Brief assessment of male depression in clinical care: Validation of the Male Depression Risk Scale short form in a cross-sectional study of Australian men

Danielle Herreen 1,2, Simon Rice3,4, Ian Zajac2

ABSTRACT

Objectives To develop and validate a short form of the Male Depression Risk Scale (MDRS-22) for use in primary care, examining associations with prototypic depression symptoms, psychological distress and suicidality.

Design Cross-sectional study with 8-month follow-up.

Setting Community-based.

Participants A community sample of younger (n=510; 18–64 years) and older (n=439; 65–93 years) men residing in Australia (M age=58.09 years, SD=17.77) participated in the study. A subset of respondents (n=159 younger men; n=169 older men) provided follow-up data approximately eight months later.

Primary and secondary outcome measures Quantitative data were obtained through a survey comprising a range of validated measures, including the MDRS-22, the Patient Health Questionnaire (PHQ-9) and the Kessler Psychological Distress Scale (K10). The MDRS-22 was refined using exploratory and confirmatory factor analysis in line with best practice guidelines. Analysis of variance and generalised linear models were conducted to explore relationships between variables.

Results The short-form MDRS consisted of seven items (MDRS-7) and captured all of the domains in the original tool. Participants with mixed symptoms (PHQ-9 ≥ 10 and MDRS-7 ≥ 5) had significantly higher risk of mental illness (K10 > 25) and current suicidality (PHQ-9 item 9 ≥ 1) than those with exclusively prototypic symptoms (PHQ-9 ≥ 10 and MDRS-7 ≤ 5). Furthermore, the MDRS-7 was shown to be effective at predicting elevated symptoms of depression at follow-up, after controlling for previous depression diagnosis.

Conclusions Findings provide preliminary evidence of the potential utility of the MDRS-7 as a screening tool for externalised and male-type symptoms associated with major depression in men. Field trials of the MDRS-7 in primary care settings may facilitate identification of men at risk of suicide and psychological distress who do not meet cut-off scores for existing measures of major depression symptoms.

INTRODUCTION

Major depressive disorder (MDD) is a common psychiatric condition and the leading cause of disability worldwide.1 2 MDD is twice as prevalent in women than men,3 and severe depression is known to significantly increase risk of suicide.4 Although men are less likely to be diagnosed with a depressive disorder,5 they are three times more likely to die by suicide compared with women.6 Current approaches to the diagnosis of depression (eg, as per International Classification of Diseases 11th Revision [ICD-11] or Diagnostic and Statistical Manual of Mental Disorders 5th Edition [DSM-5] criteria) emphasise symptoms including persistent sadness, loss of interest or pleasure in previously enjoyable activities, as well as changes in affect, cognition and neurovegetative functioning.7 8 However, a growing number of studies suggest that a significant proportion of men suffering from depression might experience a distinct phenotype.9–11 Congruent with masculine role norms, this male-typical phenotype includes anger, substance misuse, emotion suppression and risk-taking domains.10 12 However, these putative symptoms are not currently included in standard diagnostic criteria or screening measures, and it has been suggested that this might account in part for the underdiagnosis of male depression cases, and therefore under-recognition of (and treatment for) men at heightened risk of suicide.13

While men are often regarded as being less likely to seek help than women, recent...
statistics largely do not support this claim. In Australia, around 89% of men attend primary care annually. Among men experiencing mental health difficulties, annual primary care attendance is similarly high with estimates of 80%–96% of men with symptoms of depression reporting a visit to primary care within the previous 12 months. Similarly, findings from the UK demonstrate that although men are overall less likely to attend primary care compared with women, attendance rates in men and women with comparable underlying morbidities, including depression, are similar. Furthermore, findings from a population study of healthcare contacts among Canadian suicide decedents demonstrated that over 60% (n=1792) of men who died by suicide accessed professional mental healthcare in the year before their death. These findings highlight the essential role of primary care physicians in identifying depression and suicide risk in men in order to facilitate effective treatment.

