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Depressive symptoms at old age : proactive management in general practice

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Chapter 5

The influence of the administration method on scores of the 15-item Geriatric Depression Scale in old age

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ABSTRACT

Aim Many rating scales can be self-administered or interviewer-administered, and the influence of administration method on scores is unclear. We aimed to study this influence on scores of the Geriatric Depression Scale (GDS-15), used as a screening instrument in general practice.

Method In two general practices 376 registered patients aged 75 years and older were asked to participate. Exclusion criteria were dementia and current treatment for depression. GDS-15 was administered twice within one month: self-administered by mail, and interviewer-administered during home visits. The sequence of administering the methods was different for the two practices. We analysed differences in total and item GDS-scores.

Results Of 141 subjects that participated (response rate 55%) 59 were men (42%). Mean age was 81.4 years (SD 4.8). When GDS-15 was self-administered, 33 subjects (23.4%) left items unanswered. There were no items unanswered when GDS-15 was interviewer-administered. On average the self-administered total GDS-scores were 0.70 points higher than interviewer-administered scores (95% CI 0.41-0.98), with a large range of variation in the scores (limits of agreement -2.69 to 4.08). Item-item comparisons showed high percentages of agreement. Chance-corrected agreement (kappa) was moderate to fair, but three items showed only slight agreement (kappa values <0.21).

Conclusion Compared to interviewer-administered scores, scores on self-administered GDS-15 were higher. The method of administration should be taken into account when interpreting scores.

INTRODUCTION

Although depressive symptoms in old age have serious negative consequences and effective treatment is available, depressed older subjects are often not treated. Combined screening and treatment programs are being advocated to enhance recognition and to treat depressive symptoms in general practice more adequately.¹

The Geriatric Depression Scale (GDS) is used frequently to screen for depressive symptoms in old age. It was originally developed as a 30-item self rating scale with answers in simple yes/no format.² The shortened 15-item version is considered to be more acceptable as a screening tool, given the shorter administration time.³ In a recent meta-analysis of the diagnostic validity and added value of the GDS in primary care, the GDS-15 had adequate sensitivity and specificity and had good clinical utility as a screening test.⁴ The original instruction of Yesavage et al. was a combination of two methods of administration, stating that ‘patients who cannot complete the questionnaire unaided, have the questions read out to them’.

Little is written about the common practice of method of administration. In a literature search we found 12 studies in primary care in which the GDS-15 was used for screening purposes among persons aged 65 and over. In 9 of these studies the GDS-15 was administered by interview⁵⁻¹³ and in 3 studies by mail.¹⁴⁻¹⁶ Since several studies suggest that the method of administration influences the scores of scales¹⁷⁻¹⁹, we questioned whether self-administration of GDS-15 by mail would give comparable results as interviewer-administration when used to screen older subjects in general practice. Therefore we studied the influence of administering method on item and total scores of the GDS-15 among subjects aged 75 years and older in general practice.

METHODS

Subjects

In two general practices in the Netherlands, in the cities Leiden and Katwijk, registered patients aged 75 years and over were asked to participate. General practitioners (GPs) excluded patients with current treatment for depression, severe cognitive dysfunction (diagnosis of dementia or Alzheimer disease or clinically known MMSE-scores < 19 points), loss of partner or child within the last three months, a life expectancy of less than three months, or patients who do not speak Dutch. All participants were visited at their own home, during which all exclusion criteria were checked. For this study, we further excluded all participants with MMSE-scores below 24 points to minimize the influence of cognitive dysfunction.^{20 21}

Measurement of depressive symptoms

To screen for the presence of depressive symptoms the 15-item Geriatric Depression Scale (GDS-15) was used.³ The answers are in a yes/no response format. The total depression score ranges from 0 to 15 points, with higher scores indicating more depressive symptoms. In this study a score of 5 points or higher was considered as clinically relevant.⁸

The GDS-15 was administered twice: self-administered by mail, and interviewer-administered during home visits. The sequence of administering methods was different for the two practices. In the first general practice, subjects were invited by mail to complete and return the GDS, with one postal reminder after two weeks. After the self-administered GDS-15 was returned by mail, subjects were contacted for a home visit in which trained interviewers administered the GDS-15 a second time. The interviewers were kept blind for the scores on the self-administered GDS. In the interviews all questions were read out to the participant, and on request some additional explanation was given, for example as to time frame ('last month' as stated in the introduction of both written and interviewer version) or reference group ('of same age', not stated in the introduction). In the second general practice, subjects were invited by letter to participate, with one postal reminder after two weeks. When the response card was returned by mail, subjects were contacted for a home visit in which interviewers administered the GDS-15. Two weeks after this home visit, subjects were asked by mail to complete and return the GDS-15 with one postal reminder after two weeks.

