Initial Experience of a Digital Training Resource for Modified Rankin Scale Assessment in Clinical Trials

Terence J. Quinn, MRCP; Kennedy R. Lees, MD; Hans-Goran Hardemark, PhD; Jesse Dawson, MRCP; Matthew R. Walters, MD

Background and Purpose—The modified Rankin Scale (mRS) is the preferred measure of disability in cerebrovascular clinical trials, but its value is restricted by interobserver variability. Poor reliability reduces the statistical power of clinical trials and leads to underestimation of effect size. Strategies to improve mRS grading are required. Video training has previously improved application of the National Institutes of Health Stroke Scale in clinical research. We developed an mRS training resource in an attempt to minimize interobserver variability.

Methods—We produced a complete training resource comprising an instructional DVD with accompanying written materials and assessment recordings of patient interviews. Formal assessment of training involved grading of real-life cases. Results of initial training and recertification were collected centrally and scored.

Results—Data from 1564 assessments are presented. The majority of assessors were participating in 2 large prospective clinical stroke trials. Assessors represented a mixed group of disciplines and nationalities. After training, most trainees (90%) achieved certification in mRS assessment. The majority (85%) of investigators who did not reach an acceptable score on initial testing achieved certification after further exposure to the package.

Conclusions—Mass training in mRS assessment for clinical trials is possible. We outline the development of a video-based training package, including technical issues, patient selection procedures, and methods of scoring and assessment. Certification results suggest that use of the resource can improve mRS grading. Acceptability of the training has been demonstrated by its successful use in 2 international acute stroke trials, SAINT 1 and CHANT. (Stroke. 2007;38:2257-2261.)

Key Words: cerebrovascular accident ■ medical education ■ modified Rankin Scale ■ outcome assessment ■ stroke treatment ■ video recording
without precedent. A well-validated video-based training program and certification procedure exist for the National Institutes of Health Stroke Scale (NIHSS), and it is now a requirement that clinical trial investigators complete this training and undergo certification.13 The strengths of the NIHSS program, in particular its mix of didactic teaching, video explanation, and assessment procedure, could be easily applied to the mRS.

We developed a training DVD and accompanying explanatory booklet, which include recordings of real Rankin assessments, certification cases, and further recertification cases. This has been used successfully in 2 large-scale clinical stroke trials.14,15 Although brief reference to certification scoring has been made in a previous review,16 to date there has been no detailed description of the training package, its development, and the initial experience of its use. We present this here along with the results of the certification program.

### Materials and Methods

#### Development of the Training Package: Audiovisual Issues

A criticism of the early NIHSS video training was poor image quality, an important consideration, as subtle clinical signs may have a substantial effect on final scoring.16 In an interview-based assessment such as the mRS, high-quality sound recording is of equal importance. We enlisted expert technical help (Media Services Department, University of Glasgow). A recording studio was used and, where patient disability made travel to the studio inappropriate, the Media Services mobile crew filmed in the hospital.

We chose a DVD-based format. This reflects the gradual replacement of conventional VHS recordings with DVD and the improved clarity of sound and vision afforded by it. Digital recording ensures optimal quality, even after mass reproduction. Finally, this allows for ease of transfer to a Web-based server. Internet-based NIHSS training has facilitated cost-effective global dissemination of the training package.

#### Patient Selection

The stroke liaison team based in the Western Infirmary, Glasgow, selected suitable patients from recent admissions to the acute stroke unit or local inpatient rehabilitation facility. The stroke unit accepts all patients presenting within 72 hours of onset of suspected stroke, irrespective of age or severity of the neurologic deficit; thus, a cohort of patients with varying degrees of disability and background comorbidity was available. Our intention was to include at least 1 patient from each potential mRS category. Thirteen patients were selected, of whom 4 were designated as training cases, 5 were used for initial assessment, and 4 were used for recertification. To reflect clinical practice, in 1 case disability was sufficiently severe that answers were provided by a caregiver. Although the final selection of patients was a sample of convenience, they were thought to be representative of a “real-life” cross section of poststroke outcomes and to be suitable for inclusion in the training package. We obtained a consent for videotaping and use for training and research purposes from all patients, in line with national and local protocols.