Growing interest in gender-sensitive assessment of men’s depression has seen the development of male-specific screening tools to identify symptoms that align with men’s socialisation and gender norm processes (eg, Brownhill et al, Zierau et al, Magovcevic and Addis). One recently developed and widely validated measure for assessing externalising and male-type symptoms in men is the Male Depression Risk Scale (MDRS-22). The MDRS-22 consists of 22 items assessing six symptom domains including emotion suppression, drug use, alcohol use, anger and aggression, somatic symptoms, and risk-taking. Recently, Zajac and colleagues demonstrated that this tool, used in conjunction with a measure of prototypic depression symptoms (PHQ-9), was able to stratify men into three distinct risk groups: (1) prototypic symptoms (consistent with current MDD diagnostic criteria), (2) externalising symptoms consistent with masculine socialisation and (3) mixed depressive symptoms, reflecting both internalised (i.e., prototypic) and externalised symptomology. Further analyses showed that men in the externalising only group—men who are arguably missed when using measures of internalising symptoms—were at significantly increased risk of suicide compared with non-depressed men. Moreover, those with elevated externalised and prototypic symptomology were at highest risk of mental illness as well as suicide, highlighting the potential early identification and intervention benefits of leveraging male-specific tools in primary care settings.

Two-stage screening methods are commonly used in primary care and have been shown to be effective for increasing the recognition of depression. However, many primary care physicians report that time is a limiting factor in their capacity to comprehensively assess psychological issues, including depression, despite management of common mental disorders rating as a top reason for general practice attendance. To help address this issue, brief screening tools consisting of 15 items or less are often used, given their completion time is usually just a couple of minutes. Examples include the PHQ-9, the Kessler Psychological Distress Scale (K10) and the Beck Depression Inventory for Primary Care (BDI-PC).

To date, the MDRS-22 has demonstrated excellent psychometric properties as well as the ability to detect different groups of men who may be at increased risk of suicide and mental illness (eg, Zajac et al, Rice et al, Rice et al). However, given time constraints in primary care settings, the length of the current MDRS-22 is arguably impractical. The purpose of the present study was to develop a short form of the MDRS-22 to facilitate its use as a screening tool in busy and time-pressured healthcare settings. We also aimed to establish an initial set of cut-off scores for interpretive purposes. If the MDRS short form is to have clinical utility, it needs to be able to identify broader aspects of psychopathology. Thus, a secondary aim was to explore current and longitudinal risk of suicidality and mental illness by adopting a previously utilised categorisation according to cut-off scores on the MDRS and the widely used PHQ-9, which assesses prototypic depression symptoms. Furthermore, as adherence to masculine gender norms has been found to decline as men get older, younger and older men were examined separately to examine the utility of the tool across age groups.

METHOD

Participants and procedure

This cross-sectional study included baseline data from a community sample of 510 younger men aged 18–64 years (M=45.43, SD=14.56) and 439 older men aged 65–93 years (M=72.79, SD=5.88). A subset of respondents (n=159 younger men; n=169 older men) participated in the follow-up component. On average, 35 weeks (M=248.56 days, SD=24.59 days) elapsed between the provision of time 1 and time 2. The mean age for the overall sample was 58.09 years (SD=17.77). Eligible participants were Australian male residents over the age of 18 years who considered themselves fluent in English. Participants were recruited via paid advertisements displayed to Australian members of the Facebook social networking site (n=601; 63.3%) and through promotion of the study to community organisations (eg, Rotary, Men’s Shed). Time 1 data were collected between August and November 2019 using an online questionnaire. However, participants from local community organisations were provided with the option to complete a paper version of the survey to ensure inclusivity and accessibility of the sample and n=5 participants completed a paper version. Ethics approval was obtained from the University of Adelaide Human Research Ethics Committee and the CSIRO Health and Medical Human Research Ethics Committee (approval number H-2019-109). All participants provided informed consent. Reporting adhered to the STROBE cross-sectional guidelines. Table 1 presents a summary of the characteristics of the study participants at time 1 and time 2.
Open access

Public involvement
Participants were not involved in the design or conduct of this research; however, participants could nominate to receive updates on the results of the study.

Measures
Demographics
Participants reported their age, gender, relationship status, employment status, level of education and household income. They also reported whether they had previously been diagnosed with depression.

Male Depression Risk Scale (MDRS-22)
Externalising and male-type depression symptoms were assessed by the MDRS-22. The MDRS-22 contains 22 self-report items designed to assess six broad domains of externalising and male-type depression symptoms present in the last month including anger and aggression, drug use, alcohol use, emotion suppression, risk-taking and somatic symptoms using the condensed five-point Likert scale response format ranging from 0 (none of the time) to 4 (all of the time). Cronbach’s alphas for the MDRS are reported in Table 2 for both age groups and for the overall sample and are considered adequate.