There were 5 well-trained interviewers, 3 of them performed the majority of interviews in both practices (for practice A: 41%, 18% and 27%; and for practice B: 31%, 16% and 53%), and 2 others only performed interviews in the first practice (resp. 1% and 13%).

Further measurements

Cognitive functioning was measured using the Mini Mental State Examination (MMSE) during the home visit mentioned above.²² Scores range from 0 to 30 points, with lower scores indicating increasing cognitive impairment. A score below 24 points indicates cognitive impairment.²³ Finally, questions were added about education, income and living situation.

Statistical analyses

To compute the GDS-15 total score, first we interpreted all missing items as 'not-depressed' (0 points). We chose to do so because the GDS-15 total score generally has a low positive predictive value, which would even be lower by interpreting missing items as 'depressed'. Secondly, we computed the GDS-15 total scores using prorating of scores to check whether this would make a difference: for each missing item the average score of completed items per individual was imputed and was added to the total score of completed

items (<http://www.stanford.edu/~yesavage/GDS.html>; visited November 14th 2010). This imputation assumes that items are ‘missing at random’.

We calculated Cronbach’s alpha as a measure of internal coherence of the questionnaire.²⁴ For visual judgment of agreement of the two administering methods, differences in scores were plotted using the Bland-Altman method.²⁵ In this plot the x-axis represents the average between the first and second GDS-15 total score, and the y-axis represents the mean difference in scores for the whole sample with 95% limits of agreement (mean difference $\pm$ 1.96 standard deviation of the mean difference). Since we did not randomly assign sequence of administration (and interviewer) to subjects, we corrected for an uneven distribution of patient characteristics by stratified analysis of difference in scores on age, education, cognitive functioning (MMSE score 24-26 points versus 27-30 points), sequence of administration, time between measurements, and interviewer.

Differences in item scores were analysed with percentage of agreement (unadjusted agreement) and kappa. Kappa takes into account the agreement occurring by chance, thus representing a measure of agreement beyond chance. Kappa varies from -1 to +1, and agreement is considered poor when kappa is less than 0.00, slight 0.00-0.21, fair 0.21-0.40, moderate 0.41-0.60, substantial 0.61-0.80, and almost perfect 0.81-1.00.²⁶

RESULTS

Study population

In general practice A, 69 (32%) out of 218 enlisted older subjects were excluded and 82 out of 149 remaining subjects participated (response rate 55%). In general practice B, 46 (29%) out of 158 enlisted older subjects were excluded, and 59 out of 112 remaining subjects participated (response rate 53%). The mean age of all participants was 81.4 years (SD 4.8), 59 (42%) were male, and 82 (58%) lived alone. Low cognitive functioning according to an MMSE-score below 27 points was found in 23 (16%) subjects. Comparing both general practices, subjects in general practice A were older, and were less well-educated (Table 1). The time between administering the two GDS-15 scales was at average 30 days (95% CI 27-33). For 45% of the subjects time between both tests was within 21 days: in practice A this was 28% and in practice B 68%.

Table 1. Sociodemographics and cognitive functioning of all study subjects (n=141).

	Practice A Self/ Int-adm. (n=82)		Practice B Int./ self-adm. (n=59)		P-value*
Sociodemographics					
- Age: ≥ 80 years	54	67%	25	49%	0.006
- Gender: male	32	39%	27	46%	
- Education: basic schooling only	15	22%	23	56%	0.006
- Income: social security only	11	16%	7	13%	
- Living situation: alone	47	58%	27	50%	
Cognitive functioning:					
- MMSE-scores 24-26	11	13%	12	20%	
- MMSE median score (IQR)		29 (28-30)		28 (27-29)	

* Chi-square test

Table 2. Characteristics of the GDS-15 among all study subjects (n=141).