For the training component of the package, 2 patients with easily categorized disability were chosen as initial “introductory” cases. The remaining 2 training cases were chosen to highlight perceived problem areas in the application of the scale. Each of the training cases was followed by an explanatory discussion of the correct mRS score and the rationale for this grading. An accompanying booklet gave background information on the general principles of mRS scoring, including detailed definitions of the categories and discussion of what is considered to be best practice in the application of the scale. In formulating this advice, we made reference to the original description of the mRS and to recent work by means of a structured interview.4 To minimize potential language problems, we provided a transcript of the text with translation into local language. Fully translated training packages, with native speakers overdubbing the interview, have been made available for Spanish, Portuguese, and Italian researchers; a subtitled Chinese version has also been produced. Assessment for certification was performed in a variety of settings, including individual viewing of the cases, group viewing within a center, and supervised group viewing sessions at formal training meetings.

#### Recording and Scoring of the Assessments

Recordings of the interviews were analyzed and scored independently by 2 observers who were both experienced in mRS grading (K.R.L. and H.G.H.). No attempt was made to “script” the mRS assessments, and there was little postinterview editing. During a pilot run of the study, we immediately identified that some of the answers given by patients were ambiguous, and in at least 2 of the cases, debate arose as to the most appropriate category. However, a decision was made to keep the complete interviews as recorded; it was thought that scenarios artificially scripted to fit an mRS grade neatly would not have adequately prepared assessors for the difficulties inherent in grading real patients.

Scoring for the assessment component of the package took account of those patients who did not unequivocally fit a single grade. A final decision on correct grading was made by K.R.L. and H.G.H., supplemented by analysis of the results from an international pilot study involving 100 participants. We defined a correct grade as one assigned by both trainers and by >50% of trainees. A grade of an “acceptable” response was defined arbitrarily as one that was deemed by the trainers to be incorrect but that followed the basic scoring guidelines and had been assigned by a substantial minority (10% to 49%) of assessors in the pilot. Any grade offered by <10% of assessors or that clearly did not follow the accepted scoring was defined as “unacceptable.” According to these scoring rules, any candidate who graded all cases correctly, including “acceptable” answers, was awarded certification (Table 1). This scoring system was developed so that certification was awarded only to those assessors who had demonstrated a good knowledge of the mRS application but recognized that some merit should be given for “acceptable” but incorrect answers. To minimize variability, we instructed assessors to choose the more severe mRS grade when hesitating between 2 scores.

We awarded certificates of completion, along with separate confidential feedback on the actual score achieved, to all who achieved the target. Scores for individual patients were not released. We encouraged assessors who did not achieve certification on their first attempt to review the training material and resubmit an amended set of grades for the full set of certification cases. We gave no specific feedback on their errors.

### Results

Between March 2003 and May 2006, investigators from >25 countries submitted the end-of-training assessment. Data are available on 1800 assessments. The assessors were a mixed group of principal investigators and coinvestigators, study nurses, and research assistants (Table 2). The majority of respondents were part of the investigating team from countries involved in the Stroke–Acute Ischemic–NXY-059 Treatment (SAINT-I)14 trial only or in both

#### Table 1. Scoring System for Certification Process

<table>
<thead>
<tr>
<th>Grading</th>
<th>Certification</th>
</tr>
</thead>
<tbody>
<tr>
<td>All 5 cases correct</td>
<td>Pass (qualified 5)</td>
</tr>
<tr>
<td>4 correct cases and 1 acceptable</td>
<td>Pass (qualified 4)</td>
</tr>
<tr>
<td>3 correct cases and 2 acceptable</td>
<td>Pass (qualified 3)</td>
</tr>
<tr>
<td>Any other combination</td>
<td>Not qualified</td>
</tr>
</tbody>
</table>
the SAINT-II and the Cerebral Hemorrhage and NXY-059 Treatment (CHANT) study.15

The correct mRS scores for the certification cases, along with the proportion of scores assigned by observers, are shown in Table 3. To allow continued use of the training resource, the “correct” scores for assessment and recertification were not made public and are shown in a sequence different from the cases on the DVD.

