The validation of the Italian version of the GPCOG (GPCOG-It): a contribution to cross-national implementation of a screening test for dementia in general practice

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ABSTRACT

Background: The General Practitioner Cognitive Assessment of Cognition (GPCOG), a brief, efficient dementia-screening instrument for use by general practitioners (GPs), consists of cognitive test items and historical questions asked of an informant. The validity of instruments across different cultures and languages requires confirmation and so the aim of this study was to validate the Italian version of GPCOG (GPCOG-It).

Methods: The validity of the GPCOG-It was assessed against standard criteria for diagnosis of dementia (Diagnostic and Statistical Manual of Mental Disorders – 4th edition) as well as the Clinical Dementia Rating scale. The participants comprised 200 community-dwelling patients aged at least 55 years with (patient group) or without memory complaints (control group). Seven general practitioners were involved. Measurements used were the Cambridge Cognitive Assessment, Mini-mental State Examination with standard (24/25) and Italian cut-off (26/27), Alzheimer’s Disease Assessment Scale-Cognitive scale and Geriatric Depression Scale.

Results: The GPCOG-It, total score and two-stage method, were at least equivalent in detecting dementia to the MMSE using the standard 24/25 or the Italian 26/27 cut-offs. The two-stage method of administering the GPCOG-It (cognitive testing followed by informant questions if necessary) had a sensitivity of 82%, a specificity of 92%, a misclassification rate of 17.4% and positive predictive value of 95%. Patient interviews took less than 4 minutes to administer and informant interviews less than 2 minutes, half the time needed for MMSE administration.

Conclusions: GPCOG-It maintains the same psychometric features and time efficiency as the original English version. Despite methodological limitations (i.e. use of defined samples), the GPCOG-It performed well in detecting clear cut and borderline cognitively impaired patients and can be introduced in the daily practice of Italian GPs.

Key words: dementia screening, general practice, brief psychometric test, mild cognitive impairment

Introduction

The worldwide prevalence of dementia exceeds 24 million and this is set to exceed 40 million by 2020 and 80 million by 2040 (Ferri et al., 2005). Dementia is a tragedy for those affected, their families and for society in general, imposing a significant economic burden on society (Jonsson et al., 2006).

While there is no cure for most forms of dementia, early diagnosis has the potential to ameliorate stress, allows patients and families to prepare for the future, and enables the development of management plans (Ashford et al., 2007). For most patients in most countries, general practitioners (GPs) or primary care physicians are the first health professional that patients or their families contact if concerned about memory loss (Valcour et al., 2000). Even so, the majority of cases of dementia remain undiagnosed in general practice (Brodaty et al., 1994). In response to GPs’ complaints about lack of suitable instruments and time to assess patients’ cognitive...
state (Valcour et al., 2000), an instrument was designed specifically for primary care that was quick, efficient and had psychometric properties at least equivalent to the Mini-mental State Examination (MMSE) (Folstein et al., 1975), the most widely used screening tool for cognitive impairment. The General Practitioner Cognitive Assessment instrument (GPCOG) (Brodaty et al., 2002) was developed for this purpose. Unlike most screening instruments, it combines cognitive testing and informant reports. Additionally, it takes only about four minutes to complete, is easy to learn how to use, requires no specific forms or equipment, has been successfully translated into a number of languages (http://www.gpcog.com.au/) and has diagnostic accuracy equivalent or better than the standard MMSE (Brodaty et al., 2006).

The GPCOG was developed from three principal sources: the Cambridge Cognitive Examination (CAMCOG) (Roth et al., 1988), the Psychogeriatric Assessment Scale (Jorm et al., 1995) and the Instrumental Activities of Daily Living Scale (Lawton and Brody, 1969). It consists of a cognitive section, based on nine items that deal with cognitive functions (score range: 0–9), and an informant section covering six functional items that are clinically relevant and related to pivotal activities of daily living (score range: 0–6). Higher scores indicate better functioning on both subscales. The GPCOG subscales were developed to allow a two-step testing procedure in order to maximize efficiency for GPs. In the first step, the patient’s cognition is tested: a score of 9/9 indicates intact cognitive performance. In both situations, the diagnosis is about 90% accurate, making the second step, informant questioning, unnecessary (Brodaty et al., 2002). However, intermediate scores, i.e. 5–8/9, require an informant interview to clarify whether there is significant impairment. If the informant affirms that there has been no decline in four or more of the six questions, the patient is considered cognitively well, though repeat testing in six months is advised. For patients whose informant affirms lack of decline in three or fewer items (i.e. decline in four or more items), cognitive impairment is diagnosed.

Our aim was to evaluate the psychometric characteristics of the Italian version of the GPCOG (GPCOG-It), in particular as a screening tool capable of correctly identifying patients in need of further evaluation in memory clinics. While the novelty of this tool lies in the inclusion of informant and cognitive testing items, this can give rise to complex issues with respect to the sources of information and culture. For this reason, formal validation in Italian was deemed necessary.

