT and the associated Brody score along with the new chest CT and new Brody score and perhaps newly apply the Brody scoring system to both.

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