Reproducibility of Automated Voice Range Profiles, a Systematic Literature Review

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Summary: Objective. Reliable voice range profiles are of great importance when measuring effects and side effects from surgery affecting voice capacity. Automated recording systems are increasingly used, but the reproducibility of results is uncertain. Our objective was to identify and review the existing literature on test-retest accuracy of the automated voice range profile assessment.

Study design. Systematic review.


Methods. We conducted a systematic literature search of six databases from 1983 to 2016. The following keywords were used: phonetogram, voice range profile, and acoustic voice analysis. Inclusion criteria were automated recording procedure, healthy voices, and no intervention between test and retest. Test-retest values concerning fundamental frequency and voice intensity were reviewed.

Results. Of 483 abstracts, 231 full-text articles were read, resulting in six articles included in the final results. The studies found high reliability, but data are few and heterogeneous.

Conclusion. The reviewed articles generally reported high reliability of the voice range profile, and thus clinical usefulness, but uncertainty remains because of low sample sizes and different procedures for selecting, collecting, and analyzing data. More data are needed, and clinical conclusions must be drawn with caution.


INTRODUCTION

When treating voice disorders, measurement of outcome as well as side effects is important, and objective methods of measurement are of importance in ear-nose-throat departments and in speech-language therapy clinics.1 Knowing the reliability of the different assessment methods and types must be considered a minimum requirement if treatment results are to be correctly interpreted. The European Laryngological Society1,2 and the Union of European Phoniatricians’ recommend the use of voice range profile (VRP) when assessing the voice. This measures the maximum voice capacity in terms of limits in vocal fundamental frequency (f0) and intensity—parameters that can be changed by disease and by treatment, and are of great significance for the functionality of the voice. Knowledge of VRP assessment reliability is sparse. Most likely, many possible influencing sources cause variation in the assessment, for instance, natural variation in the voice from day to day, different times of the day, with and without vocal warm-up, clinician’s motivation and elicitation strategies, preciseness of the protocol, and more.1,4–18

Previously, VRPs were recorded by manual procedures, where the patient had to match and hold a tone for up to 3 seconds, while the clinician evaluated the f0, and read the sound level from a sound level meter.19 The reliability of these manual procedures has been investigated in test-retest studies of healthy voices, where studies find the test-retest variation varying from 1 to 10 dB in intensity range and 1–4 semitones in frequency range.13,16,19,20

At present, the measurement is automated by the use of computer programs and corresponding equipment, which facilitate the process for both patients and clinicians.21 Although there is still a need for a consistent clinician and protocol, the demand for the patients to match their pitch to a musical note and hold it steady for up to 3 seconds is no longer required, as some of the new automated methods require only very short tone durations.21,22 Nowadays, very short phonation times will be detected and the voice is recorded and analyzed precisely in real time, rendering direct comparability between the new and the old methods very difficult. In addition, former data of variability and reproducibility cannot be considered representative for the automated VRP.5 It would be reasonable to assume larger SPL variation, and thus decreased reliability, of the automated method, when the vocal production needs only to last milliseconds. However, the programs typically do not register all these very short phonations. Instead, they accumulate them, and only include them in the voice analysis when a certain time threshold has been reached, for example 0.1 sec.21,23

It is important to note that only the reliability of the automated VRP is assessed, and not the validity. Whereas reliability concerns the difference between two equal measurements (the same clinician measuring VRP on the same subjects, under the same recording conditions, using the same protocol), the validity concerns the amount of measurement error, and the preciseness of the results reflects reality.24

Based on a systematic literature review, we aimed to identify differences between test and retest of VRP in normophonic,
healthy voices using automated measurement, thus achieving a clearer insight into the assessment variation. In the present study, we use the term *automated* to cover VRPs recorded with computer program with a clinician or experimenter providing guidance, coaching, and encouragement to the patient.

**METHODS**

**Study design**
A systematic literature review was conducted. We adhered to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses checklist and statement recommendations.  