There was a spread of opinion on all of the cases, with submitted answers spanning 3 to 5 mRS grades for each answer. For 3 of the cases, the majority of respondents opted for the correct grading; in 2 of the cases, opinion was split, with a substantial proportion of assessors (39.2% and 40.4%) choosing a lower mRS grade than the correct answer. We accounted for this variation in opinion in the final scoring, and the lower grade was defined as an “acceptable” answer. Twenty-three assessors gave 2 scores, despite explicit instructions to choose the best single score. If the scores given were the correct and an acceptable score, then a “pass” was allowed; otherwise, the grading was considered invalid.

Percentages of respondents achieving acceptable scores for the certification assessment are given in Table 2. The majority of assessors (1464, 81.3%), achieved a “pass” on the certification exercise. However, only 38% of these individuals graded all of the 5 cases correctly. The remainder of the group comprised those assessors who wrongly assessed 1 or both of the previously described equivocal cases but whose assessment was still defined as acceptable. Of the 336 who did not achieve certification on their first attempt, 85% scored a “pass” on a second attempt.

Demographic data on assessors submitting for certification were collated. Results of the certification assessment are presented by training (Table 2) and by country of origin (Figure). We intended the recertification process to be undertaken 1 year after the initial training, as such full data on recertification are presently limited to 370 results. Of these, only 6.5% assessors failed to achieve a satisfactory score (Table 3).

**Discussion**

Consistency in grading of poststroke disability is crucial both in daily clinical work and in the context of a clinical trial. For trial purposes, consistency is more important than accuracy. The potential for significant variation in application of the mRS is now apparent.6 We developed a digital training resource for mRS grading in an attempt to improve this situation. Our results show that mass training of observers in use of the mRS is achievable in the context of a clinical trial via the use of a novel DVD training package.

Several issues arose during development of the DVD that deserve comment. We used no specific criteria to select patients for the training or certification components of the package, although patients used for assessment were judged to be suitably taxing to allow a valid assessment of ability. A clustering of grades around the midrange was noted in the cases selected to be used for the certification process. It has been shown that clinicians are comfortable to assign grades at the extremes of the mRS, possibly because these grades are well-defined or because deviation can be in 1 direction only. Therefore, we decided to proceed to use this relatively biased sample.

Even with a training system, it may be impossible to completely remove interobserver variability for the “difficult” midrange grades. For assessment cases 3 and 9, respondents were almost evenly split between mRS grades 2 and 3. This is not a failure of the training, rather an example of the complexity of assigning fixed grades to real patients. We anticipate that those video cases that have divided opinion will be reviewed again by trainees, and discussion of these cases with other investigators will facilitate further improvement in mRS application. It is in the midrange of mRS grades that the reliability of the scale assumes the greatest impor-

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**Table 2. mRS Gradings Submitted for Certification and Recertification, %**

<table>
<thead>
<tr>
<th>Case</th>
<th>mRS 0</th>
<th>mRS 1</th>
<th>mRS 2</th>
<th>mRS 3</th>
<th>mRS 4</th>
<th>mRS 5</th>
<th>Correct grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.2</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>0.0</td>
<td>0.9</td>
<td>26.5</td>
<td>2.7</td>
<td>0.0</td>
<td>0.0</td>
<td>2</td>
</tr>
<tr>
<td>3</td>
<td>0.7</td>
<td>55.0</td>
<td>72.6</td>
<td>39.2</td>
<td>0.0</td>
<td>1.5</td>
<td>3</td>
</tr>
<tr>
<td>4</td>
<td>6.3</td>
<td>44.1</td>
<td>0.7</td>
<td>58.1</td>
<td>0.0</td>
<td>3.7</td>
<td>3</td>
</tr>
<tr>
<td>5</td>
<td>92.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>14</td>
<td>93.8</td>
<td>7.3</td>
</tr>
<tr>
<td>6</td>
<td>0.9</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>86</td>
<td>0.9</td>
<td>0.0</td>
</tr>
<tr>
<td>7</td>
<td>4</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>5</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>8</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>9</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.1</td>
</tr>
</tbody>
</table>

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**Table 3. Certification Scores by Background Training**

