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Low agreement between mMRC rated by patients and clinicians – implications for practice

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Abstract:	



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Low agreement between mMRC rated by patients and clinicians — implications for practice

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To the editor,

Chronic breathlessness [1] causes immense suffering in cardiorespiratory diseases. The functional impact of activity-related breathlessness, measured on the modified Medical Research (mMRC) scale [2], is highly prognostic, informs disease evaluation and management including in chronic obstructive pulmonary disease (COPD) [3], and is widely used for determining eligibility in clinical trials.

In clinical practice, mMRC is often rated by physicians based on the patient's medical history. It is unknown to what extent mMRC ratings differ when administered by clinicians compared with patient self-report. The ratings may be influenced by other clinical characteristics, such as the patient's functional status. The New York Heart Association (NYHA) scale, which is similar to mMRC and is key for management of heart failure, is associated with functional status, measured using the Australia-modified Karnofsky Performance Status (AKPS) [4], but discriminates poorly between clinically important performance states in people with advanced disease [4].

The primary aim of this study was to evaluate the agreement between clinician- and patientreported mMRC scores. Secondary aims were to evaluate whether the agreement differed by severity of activity-related breathlessness and how clinicians' and patients' ratings correlated to the patient's functional status.

This was a pooled analysis of two randomised, placebo-controlled trials of morphine [5] and sertraline [6] for chronic breathlessness. Only data at screening and baseline were used (before any study treatment was initiated). Patients had severe life-limiting illnesses and chronic breathlessness defined as a clinician-rated mMRC ≥ 2 at screening despite optimal treatment for the underlying cause(s), as detailed elsewhere [5, 6]. Participants with missing data on clinician- or patient-reported mMRC (n=68) were excluded. No data were imputed.

mMRC was rated by clinicians at screening and was then self-reported by patients in their study diary at baseline (before randomisation). Patients' functional status was rated by clinicians at baseline using AKPS [7]. The primary analysis compared clinician and patient mMRC ratings conducted within three days or less. A sensitivity analysis was performed using ratings performed four days or more apart. Agreement was analysed using quadratic-

 weighted Cohen's Kappa, categorized according to Landis *et al.* [8]: 0 = no (chance) agreement; 0.01-0.2 = slight; 0.21-0.40 = fair; 0.41-0.60 = moderate; 0.61-0.80 = substantial; $\geq 0.81 = high$ agreement. Associations between the mMRC ratings and patients' functional status (AKPS) were analysed using Kendall's tau. The study was approved by relevant human research ethics committees and all participants provided written, informed consent. Statistical analyses were conducted using the Statistical Package for the Social Sciences (SPSS) software Version 24.0 (Armonk, NY: IBM Corporation).

In total, 464 patients (294 from the morphine and 170 from the sertraline trial) had paired clinician and patient mMRC ratings. The time between clinician and patient mMRC ratings was a median 0 (IQR, -1, 0) days; 312 patients had ratings within 3 days (by 42 physicians) and were included in the primary analysis: mean age 73.8 (standard deviation [SD] 8.8); 63.5% men; most common diagnoses were COPD (70.5%), interstitial lung disease (17.3%), lung cancer (13.8%) and heart failure (4.8%); and patients were ambulatory with a mean AKPS of 61.5 (SD 10.1). Characteristics were similar between patients who were included and excluded from the primary analysis.

Agreement between clinician- and patient-reported mMRC (scored within 3 days; n=312) is shown in Figure 1. The ratings differed considerably and the agreement for all categories was slight to fair, Cohen's kappa 0.238 (95% confidence interval [CI] 0.143, 0.326). The rate of under- and over-estimation by clinicians compared to patient self-reports was similar (Figure 1). However, agreement was better for higher mMRC scores (25% for clinician mMRC 2; 31% for mMRC 3; and 61% for mMRC 4; p < 0.001 using Mantel–Haenszel chi-square test). Functional status was more closely related to clinician-rated mMRC (tau=-0.42; p<0.001) than patient-rated mMRC (tau=-0.22; p<0.001). For scores more than three days apart (n=152), agreement was slightly lower, Cohen's kappa 0.154 (95% CI 0.047, 0.260), but findings were otherwise similar.