**Review objective**
The electronic search strategy was guided by the study question: identify frequency and/or intensity differences between test and retest in the automated VRP of healthy voices. The variables of interest were highest and lowest frequency and intensity, frequency and intensity ranges, and area (number of cells; VRP size) (Table 1, applied abbreviations).

**Literature search**

**Information sources and search**
The systematic literature search was constructed as a block search and conducted electronically on June 7, 2016. It was supervised by The Medical Research Library, the medical special library for The University Library of Southern Denmark. Six databases, including PubMed, MedLine, and Embase (Table 2), were searched for relevant articles in the time period from 1983 to 2016. Studies before 1983 were considered irrelevant, as this was the promotional year for the first automated VRP technology. We applied a core set of key words and reviewed search terms pertaining to the VRP (phonetogram, voice range profile, acoustic voice analysis, voice capacity assessment, etc). The specific use of search terms and truncations is provided in Table 1. Reference lists were reviewed for relevant literature not included in the database search. All titles and abstracts were downloaded onto the reference management database EndNote X6, Thomson Reuters, New York, NY. Duplicates and references that clearly deviated from the subject were removed.

**Inclusion process**
Two independent raters (TR and TP) assessed abstracts for further inclusion. In cases of disagreement, discussions between raters led to agreement. At full-text ratings, “reason for exclusion” codes were used. Discussions were conducted at all disagreements, including incongruence in the codes. The two investigators read the full text together and discussed whether the codes were correct. A third investigator, either author A-KD or ÅMG, was involved in case of doubt or disagreement of technical or statistical and other questions, respectively. Here, the issue in question was discussed informally, yet in accordance with the eligibility criteria until agreement was reached.

**Eligibility criteria**
For an article to be included, it was required to:

- measure VRP with the automated measurement
- present quantitative assessment of data, for instance means and standard deviations or intraclass correlation coefficient on at least one of the following parameters:
  - maximum intensity measured in dB SPL
  - minimum intensity measured in dB SPL
  - SPL range measured in dB (lowest to highest SPL)
  - maximum f0 measured in Hz or ST (highest tone)
  - minimum f0 measured in Hz or ST (lowest tone)
  - semitone range measured in Hz or ST (lowest tone to highest tone)
  - area, measured in cells (size of the VRP)
- measure healthy voices with no history of voice intervention or treatment
- report no intervention, treatment or other possible influencing factors between the two tests
- have uniform recording conditions—both under and between test and retest
- be written in Danish, Norwegian, Swedish, English, or German

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**TABLE 1.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Abbreviation</th>
<th>Measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Highest intensity</td>
<td>Max SPL</td>
<td>dB SPL</td>
</tr>
<tr>
<td>Lowest intensity</td>
<td>Min SPL</td>
<td>dB SPL</td>
</tr>
<tr>
<td>Intensity range (lowest to highest SPL)</td>
<td>SPL range</td>
<td>dB</td>
</tr>
<tr>
<td>Lowest frequency</td>
<td>Min f0</td>
<td>Hz/ST</td>
</tr>
<tr>
<td>Highest frequency</td>
<td>Max f0</td>
<td>Hz/ST</td>
</tr>
<tr>
<td>Frequency range (lowest to highest tone)</td>
<td>ST range</td>
<td>ST</td>
</tr>
<tr>
<td>Area (semitones times decibels/number of cells)</td>
<td>Area</td>
<td>(ST x dB)/cells</td>
</tr>
</tbody>
</table>

**Abbreviations:** dB SPL, decibel sound pressure level; Hz, hertz; ST, semitones.

**TABLE 2.**

<table>
<thead>
<tr>
<th>Databases</th>
<th>Search Terms</th>
<th>Truncations</th>
</tr>
</thead>
<tbody>
<tr>
<td>PubMed</td>
<td>Phonetogram</td>
<td>f0: fundamental frequency</td>
</tr>
<tr>
<td>Cochrane Library</td>
<td>Phonetography</td>
<td>SPL: sound pressure level</td>
</tr>
<tr>
<td>ComDisDome</td>
<td>Voice range profile</td>
<td></td>
</tr>
<tr>
<td>Embase</td>
<td>SPL</td>
<td></td>
</tr>
<tr>
<td>CINAHL (EBSCO)</td>
<td>Acoustic voice analysis</td>
<td></td>
</tr>
<tr>
<td>Scopus</td>
<td>Voice evaluation</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Voice capacity</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Voice assessment</td>
<td></td>
</tr>
</tbody>
</table>