The Patient Health Questionnaire (PHQ-9)
The PHQ-9 is a self-report depression screening tool for use in primary care that assesses nine symptoms consistent with the DSM-5 diagnostic criteria for MDD. Participants endorse how often they have experienced each symptom (eg, ‘Feeling down, depressed, or hopeless’) during the preceding 2-week period using a four-point Likert scale ranging from 0 (not at all) to 3 (almost every

Table 1 Sociodemographic characteristics of participants

<table>
<thead>
<tr>
<th>Variable</th>
<th>Younger men (&lt;65 years)</th>
<th>Older men (&gt;65 years)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Time 1 (n=510)</td>
<td>Time 2 (n=159)</td>
</tr>
<tr>
<td>Age range (years)</td>
<td>18–64</td>
<td>65–93</td>
</tr>
<tr>
<td>Age, M (SD)</td>
<td>45.43 (14.56)</td>
<td>72.79 (5.88)</td>
</tr>
<tr>
<td>Relationship status, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Single (never married)</td>
<td>118 (23.1)</td>
<td>27 (17.0)</td>
</tr>
<tr>
<td>Widowed/divorced/separated</td>
<td>68 (13.3)</td>
<td>19 (11.9)</td>
</tr>
<tr>
<td>Married/de facto</td>
<td>322 (63.1)</td>
<td>113 (71.1)</td>
</tr>
<tr>
<td>Prefer not to say</td>
<td>2 (0.4)</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>Employment status, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Employed full-time</td>
<td>227 (44.5)</td>
<td>66 (41.5)</td>
</tr>
<tr>
<td>Employed part-time</td>
<td>37 (7.3)</td>
<td>11 (6.9)</td>
</tr>
<tr>
<td>Employed casually</td>
<td>67 (13.1)</td>
<td>19 (11.9)</td>
</tr>
<tr>
<td>Not employed or unpaid work</td>
<td>94 (18.4)</td>
<td>24 (15.1)</td>
</tr>
<tr>
<td>Retired</td>
<td>73 (14.3)</td>
<td>39 (24.5)</td>
</tr>
<tr>
<td>Prefer not to say</td>
<td>12 (2.4)</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>Household income, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;$A35 000</td>
<td>136 (26.7)</td>
<td>28 (17.6)</td>
</tr>
<tr>
<td>$A35 000–$A65 000</td>
<td>91 (17.8)</td>
<td>32 (20.1)</td>
</tr>
<tr>
<td>$A65 000–$A105 000</td>
<td>100 (19.6)</td>
<td>44 (27.7)</td>
</tr>
<tr>
<td>$A105 000–$A160 000</td>
<td>97 (19.0)</td>
<td>26 (16.4)</td>
</tr>
<tr>
<td>&gt;$A160 000</td>
<td>65 (12.7)</td>
<td>20 (12.6)</td>
</tr>
<tr>
<td>Prefer not to say</td>
<td>21 (4.1)</td>
<td>9 (5.7)</td>
</tr>
<tr>
<td>Highest level of education, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Year 11 or below</td>
<td>49 (9.6)</td>
<td>11 (6.9)</td>
</tr>
<tr>
<td>Year 12</td>
<td>52 (10.2)</td>
<td>10 (6.3)</td>
</tr>
<tr>
<td>Certificate/diploma</td>
<td>154 (30.2)</td>
<td>55 (34.6)</td>
</tr>
<tr>
<td>Bachelor's degree</td>
<td>139 (27.3)</td>
<td>44 (27.7)</td>
</tr>
<tr>
<td>Graduate certificate/diploma</td>
<td>43 (8.4)</td>
<td>15 (9.4)</td>
</tr>
<tr>
<td>Postgraduate degree</td>
<td>72 (14.1)</td>
<td>23 (14.5)</td>
</tr>
<tr>
<td>Prefer not to say</td>
<td>1 (0.2)</td>
<td>1 (0.6)</td>
</tr>
</tbody>
</table>

Percentage may not equal 100% due to rounding.
¹ Household income reflects Australian Dollars.
day). A score of 10 and above is indicative of clinically significant depressive symptoms. In addition to utilising total PHQ-9 scores, we used item 9 as a measure of suicidality: ‘Over the past two weeks, how often have you been bothered by thoughts that you would be better off dead, or of hurting yourself in some way?’ We deemed those who scored 1 or more on this item to be currently experiencing suicidal ideation. Internal consistency of the PHQ-9 in the present study for the overall sample was high (α=0.93).