This study for the first time evaluated the agreement between clinician- and patient-rated mMRC. The main finding was that only a minority of ratings agreed, with similar rates of clinician under- and over-estimation. These findings are consistent with reported disagreement between clinicians' and patients' ratings of subjective measures including symptom intensity [9] and quality of life [10]. Our study is the first indication of substantial

disagreement between clinicians and patients when assessing even a relatively objective measure such as when breathlessness limits exertion.

Secondly, a novel finding was that as activity-related breathlessness worsened, agreement between patients and their clinicians improved. The subjective symptom of breathlessness might be under-detected by the clinician until becomes visible as a "clinical sign" of reduced function. Functional status was more closely related to clinician-rated than patient-rated mMRC. This could reflect that patients reduce or avoid physical activities to limit their breathing discomfort – which could lead to patients under-estimating their activity-related breathlessness (as they become more inactive) – contributing to symptom under-report. Clinicians may also incorporate other clinical information when rating breathlessness such as the patient's disease severity and functional status. In fact, this could make the clinician-ratings even more predictive than the self-report of future clinical outcomes, which should be evaluated in studies with long term outcome data.

A strength of the analysis was the large sample of patients with chronic breathlessness, with ratings using standardised scales in the setting of randomised controlled trials. A potential limitation was the time between the ratings, hence the primary analysis included ratings done within three days. Given that mMRC only has five levels that are quite broad and the chronicity of breathlessness in the study population, mMRC scores should be stable within time periods longer than three days. As a clinician rated mMRC of 2-4 was an eligibility criteria, findings pertain mostly to moderate to severe chronic breathlessness. The improved agreement for higher mMRC scores might be partially related to getting closer to the upper limit of the scale. Higher agreement might also be found near the lower limit (mMRC 0-1) giving a U-shaped agreement for mMRC, which should be further explored. There were no data on how each clinician established a patient's mMRC. Involvement of patients in the clinician-rating is possible but would in fact make their scores more similar and overestimated the agreement.

The low agreement between clinician- and patient-rated mMRC has direct clinical implications, as mMRC is widely used to assess disease severity and prognosis, guide patients' management, and select participants for interventional symptom trials [3, 11]. The findings highlight that activity-related breathlessness is a subjective experience that is only weakly related to other commonly measured clinical parameters (including functional status),

and that symptom assessment should include self-report whenever possible to accurately capture patients' experiences [1, 12]. At the same time, given the complexity of chronic progressive diseases, comorbidities and symptoms, assessment necessitates clinician's involvement, which may also mitigate symptom under-reporting by patients. Training of clinicians to adequately assess breathlessness and gain a better proxy mMRC where self-report is not possible, would give more accurate representation of patient status, which is important in cardiorespiratory disease.

Improved method to assess exertional breathlessness is needed for use in clinical care, for selecting participants to clinical trials and to measure treatment effects. The mMRC might under-report symptoms in patients with milder disease and who have become less active due to breathlessness [13], and is too unresponsive to detect change. Standardised tests for measuring changes in activity-related breathlessness have been validated in COPD [14, 15].

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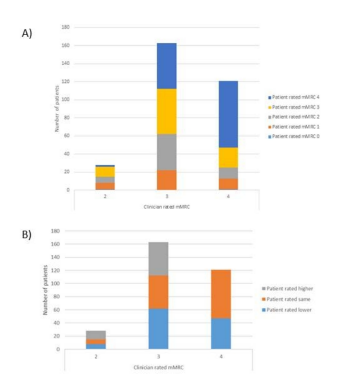
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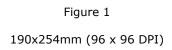
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Figure 1. Agreement between A) clinician- and patient-rated modified Medical Research Council (mMRC) breathlessness scores; B) distribution of lower, similar and higher patient vs. clinician ratings. Agreement was relatively low between patient and clinican rated mMRC, with an even distribution of under and over ratings for mMRC 2-3.





Low agreement between mMRC rated by patients and clinicians when assessing eligibility in randomised controlled trials — implications for practice

Magnus Ekström ^{1,2} Sungwon Chang ² Miriam J Johnson ^{2,3} Belinda Fazekas ² Slavica Kochovska ⁻² Chao Huang ³ David C Currow ^{2,3}

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