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4.26,27

14,26,27
Studies that met these criteria were assessed with the Critical Appraisal Skills Programme’s “Case control checklist” from May 31, 2013, which assesses level of evidence including risk of bias. During the selection process, all subsections of articles were of interest. If a small test-retest study was part of a clinical study, for instance a case-control study presenting a control group with no intervention between test and retest, this was included in our study to ensure that as much relevant test-retest data from the literature as possible were included. The checklists were completed, considering these minor reliability tests as the “primary focus” of the article in question.

Statistics
Kappa statistics were used to calculate the level of interrater agreement using STATA/IC 13.1 (StataCorp LP, College Station, TX).

RESULTS
Systematic literature search
Study selection and data extraction
The initial electronic search resulted in 5455 titles and abstracts (Figure 1). After removing duplicates, 483 abstracts remained. Of these, 231 were included for further full-text
assessments. Kappa statistics for interrater agreement (abstract selection) was 0.82, equaling “substantial” interrater agreement ($P < 0.0001$). In full-text rating, two criteria were explored: (1) whether the study applied automated VRP equipment, and (2) whether the study held test and retest VRP data of healthy voices. These criteria were proven to be the two most frequent reasons for exclusion (180 full texts, comprising 77.9% of the studies). Six papers met the criteria for inclusion in the review (Tables 3 and 4). Kappa statistics for interrater agreement in full-text selection were 0.66 ($P < 0.0001$).

Six articles fulfilled all inclusion criteria (Tables 3 and 4). Together, they presented data for a total of 66 adult participants aged between 19 and 70 years with healthy voices. In one study, the voices were assessed in the morning and again in the afternoon; this was repeated 4 weeks later. Between the morning and the afternoon recordings, all participants worked in a call center: more than 50% of them talked more than 8 hours a day, which has most likely put some strain on their voices. Therefore, the afternoon assessments were excluded from our review, whereas the set of morning data is included here. Automated VRPs and clinician guiding were used in all articles.

Four articles stated that test-retest was part of the aim and focus of the study\cite{5,11,31,32}; the last two studies had different foci.\cite{29,30} One tested voice use at work, and had two recordings well suited for our purpose\cite{29}; the other was a therapy study, where only the control group receiving no intervention between the two tests was included in the review.\cite{29} In only one study, both gender and specific age range of the test-retest participants was provided;

<table>
<thead>
<tr>
<th>Author (year)</th>
<th>n</th>
<th>Gender and Age</th>
<th>Trained or Untrained Voices</th>
<th>Aim and Focus of Study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sanchez et al (2013)\cite{5}</td>
<td>6</td>
<td>3 males, 3 females(*)</td>
<td>Untrained healthy Australian voices</td>
<td>(1) To add to the body of knowledge about automatic phonetograms (2) Investigate the test-retest reliability of VRP data</td>
</tr>
<tr>
<td>D’Haeseleer et al (2013)\cite{29}</td>
<td>7</td>
<td>Gender N/S (mean age 21.4, SD 1.8 y, range 19–25 y)</td>
<td>Students from a bachelor’s program in music and performing musical art</td>
<td>Investigate the impact of manual circumlaryngeal therapy on the vocal characteristics of future elite vocal performers (the control group received no therapy, and is included in the present review)</td>
</tr>
<tr>
<td>Schneider-Stickler et al (2012)\cite{30}</td>
<td>30</td>
<td>21 females, 16 males (mean age 29.6 y ± 8.5 y) (7 drop outs, leads to n = 30)</td>
<td>Untrained, but professional: employees at a telecommunication company</td>
<td>Examine the voice use at work and introduce biofeedback software into real-life workplace situation to improve vocal performance</td>
</tr>
<tr>
<td>Hallin et al (2012)\cite{11}</td>
<td>3</td>
<td>Males(*)</td>
<td>Untrained Swedish speakers</td>
<td>Suggest protocols for recordings and analyses of speech range profiles and voice range profiles</td>
</tr>
<tr>
<td>Chen (2008)\cite{31}</td>
<td>10</td>
<td>N/S(*)</td>
<td>Untrained Taiwanese Min or Mandarin speakers Hospital employees and university students</td>
<td>(1) Investigate the physiological frequency and intensity ranges of the tonal dialect of Min (2) Compare the physiological frequency and intensity ranges of Min with those of nontonal languages</td>
</tr>
<tr>
<td>Behrman et al (1996)\cite{32}</td>
<td>10</td>
<td>8 females, 2 males (age 19–70)</td>
<td>Untrained hospital employees and speech-language pathology students</td>
<td>(1) Determine the important features of the contours of the VRPs of patients with organic pathology (2) Determine if the VRP is a clinically useful, within-subject measure of change in vocal function as a result of surgical intervention</td>
</tr>
</tbody>
</table>