<table>
<thead>
<tr>
<th>Position</th>
<th>Not Qualified</th>
<th>Qualified 3</th>
<th>Qualified 4</th>
<th>Qualified 5</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coinvestigator</td>
<td>125 (16.7%)</td>
<td>96 (12.9%)</td>
<td>224 (30.0%)</td>
<td>302 (40.4%)</td>
<td>747</td>
</tr>
<tr>
<td>Principal investigator</td>
<td>28 (17.6%)</td>
<td>18 (11.3%)</td>
<td>48 (30.2%)</td>
<td>65 (40.9%)</td>
<td>159</td>
</tr>
<tr>
<td>Study nurse</td>
<td>53 (25%)</td>
<td>29 (13.7%)</td>
<td>64 (30.2%)</td>
<td>66 (31.1%)</td>
<td>212</td>
</tr>
<tr>
<td>Other</td>
<td>130 (19.0%)</td>
<td>71 (10.4%)</td>
<td>228 (33.4%)</td>
<td>253 (37.2%)</td>
<td>682</td>
</tr>
<tr>
<td>Total</td>
<td>336</td>
<td>214</td>
<td>564</td>
<td>686</td>
<td></td>
</tr>
</tbody>
</table>

*For explanation of “Qualified 5, 4, 3,” see Table 1.
distance; mRS-based outcomes are frequently dichotomized, with mRS ≥4 defining poor outcome (STICH trial)\(^3\) or scores ≤2 defining good outcome (ECASS II).\(^2\) Given the problems with midrange mRS grades highlighted by our data, increasing use of nondichotomized trial end points should be encouraged. To this end, the SAINT and CHANT trials examined changes in mRS across the continuum of possible grades.

It is difficult to adequately measure the “success” of a multidimensional intervention such as an educational resource. Analyzing only “pass” rates for the assessment is a relatively crude measure, as it is likely that even those viewers who failed the initial certification will have gained improved knowledge of the mRS scores and assessments. Despite this, the high rate of satisfactory scoring on the certification exercise is reassuring.

We gave assessors confidential feedback on their total score. This allowed all users to review the cases and perhaps to correct any grading errors. Data are not available for all of the second attempts at certification, which raises the possibility of sample bias, but the high “pass” rate seen provides further support for utility of the training package. Given that our training package is the only educational resource available for training, it is safe to assume that this improvement was achieved with no extra tuition other than repeated viewing of the package and knowledge of previous scoring.

Purists will argue that our data do not conclusively prove the benefit of the training package. The primary purpose of the package was to improve reliability of mRS grading in large clinical trials, so we did not design a “control” arm of assessors not exposed to training. Given that our training package is the only educational resource available for training, it is safe to assume that this improvement was achieved with no extra tuition other than repeated viewing of the package and knowledge of previous scoring.

It is recognized that some questions as to optimal delivery of the package remain unanswered, such as how best to address the issue of repeated failure to achieve certification and whether training should be performed alone or in a group setting with the opportunity to discuss the cases and content. Work is ongoing to answer these questions.\(^7\) Already there is scope for further improvements; for instance, making the training available on the Internet could ease dissemination to the target audience.
provide a powerful tool for better definition of the properties of the scale. Analysis of interobserver and intraobserver variability of the scale with further subanalysis of individual components of the scale and relations to country of origin and level of training are ongoing.

We live in a digital age, and using the available technology to deliver educational resources makes scientific and economic sense. Strategies to improve the reliability of the mRS are needed, and it is likely that electronic dissemination of teaching material to participants in multicenter clinical trials will be widely used in the future. We have demonstrated that digital training in poststroke assessment is feasible and accepted by most potential assessors. Further work to quantify the potential impact of such training on the quality of future stroke trials is required.

Acknowledgments

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Source of Funding

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Disclosures

K.R.L. was the international principal investigator for the GAIN International and SAINT I trials and chairs the steering committee for the CHANT and SAINT I and II trials. He has published data in support of the mRS as the optimal end point for acute stroke trials. He has received fees, expenses, and institutional grants relating to these and other trials from GlaxoSmithKline, AstraZeneca, and several other pharmaceutical companies that have or are developing treatments for stroke. H.G.H. is an employee of AstraZeneca. K.R.L., M.W., J.D., and T.J.Q. have applied for academic grant support to continue work on developing stroke outcome assessments with the mRS.

References

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