	Study subjects	
GDS total score		
- self-administered#, median (IQR)	1.0	(0-2)
- interviewer-administered, median (IQR)	0.0	(0-2)
Difference between self-administered and interview-administered GDS total score		
- mean difference (+/- 1.96 S.E. = 95% CI)	0.70	(0.41-0.98)
- mean difference (+/- 1.96 S.D.= <i>limits of agreement**</i>)	0.70	(-2.69-4.08)
GDS total score ≥ 5		
- self-administered#	<i>n=16</i>	11.3% (6.1-16.7)
- interviewer-administered	<i>n=5</i>	3.5% (0.0-7.0)

IQR = interquartile range

95% CI = 95% confidence intervals (mean $\pm$ 1.96 S.E.)

to compute the total score, missing items were interpreted as 'not-depressed'.

** according to Bland and Altman²⁵**Influence of administration method on GDS-scores**

When GDS-15 was self-administered, 33 subjects (23.4%) left some items unanswered, of which 4 subjects (2.5%) left five or more items unanswered. There were no items unanswered when GDS-15 was interviewer-administered.

The internal coherence (Cronbach's alpha) was 0.77 for the self-administered GDS-15 and 0.69 for the interviewer-administered GDS-15. Table 2 shows that the mean difference in total GDS-15 score was 0.70 points (95% CI 0.41-0.98), i.e. total GDS-15 scores were on average 0.70 points higher when GDS was self-administered compared to interviewer-administered. Using a GDS-15 cut-off score of ≥ 5 points, 16 subjects (11.3%, 95% CI

6.1-16.7) were considered to have clinically relevant depressive symptoms on the self-administered GDS-15 and 5 subjects (3.5%, 95% CI 0.0-7.0) on the interviewer-administered GDS. Prorating of scores, instead of interpreting missing items as ‘not-depressed’, gave almost similar results: median GDS total score was 1.0 (IQR 0-2.5) when self-administered, the mean difference between self-administered and interviewer-administered was 0.76 (95% CI 0.47-1.05), and GDS total score ≥ 5 for 12.1% of subjects (n=17).

The Bland-Altman plot (Figure 1) shows a large range of agreement in individual scores (SD = 1.73; 95% limits of agreement -2.69-4.09). The mean difference lies above the null line, indicating a bias towards a higher score on self-administered GDS-15 total score. Eight subjects could be seen as ‘outliers’, as they fell outside the limits of agreement. Their characteristics did not differ from the characteristics of the total group; 7 of these 8 subjects had MMSE-scores above 26. Stratified analyses on age, education, cognitive functioning, sequence of administration, time between measurements, and interviewer showed that GDS-15 total scores were higher when self administered in all defined subgroups (Table 3).

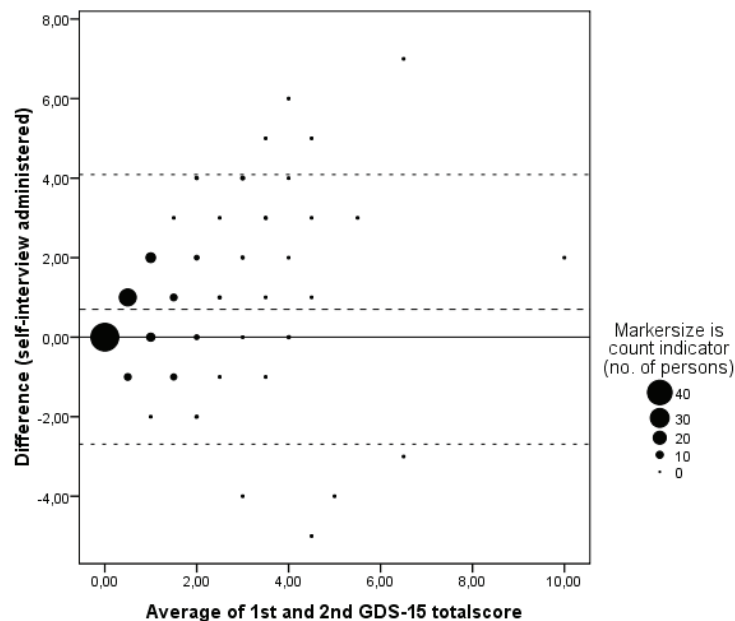


Figure 1. Plot of mean of self-administered and interviewer-administered GDS-15 total score (x-as) and difference between self-administered and interviewer-administered total score (y-axis). Dotted lines are limits of agreement, i.e. 95% confidence interval of mean difference in scores (± 1.96 SD).

Table 3. Stratified analyses for differences in GDS-15 total scores between self-administered and interviewer-administered.