* Specific age range for test-retest participants not stated.
The mean age was, however, not stated. The most frequent interval between test and retest was 1–4 weeks; however, in one study, the retest was performed after 20 minutes with complete vocal rest in between and in another after 3–4 months. Recoding protocols were somewhat similar, although variations in use of glissandi or tone-by-tone methods were applied. One study had a recording time limit.

Three types of equipment were used in the studies, Kay Elemetrics’ Voice Profiler being the most frequent. With only six studies, it is not possible to analyze whether the reproducibility of these is alike.

Table 4 shows the VRP reproducibility reported by the six studies included in the review. Three studies provided reliability measures as correlations ($r$), and found high reliability, although one study found lower reliability in ST range and area. The other three studies reported results in means or medians. None of these provided data on all seven VRP variables. Differences from test to retest were 4 dB ($P = 0.107$) in max dB; and 1 dB in min SPL. Two studies reported max $f_o$ differences of 78 Hz ($P = 0.500$) and 1 ST, respectively. Min $f_o$ differences were 8 Hz ($P = 0.581$) and 2 ST. Differences in semitone range varied from almost no difference to 3 ST (±5 ST). No studies reported specific differences in dB range or area.

**Biases and limitations of the included studies**

A risk of bias assessment was conducted by TR and TP (Table 6). Three types of bias were found: (1) lack of randomization, (2) lack of effectiveness of blinding, and (3) limitations in recording time. In one study, a limitation of the test was set to 20 minutes including questionnaires, voice recordings for acoustic analyses, and VRP measurements. The limit of the retest was 10 minutes, including both voice recordings for acoustic analyses and VRP measurements. Another study provided only a 20-minute break for the voice to recover before retesting. Neither of the studies used random selections of groups. Two studies included supra normal voice users, in the form of call center agents and students from a bachelor’s program in music and performing musical art, which might lead to testing bias. In at least four of the studies, participants worked or studied in the same company or class; accordingly, a risk of ineffective blinding due to discussion of the study purpose was present.

One study referred to the possible impact of giving the participants detailed information about study goals and vocal risk factors as well as the repetitions of VRP measurements as a potential bias of the study. Although not mentioned, the latter might also apply to the other studies. Other potential biases emphasized in the studies are cooperation and motivation of the participants and lack of vocal warm-up before elicitation of the VRPs.

**DISCUSSION**

To the best of our knowledge, this is the first systematic review evaluating reproducibility in the automated VRP assessment. The precision of the assessment is essential when it should be trusted for clinical and research application.
### TABLE 5.
Test-Retest Results of Included Studies