Kessler Psychological Distress Scale (K10)
The Kessler Psychological Distress Scale (K10) is a widely used measure in both research and primary care settings. It comprises 10 questions assessing a person’s negative emotional state in the preceding 30 days (eg, ‘About how often did you feel so nervous that nothing could calm you down’). Responses are based on a five-point Likert scale ranging from 1 (none of the time) to 5 (all of the time). In addition to examining K10 total scores, we created a binary variable with scores ≥2 indicating probable mental illness, consistent with published cut-off scores for the K10. Internal consistency of the K10 in this study for the overall sample was high (α=0.95).

Analytic sample
A total of 1114 participants commenced the study. However, 156 participants were not included in the analyses due to substantial missing data. Thus, N=949 participants who provided complete data for the items comprising the MDRS-22 were included in the item reduction process described below. Of this sample, n=29 did not provide complete data for the PHQ-9 or K10 items. Thus, models using these variables comprised n=920 participants.

Table 2  Item loadings derived from exploratory factor analysis (maximum likelihood estimation)

<table>
<thead>
<tr>
<th>Domains</th>
<th>Items</th>
<th>18–64</th>
<th>65+</th>
<th>Overall</th>
<th>Time 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emotion suppression</td>
<td>I bottled up my negative feelings</td>
<td>0.67</td>
<td>0.72</td>
<td>0.71</td>
<td>0.68</td>
</tr>
<tr>
<td>Alcohol use</td>
<td>I needed alcohol to help me unwind</td>
<td>0.44</td>
<td>0.45</td>
<td>0.48</td>
<td>0.37</td>
</tr>
<tr>
<td>Somatic symptoms</td>
<td>I had unexplained aches and pains</td>
<td>0.56</td>
<td>0.59</td>
<td>0.58</td>
<td>0.63</td>
</tr>
<tr>
<td>Aggression</td>
<td>I overreacted to situations with aggressive behaviour</td>
<td>0.69</td>
<td>0.74</td>
<td>0.71</td>
<td>0.30</td>
</tr>
<tr>
<td>Anger</td>
<td>It was difficult to manage my anger</td>
<td>0.75</td>
<td>0.74</td>
<td>0.75</td>
<td>0.65</td>
</tr>
<tr>
<td>Drug use</td>
<td>Using drugs provided temporary relief</td>
<td>0.36</td>
<td>0.44</td>
<td>0.42</td>
<td>0.44</td>
</tr>
<tr>
<td>Risk-taking</td>
<td>I stopped caring about the consequences of my actions</td>
<td>0.63</td>
<td>0.62</td>
<td>0.65</td>
<td>0.80</td>
</tr>
</tbody>
</table>

Note: Time 2 loadings derived using confirmatory factor analysis in the combined sample. All correlations were significant at p<0.001.

Statistical analyses
Data for the present study were analysed using IBM SPSS Statistics (V.26.0) except for the confirmatory factor analysis (CFA) undertaken in JASP (V.0.13.1). Various recommendations exist for the selection of items for short-form surveys including both Rasch analysis and descriptive approaches. Broadly speaking, the focus is on selecting items with maximum variability and which retain the theorised underlying construct—as well as subdomains—measured by the long-form scale. Therefore, we calculated descriptive (means, SD and skewness) and relational statistics (correlations) for each item (see online supplemental file 1). Items were then scored based on each statistic within its corresponding domain (ie, largest SD, strongest correlation) and summed across the different descriptive indices to derive a total performance score for each item. Parallel analysis consisting of 1000 permutations of the original raw data was used to determine thresholds for retaining factors. Exploratory factor analysis of the best performing items was performed with maximum likelihood estimation within each age group and in the combined sample. Stability of this solution was then established using CFA of time 2 data (n=328). Fit indices reported include: comparative fit index (CFI); the Tucker-Lewis index (TLI); the root mean square error of approximation (RMSEA) and the standardised root mean residual (SRMR). Interpretation of these indices were guided by the recommendations of Hu and Bentler.