	N	Difference in total score between self-administered and interviewer-administered	
		Mean Difference	(95% CI)
Age group			
-75-79 years	62	0.39	(0.001-0.71)
-80 years and over	79	0.94	(0.52-1.35)
Gender			
-male	59	0.78	(0.43-1.13)
-female	82	0.63	(0.20-1.07)
Education			
-basic level (max 6 years)	38	1.18	(0.55-1.82)
-more then basic level	103	0.51	(0.20-0.83)
Cognitive functioning			
-MMSE-scores 24 to 26	23	1.48	(0.72-2.24)
-MMSE-scores 27 or higher	118	0.54	(0.23-0.85)
Interviewers			
-AC	51	0.55	(0.13-0.97)
-CM	1	3.00	
-EH	24	0.71	(-0.30-1.72)
-IM	55	0.60	(0.20-1.00)
-PT	10	1.70	(0.19-3.21)
Order of administration			
-1 st self, 2 nd interviewer (practice B)	82	0.80	(0.40-1.21)
-1 st interviewer, 2 nd self (practice A)	59	0.54	(0.14-0.94)
Time between test 1 and test 2			
-within 3 weeks	78	0.50	(0.10-0.90)
-longer than 3 weeks	63	0.94	(0.52-1.36)

Table 4 shows for self-administered GDS per item the percentage of missing answers, the percentage of depressive answers per administration method and agreement between administration methods per item. On items 2, 9 and 10 subjects gave far more often a depressive answer when self-administered than when interviewer-administered; for example on item 10 'Do you feel you have more problems with memory than most other people', 30 subjects (21.3%) gave a depressive answer when self-administered, but only 10 subjects (7.1%) when interviewer-administered. Item-item comparisons showed high unadjusted percentages of agreement. Kappa values showed that chance-corrected agreement was moderate to fair, but items 1, 3, and 12 showed poorest agreement (values < 0.21). Kappa is dependent on observed prevalence rates of marginal totals per item, and therefore kappa values inherently vary across the items. Consequently, for these items,

which had a high agreement, one should interpret the value of kappa with caution, because of the so-called high agreement but low kappa paradoxes.²⁷

Table 4. Answers on each GDS-15 item self-administered or interviewer-administered (n=141), with unadjusted percentage of agreement and kappa. Items are ordered by kappa.

		Self-administered		Interviewer-administered	Item-item agreement	
		No answer (%)	Depressive Answer (%)	Depressive Answer (%)	Unadjusted percentage agreement	Kappa *
14	Do you feel that your situation is hopeless?	2	4	5	96	0.48
13	Do you feel full of energy?	7	27	22	79	0.45
11	Do you think it is wonderful to be alive now?	1	2	4	96	0.43
8	Do you often feel helpless [hopeless]?	4	4	2	96	0.43
6	Are you afraid that something bad is going to happen to you?	6	22	18	81	0.41
4	Do you often get bored?	2	1	2	98	0.39
5	Are you in good spirits most of the time?	3	4	1	96	0.28
10	Do you feel you have more problems with memory than most?	7	21	7	82	0.27
2	Have you dropped many of your activities and interests [last month]?	1	16	4	87	0.25
9	Do you prefer to stay at home, rather than going out and doing new things?	7	45	15	65	0.23
15	Do you think that most people are better off than you are?	4	8	4	92	0.21
7	Do you feel happy most of the time?	4	6	5	91	0.21
3	Do you feel that your life is empty?	1	6	8	89	0.16
12	Do you feel pretty worthless the way you are now?	2	9	4	89	0.07
1	Are you basically satisfied with your life?	3	3	7	91	-0.03

* Kappa: poor < 0.00, slight 0.00-0.21, fair 0.21-0.40, moderate 0.41-0.60

DISCUSSION

We questioned whether self-administration of GDS-15 by mail would give comparable results as interviewer-administration when used to screen subjects aged 75 years and over in general practice. On average total depression scores were 0.70 points higher when GDS-15 was self-administered than when interviewer-administered, with a large variation between subjects (limits of agreement -2.69-4.09). On some items subjects gave a depressive answer far more often when self-administered than when interviewer-administered. Apart from three items, item-item comparisons showed fair to moderate agreement.