<table>
<thead>
<tr>
<th>Study or Independent Variable</th>
<th>Max SPL (dB)</th>
<th>Min SPL (dB)</th>
<th>SPL range</th>
<th>Max ( f_0 )</th>
<th>Min ( f_0 )</th>
<th>ST range</th>
<th>Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sanchez et al (2013)(^5)</td>
<td>High reliability(^*)</td>
<td>High reliability(^*)</td>
<td>High reliability(^*)</td>
<td>High reliability(^*)</td>
<td>High reliability(^*)</td>
<td>High reliability(^*)</td>
<td>High reliability(^*)</td>
</tr>
<tr>
<td>D’Haeseleer et al (2013)(^2)</td>
<td>Test: median: 108.0 dB</td>
<td>53.0 dB</td>
<td>–</td>
<td>Test: median: 1396.9 Hz</td>
<td>138.6 Hz</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Retest: median: 112.0 dB</td>
<td>54.0 dB</td>
<td>( P = 0.107^* )</td>
<td>Retest: median: 1318.5 Hz</td>
<td>130.8 Hz</td>
<td>( P = 0.500^* )</td>
<td>( P = 0.581^* )</td>
</tr>
<tr>
<td>Schneider-Stickler et al (2012)(^2)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Hallin et al (2012)(^1)</td>
<td>( r = 0.83 ) and ( 0.92 )</td>
<td>( r = 0.83 ) and ( 0.92 )</td>
<td>( r = 0.99 )</td>
<td>–</td>
<td>–</td>
<td>( r = 0.69 )</td>
<td>( r = 0.84 )</td>
</tr>
<tr>
<td></td>
<td>Test: mean: 64.2 (mean SD 4.3)</td>
<td>Test: mean 40.0 ST</td>
<td>Test: mean: 19.0 ST</td>
<td>Retest: mean: 39.0 ST</td>
<td>Retest: mean: 21.0 ST</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chen (2008)(^3)</td>
<td>( r = 0.83 ) and ( 0.92 )</td>
<td>( r = 0.83 ) and ( 0.92 )</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Test: mean: 64.2 (mean SD 3.7)</td>
<td>Test: mean 30.3 (mean SD 6.9)</td>
<td>Test: mean: 30.4 (mean SD 6.2)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Results are stated as they are reported in the respective papers according to the different statistic tests.

* No significant difference between test and retest, and thus indicates high reliability.
knowledge, it remains unclear to what extent differences before and after treatment can be ascribed to changes in the voice as a result of the treatment or to general variability in the assessment. The majority of the literature analyzed addressed issues other than test-retest variance. Six articles were included in the final analyses.

We included only studies of healthy voices, as we were testing the reliability of the assessment, and not overall variation in dysphonic voices, which are very likely to vary more from test to retest.13,27,35 This larger variation should be considered when applying the VRP (and all other voice assessments) clinically. Moreover, we included only physiological VRPs, as recommended by Pabon and the Voice Profiler Users Group.29 In the physiological VRP, the aim is to detect the physiological boundaries, or extremes, and not only the most beautiful tones. For this reason, dB max and range results from one study were not included.32 One could argue that the \( f_o \) ranges should also have been excluded, as \( f_o \) and intensity are related, and the participants in this article might not have reached their highest \( f_o \), as only the maximal comfortable level, and not the extreme intensity level, was pursued. Including these figures would raise a question regarding the validity of the automated VRPs, which in this case might be reliable, but may not be valid in the sense of testing the extremes of the vocal range, which should be the aim, when assessing the VRP contour. Here, experimenter guidance and motivation, as well as a stable protocol, play a great role.23

The difference of 3 ST in semitone range when using automated VRP assessment is in accordance with the test-retest differences reported for the manual procedures.19,26 However, in regard to measuring the dB level, the automated procedures, in general, seem to have a better reliability than the manual methods.13,16,20,35 One possible explanation is the accumulation of time in the cells. The automated systems register the voice only when it hits a cell several times and then reach a predefined accumulated time.23 Another explanation is the threshold for registering the min SPL value. The automated systems have different thresholds. For instance, Chen31 and Behrman et al12 used the Kay Elemetrics, Voice Range Profile, Model 4326 with a 50 dB SPL minimum threshold. Generally, healthy voices can phonate softer than the 50 dB SPL at a 30-cm measuring distance,37 and therefore reach the 50 dB SPL threshold repeatedly during a recording. This might fictively improve reliability as the min SPL threshold is reached repeatedly, thus not showing the true variance of the voice. The differences in results support our assumption that reliability data for the manual VRP cannot be considered representative for the automated VRP.