In order to investigate the clinical utility of the reduced item scale, cut-off scores were determined for low (0–5), moderate (6–7), severe (8–12) and extremely severe (13+) symptom severity groups. The corresponding cumulative percentiles (cum5) at the upper boundaries of these categories were: low (cum5=63.5), moderate...
(cum% = 77.5), severe (cum% = 95.5), extremely severe (cum% = 100.0). These category scores were determined using previously reported cumulative percentiles that represented differing degrees of increased risk of recent suicide attempt for the MDRS-22. A 2x2 analysis of variance was conducted to explore the effect of age group differences and MDRS-7 symptom categories on prototypic depression (PHQ-9) and psychological distress (K10). We classified individuals into depression groups using the MDRS-7 in combination with the PHQ-9 based on previous research with groups referred to as: not depressed (PHQ-9 < 10 and MDRS-7 ≤ 5), prototypic depression features (PHQ-9 ≥ 10 and MDRS-7 ≤ 5), mixed features (PHQ-9 ≥ 10 and MDRS-7 > 5) and externalising and male-type features (PHQ-9 < 10 and MDRS-7 > 5). In addition, we used the K10 to determine those individuals suffering a moderate mental illness (K10 ≥ 25) from those without a mental illness (K10 < 25), and current suicidality was ascribed based on scores ≥ 1 on PHQ-9 item 9: ‘Over the past two weeks, how often have you been bothered by thoughts that you would be better off dead, or of hurting yourself in some way?’ Based on these classifications, generalised linear models (GLMs) were used to determine the risk of mental illness and suicidality based on depressive symptom groupings while controlling for previous diagnosis of depression.

An additional GLM examined risk of depression at time 2 (PHQ-9 ≥ 10) as a function of MDRS-7 categories at time 1. Assumptions of GLMs were considered through inspection of scatter plots and histograms of residuals and predicted values, with model results reported as standardised betas.

RESULTS
Sample characteristics
Table 1 presents the characteristics of the participants at time 1 and time 2. As expected, there was a higher proportion of older participants who reported themselves as married/de facto or widowed/divorced/separated, in comparison with younger men. Regarding education, the majority of older participants completed year 11 or below, while the proportion of participants completing a Bachelor’s degree was higher in the younger sample. In addition, household income appeared to be higher in younger compared with older men, consistent with the majority of the older sample reporting themselves as being retired. Comparisons with 2016 Australian Census data indicate that participants in the current study were more likely to be married or in a de facto relationship (63.1% vs 58.1%), more likely to have completed a Bachelor degree level or above (49.8% vs 22.0%) and less likely to be employed full-time (44.5% vs 57.7%) compared with the Australian population. This likely reflects the trend towards older men in the current study. Sample characteristics at time 1 and time 2 were mostly comparable, with a higher proportion of participants at time 2 retired.

Item reduction
Descriptive and relational statistics for each of the MDRS-22 items across younger and older age groups are displayed in online supplemental table 1. For the emotion suppression, alcohol use, somatic symptoms and drug use domains, a single highest scoring item emerged congruent across age groups. For the anger and aggression domain, two different items were retained because of their performance across the age groups. Finally, although two risk-taking items scored equally well in the younger group, only one of these loaded within the older age group, and only this item was retained. This resulted in a total selection of seven items for the short-form scale covering all of the original MDRS-22 domains. The final scale and response format is presented in Appendix A.

Factor analysis of these seven items revealed the presence of a single underlying domain that satisfied criteria determined by the parallel analysis; eigenvalues were required to exceed 1.16. As shown in table 2, all items demonstrated a moderate-to-strong loading on a single underlying factor except for those measuring alcohol and drug use, which loaded moderately. When modelling these seven items using CFA at time 2, the initial solution specifying all items loading on a single latent MDRS-7 factor was not quite adequate ($\chi^2(14)=65.85$, p<0.001, CFI=0.96, TLI=0.94, RMSEA=0.11 (0.08, 0.13), SRMR=0.10). However, allowing the errors of the two items assessing anger and physical aggression to covary resulted in acceptable model fit ($\chi^2(13)=29.04$, p<0.01, CFI=0.99, TLI=0.98, RMSEA=0.06 (0.03, 0.09), SRMR=0.09).