Our study on the GDS-15 is in concordance with other studies on depression suggesting that self-administered measures give higher scores compared to interviewer-administered measures. Among patients of a medical geriatric unit, the GDS-30 was administered twice within 5 days, once self-administered and once staff-administered in a random sequence showing that total scores on GDS-30 were on average 2 points higher when self-administered.¹⁷ Another study using the Center for Epidemiologic Studies Depression (CES-D) Scale also found higher scores when CES-D was self-administered compared to interviewer-administered, both in lower (< 70 years) and in higher age groups (over 70 years).¹⁹

Why would self-administered scores be higher (i.e. more depressive) than interviewer-administered scores? These findings could suggest that older subjects need help filling out questionnaires, and that this need for help is not restricted to the oldest old. This is affirmed by our finding that many older subjects (23.4%) left items unanswered when GDS-15 was self-administered. But what kind of help is needed? Perhaps it is help in general, such as explaining procedures and solving misunderstandings, which can not be given by mail. On the other hand, the written GDS-15 could be improved. Given the yes-no answering categories subjects are unable to score nuances, which may lead to unanswered items. Subjects may have difficulty to interpret some specific items: e.g. Segulin and Deponte²⁸ suggested rephrasing some items to make them less 'philosophical' and more concrete. Additional written information may be needed concerning the reference group ('of same age') and time frame ('last month'). Another improvement might be to shorten the GDS-15 by removing problematic items. Several shortened versions have been proposed by evaluating item correlations with depression (for example D'Ath et al⁸) or item suitability (for example Jongenelis et al²⁹). Unfortunately, in the GDS-10 and GDS-8 different items were removed and we identified again other items as problematic, except item 9 which is commonly considered as problematic. The GDS-10 and GDS-8 both still include items 1 and 3, which had a low kappa in our analyses.²⁹ Other reasons for systematic discrepancies between the two ways of administration may be rating by proxy (for example through visual problems, cognitive problems, language skills, illiteracy or lack of motivation), or social desirable responding. Some authors suggest that sensitive questions are answered more truthfully when self-administered, since the presence of the interviewer might influence scores towards social desirable answers that is not being depressed.³⁰ If this is true, this would favor a self-administration method. For some populations (such as the oldest old) one might want to include a cognitive test. This is more feasible during an interview, although this can be costly. In a preceding pilot study 12% of elderly mentioned that the postal GDS-15 was filled out by others. We assume that this might lead to somewhat higher but valid scores, as was found in studies developing an informant-version of the GDS.³¹ An alternative, to overcome

missing items and rule out filling out by proxy, is to use the GDS by telephone.³² This might still induce social desirable answers though.

To our knowledge, this study is the first to assess the difference between self-administered and interviewer-administered GDS-15 scores among older subjects aged 75 years and older in the general population. Many GDS validation studies have included patients aged 60 and older, and we showed that the method of administration had influence on scores among the older old. It could be seen as a limitation that the sequence of administration to each subject was not randomly assigned. And we did not always succeed to keep the time between the two tests to a preferred minimum. However, stratified analyses consequently showed higher scores when GDS-15 was self administered. Therefore we expect that scores will be systematically higher when self-administered. Given these limitations, however, we cannot exactly quantify the difference. The interviewers did not restrict themselves to 'read out loud' the GDS, as is often advised. To our opinion, however, it was desirable that they gave synonyms or examples to explain items to enable subjects to choose between 'yes' and 'no'. We advise other researchers to administer the GDS in the same manner.

What are the implications of our findings for research and practice? We consider a mean difference of 0.5 to 1 point on a total of 15-points could be clinically relevant, especially when a strict cut-off point is used to act on. Screening by mail is less costly and may have fewer barriers in revealing sensitive information.³⁰ On the other hand, interviews give no missing answers and help can be offered when questions are not clear to the participant or are misunderstood, resulting in more accurate answers. It is advisable to register whether persons received help. Differences between both administration methods varied a lot between individual older subjects, but we could not identify a specific subgroup for which screening by mail is not applicable or reliable. For epidemiological studies the extra costs of interviewing all subjects may be worthwhile to get more valid data. Perhaps administration of the GDS by telephone is another option.³² In clinical practice the costs may not weigh up to the benefits, for example in a combined screening and treatment program initiated by the GP.³³ In this case a two-step design, firstly mail and secondly an interview among screen positives, can be chosen.

To conclude, our study indicates that the method of administering the GDS should be carefully weighted and reported. It should be taken into account when interpreting scores, for example when comparing studies or choosing a cut-off point.

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