### Bias of included studies

Participants from two articles studied either speech-language pathology or music.29,32 Accordingly, they might have special interest in, and insight into, the study goal, resulting in smaller test-retest differences than in a broader layperson population. In one article, participant selection relied only on the participants’ subjective self-evaluation and reassurance of no previous history of dysphonia.30 Moreover, they defined hypofunctionality as the inability to reach 90 dB SPL (at 30-cm microphone distance), but there was no clear statement as to whether participants with hypofunctional voices were excluded from the results, causing a potential bias in this article. Furthermore, they allowed only a very short and limited recording time for their VRPs.30 In general, time limits in voice recordings can be problematic, owing to the variation in voice abilities, the participant’s understanding of the task, need for breaks, etc.2,5,6 This time limit might be part of the explanation for the larger test-retest differences found in this article. Another possible explanation might be that newer, and perhaps more sensitive, technology was deployed. Computer algorithms of the different automated VRP equipment are different and this could also lead to some bias. The time intervals between test and retest varied from 20 minutes29 to 3–4 months.11 The VRP can be a strenuous test for the voice,22 and a 20-minute break seems to be a relatively short time for the voice to recover. This could potentially induce a bias in that the voice is fatigued at the retest. Moreover, it is not clear whether there is a learning effect in the VRP, and how long this might last before wearing off, but it is probable. It is not possible to draw any conclusions about this from the present data, as there was no clear tendency for a learning effect, or the opposite, in the data, yet in the study that allowed the shortest break (20 minutes), “absence or minimal impact of learning effect” (page 4) was concluded.29

### Generalization of results of this study

The six studies used different VRP equipment, and because this alone might induce variations owing to differences in computer algorithms, microphone stability, headset and microphone details, and sensitivity to noise,23 generalization of the results is inadvisable, and no strong conclusions can be drawn. For the purpose of increasing the precision of the analyses, we tried

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**TABLE 6. Risk of Bias Analysis in the Studies Included**

<table>
<thead>
<tr>
<th>Risk of Bias Assessment</th>
<th>Random Selection of Subjects</th>
<th>Blinding of Participants</th>
<th>Exposure Bias</th>
<th>Bias in Assessment Method</th>
<th>Selective Reporting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sanchez et al (2013)5</td>
<td>No</td>
<td>?</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>D’Haeseleer et al (2013)29</td>
<td>No</td>
<td>?</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Schneider-Stickler et al (2012)30</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>?</td>
<td>No</td>
</tr>
<tr>
<td>Chen (2008)31</td>
<td>No</td>
<td>?</td>
<td>Yes</td>
<td>?</td>
<td>No</td>
</tr>
</tbody>
</table>
conducting a meta-analysis, but owing to limited amount of data, this was rejected.

Four of the studies outlined test-retest as a research focus. The largest study included 30 participants, but their focus was to assess voice demands in call center employees and they had a time constraint on the VRP. This was suitable for their focus, but questionable regarding our study. Excluding their data, 26 participants are left in the review, instead of the previously stated 66. Most articles present only data on selected variables, and these vary between studies. This might lead to a decrease in the power of the results, and thus the conclusions of the present study must be viewed with some caution. Estimation of disease-specific problems and the results of treatments are partly determined on the basis of VRP recording measurements, and even though clinical usefulness and reliable apparatus are indicated in this review, larger studies allowing for the clinical relevant differences in their estimation of number of participants are warranted.

CONCLUSION

This is the first literature review to specifically and systematically analyze the reliability of automated VRP assessment. The articles generally report high reliability of the VRP, and thus clinical usefulness, but uncertainty remains because of the low sample sizes and different procedures for selecting, collecting, and analyzing data. The current literature is not sufficient for clear results, and more studies with a higher level of evidence are warranted.

Acknowledgment

We acknowledge The Medical Research library at Odense University Hospital for their qualified assistance in the literature search.

REFERENCES