Cut-off scores for the short scale
The proportion of men in each of the different MDRS-7 symptom severity categories is shown in online supplemental figure 1 for the total sample and by age group. As can be seen, older men appear more likely to be in the ‘low’ category of symptoms and less likely to be in the ‘severe’ or ‘extremely severe’ categories compared with younger men. Figure 1 shows the effect of age and MDRS-7 categories on prototypic depression (PHQ-9) and psychological distress (K10). For PHQ-9, there were significant differences between all MDRS-7 groups (F(3, 912)=208.05, p<0.001) and between age groups (F(1, 912)=26.76, p<0.001), with no significant interaction between MDRS-7 and age (F(3, 912)=0.59, p=0.625). For the K10, results were similar: significant differences between all MDRS-7 groups (F(3, 912)=188.95, p<0.001) and between younger and older men (F(3, 912)=33.05, p<0.001), but no interaction between MDRS-7 and age (F(3, 912)=0.44, p=0.719).

Clinical utility of the MDRS-7
The proportion of men according to depressive classification type is shown in online supplemental figure 2. Externalised and male-type depression affected approximately 10% of younger and older men, while prototypic and mixed depressive symptoms were more common in younger men. Table 3 shows the risk of mental illness and suicidality compared
with non-depressed participants within each age group after controlling for a previous diagnosis of depression. All classifications were associated with both outcome measures.

Individuals with mixed symptoms have the highest risk of suicidality and mental illness.

A final GLM considered the likelihood of being classified as depressed at follow-up based on responses to the PHQ-9 at time 2 (ie, score ≥ 10). MDRS-7 category was entered as a predictor controlling for PHQ-9 scores at time 1, previous diagnosis of depression and age. As shown in Table 4, PHQ scores at time 1 were significantly associated with an increased risk of depression at time 2 although age and prior diagnoses were not significantly associated. Those classified as having moderate MDRS-7 symptoms at time 1 were significantly more likely than those in the low symptom category to be classified as depressed at time 2, while the severe and extremely severe categories were not associated with increased risk.

DISCUSSION

Clinical reports and emergent empirical work suggest that men’s depression may be underdetected as a result of prototypic screening tools that may be insensitive to men’s

Table 4  Odds of being classified as depressed at follow-up

<table>
<thead>
<tr>
<th>Outcome: Depressed (PHQ-9 ≥ 10) at time 2 (n=328)</th>
<th>OR</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (older)</td>
<td>1.46</td>
<td>0.69 to 3.09</td>
</tr>
<tr>
<td>Previous depression diagnosis (yes)</td>
<td>1.87</td>
<td>0.88 to 3.99</td>
</tr>
<tr>
<td>PHQ-9 (time 1)</td>
<td>1.24***</td>
<td>1.15 to 1.34</td>
</tr>
<tr>
<td>Moderate (MDRS-7)</td>
<td>3.30**</td>
<td>1.38 to 7.90</td>
</tr>
<tr>
<td>Severe (MDRS-7)</td>
<td>2.00</td>
<td>0.76 to 5.28</td>
</tr>
<tr>
<td>Extremely severe (MDRS-7)</td>
<td>1.64</td>
<td>0.28 to 9.54</td>
</tr>
</tbody>
</table>

Note: Reference category=low symptoms. 
***p<0.001, **p<0.01.

MDRS-7, Male Depression Risk Scale; PHQ-9, Patient Health Questionnaire.
gender role socialisation. The MDRS-22 assesses externalised and male-type symptoms of depression, such as substance misuse, risk-taking and anger. However, in its current 22-item form, it is impractical for rapid use in primary care, particularly when used alongside traditional depression screening tools. The present research aimed to derive a short form of the MDRS-22, examine its psychometric properties and relationships with psychological distress, depression and suicidal ideation in order to demonstrate its utility as a potential screening tool for primary and other healthcare settings.

The short form derived herein comprises seven items, representing one item for each of the original MDRS domains including emotion suppression, risk-taking, substance use, drug use, somatic symptoms and two items for the anger and aggression domain, based on criteria including variability within items, the item’s relationship to its original MDRS domain but also with the overall MDRS score. Of particular importance is our finding that the correlation between the MDRS-7 and the original MDRS-22 was near perfect (r=0.94). Five of the seven items demonstrated moderate-to-strong loadings on a single underlying construct presumed to reflect the male depression phenotype, while two items assessing alcohol and drug use loaded moderately. This likely reflects the reduced variability of participant responses on these items, with most participants reporting that these items applied to them none or a little of the time. However, these loadings still exceeded the minimum recommended factor loading of 0.32. In addition, items that tap these behaviours are important to retain, given that substance use is an important marker of depression and suicidality in men and particularly those who adhere to masculine norms. It is nonetheless important to note that substance use may reflect a comorbidity or maladaptive coping. These are important questions for future research to explore.

In the present study, externalising and male-type symptoms, either alone or in combination with prototypic symptoms, were found to be more common than exclusively prototypic symptoms. Approximately 10% of younger and older men were found to present with uniquely externalising and male-type symptoms, while 38% of younger men and 13% of older men presented with mixed symptoms. These findings are consistent with previous research using the MDRS-22 and highlight the potential utility of the MDRS-7 for detecting additional cases of men at risk. Men with exclusively externalised and male-type depression are a subset who score below threshold on measures such as the PHQ-9. However, an added benefit of using this scale alongside prototypic measures is the ability to detect men presenting with mixed symptomology whose risk of suicide and poor mental health outcomes is significantly elevated. Therefore, the clinical utility of this measure may extend beyond screening and detection and into the therapy setting where it is necessary to determine, monitor and manage differing degrees of suicidality.

Clinical implications

There is an urgent need for health services and providers to use more sensitive diagnostic tools as a means of improving the detection of depression and psychological distress in men and addressing the high rates of male suicide. The use of brief tools such as the MDRS-7 may assist with detecting unique cases of men who would score below threshold on measures such as the PHQ-9. However, an added benefit of using this scale alongside prototypic measures is the ability to detect men presenting with mixed symptomology whose risk of suicide and poor mental health outcomes is significantly elevated. Therefore, the clinical utility of this measure may extend beyond screening and detection and into the therapy setting where it is necessary to determine, monitor and manage differing degrees of suicidality.

Limitations and suggestions for future research

The methodology adopted in this study is not without limitations. The majority of participants were recruited online, which may limit the generalisability of the findings to other populations. Future research should examine measurement invariance according to factors such as education level, income and cultural background. There was also a trend towards older men in the current sample. However, items retained in the MDRS-7 were those that performed best in both younger and older men to ensure the measure was appropriate across the lifespan. Future research should examine the psychometric properties of the MDRS-7 with additional populations, including clinical samples of men across the lifespan presenting to primary care. In addition, as data was self-report,
diagnosis of depression could not be verified at clinical interview. The results of this study would be strengthened by a more rigorous assessment of psychopathology and comorbidity. It is also important to acknowledge that this study used a single item from the PHQ-9 to examine current suicidal ideation. Therefore, there is a need for additional research to examine the relationship between the MDRS-7 and other measures of suicidality, including recent suicide attempt.

CONCLUSION

The present study provides important preliminary information on the development and validation of the MDRS-7. Specifically, this study provides emerging support for the validity and reliability of the MDRS-7 as a measure of externalising and male-type depression symptoms in both younger and older men in terms of its psychometric properties as well as its relationship to prototypic depression symptoms, psychological distress and suicidality. Use of male-specific measures of depression such as the MDRS-7 may improve the detection of depression and suicide risk in men, and adjunctive use (alongside established prototypic scales such as the PHQ-9) may contribute to improved public health outcomes.

Acknowledgements

The authors thank all participants for contributing to this study. The authors would also like to acknowledge Lynn Ward for her valuable suggestions on earlier drafts of the manuscript.

Contributors

DH is the guarantor of the paper, DH, SR and IZ developed the study concept. DH and IZ performed the data analyses. DH drafted the paper and SR and IZ provided critical revisions. All authors approved the final version of the paper for submission.

Funding

DH is supported by an Australian Government Research Training Program Scholarship and a CSIRO Supplementary Scholarship. SR is supported by an NHMRC Career Development Fellowship (GNT1158881) from the Australian Government. These funding bodies had no influence on the design of the study and collection, analysis or interpretation of data or in writing the manuscript.

Competing interests

None declared.

Patient consent for publication

Not applicable.

Ethics approval

Ethics approval was obtained from the University of Adelaide Human Research Ethics Committee and the CSIRO Health and Medical Human Research Ethics Committee (approval number H-2019-109). All participants provided informed consent.

Provenance and peer review

Not commissioned; externally peer reviewed.

Data sharing statement

Data is available on reasonable request. Data is available upon reasonable request.

Supplemental material

This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

Open access

This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

REFERENCES

1. Friedrich MJ. Depression is the leading cause of disability around the world. JAMA 2017;317:1517.

ORCID ID

Danielle Herreen http://orcid.org/0000-0001-6249-8375


JASP Team. JASP (version 0.13.1), 2020