**Methods**

**Implementation of the GPCOG-It from the GPCOG**

The translation of the items was undertaken with particular attention and care being given to rendering the original level of difficulty and the original functional sense and meaning in the Italian wording. The Italian version of the GPCOG fully respects the original format and scoring system.

The validity of the GPCOG-It was assessed against standard DSM-IV criteria for diagnosis of dementia as well as the Clinical Dementia Rating Scale (CDR) (Morris, 1993). Both sets of criteria identify dementia and, moreover, CDR stage 0.5 also allows recognition of questionable dementia or an early transitional state compatible with mild cognitive impairment (Gauthier et al., 2006).

**Sample and setting**

The Ethics Committee of the Faculty of Medicine at the University of Modena approved the research protocol. Written informed consent was obtained from each participant and a family member (as potential caregiver).

The research was conducted on a sample consisting of two groups: a “patient group” and a “control group”. For the “patient group”, 182 community-dwelling individuals complaining of memory impairment were recruited over a 6-month period: 103 were referred to the Dementia Assessment Units by GPs trained in administering the GPCOG-It in their daily practice activity, whereas 79 directly contacted Dementia Assessment Units seeking medical advice. Of these 182 patients, 132 met the inclusion criteria and completed the study (Figure 1). For the “control group”, 78 individuals were recruited by the GPs during daily practice activity from well-known attendees without any subjective memory complaints, after excluding those with known cognitive impairment or comorbidity predisposing to cognitive disorders such as metabolic and cardiovascular diseases. Of these, 68 individuals met the inclusion criteria and completed the study (Figure 1).

All study participants were required to be at least 55 years old, have an available informant and provide informed consent. The following criteria precluded study participation: living in a nursing home, presence of psychiatric disorders requiring treatment, presence of other comorbidity interfering with cognitive performance such as severe chronic heart failure, chronic obstructive pulmonary disease, diabetes, anemia, impairment of auditory or visual perception or low education.
Validation of the Italian version of the GPCOG

Figure 1. Flow diagram of the recruitment of the sample.

Subjects with memory complaints (182)
perceived by himself and/or by an informant

↓

103
referred to DAUs (*) by GPs
after GPCOG-It administration

↓

assessment
(2 different raters blind one vs
the other and the GPCOG-It
results administered by GPs)

↓

26
dementia (DSM IV)
17 withdrawn or did not meet
the inclusion/exclusion
criteria (§)
9 no dementia

↓

132
“patient group”

Subjects without memory complaints

↓

79
directly referred to DAUs

↓

assessment
(2 different raters, blind one vs
the other)

↓

78 (#)
referred to DAUs
(*) by GPs without
GPCOG-It administration

↓

55
dementia (DSM IV)
24 withdrawn or did not meet
the inclusion/exclusion
criteria (§)
15 no dementia

↓

10
withdrawn or did not meet the
inclusion/exclusion criteria (§)

↓

68
“control group”

(*): DAU = Dementia Assessment Unit

(#): GPs were instructed to recruit the persons among their well known attendees without known cognitive impairment or comorbidity predisposing to cognitive disorders.

(§) The following criteria precluded study participation: living in a nursing home, presence of psychiatric disorders requiring treatment, presence of other comorbidity interfering with cognitive performance such as metabolic and cardiovascular diseases (severe CHF, COPD, diabetes, anaemia and so on) impairment of auditory or visual perception or low education level precluding test administration, lack of a reliable informant.
level precluding test administration, and lack of a reliable informant. As we were interested in comparing the GPCOG-It and MMSE, we excluded 24 patients who were found not to have dementia: 15 who contacted the Dementia Assessment Units directly without a GP referral and nine who were seen by their GP and referred to Dementia Assessment Units. Informants, defined as persons who usually lived with the patient or met with her/him at least three times a week, were contacted directly or by phone.

The GPs comprised four males and three females, aged from 38 to 55 years. They were a convenience sample of GPs practicing within Italy’s National Health System, recruited on a voluntary basis. Their characteristics with regard to age, gender, practice population and location (urban or rural) were similar and representative of Italian GPs. All had worked at least five years in general practice, with four having practiced for more than 20 years. The number of patients per practice varied from 800 (for the younger GPs) to 1500 (for the older GPs). The total number of practice-registered patients was 8,738.

The two Dementia Assessment Units (Unità di Valutazione Alzheimer – UVA) were located in a middle-sized industrial city (Modena = 300,000 inhabitants) and in a rural town (Cento, in the Province of Ferrara = 30,000 inhabitants). The Dementia Assessment Units were part of a nationwide program, labeled “Progetto Cronos” (http://www.alzheimer-cronos.org), implemented by the National Department of Health for the screening, diagnosis and treatment of Alzheimer’s disease (AD). The staff employed at the Dementia Assessment Units comprised geriatricians or neurologists, and psychologists, grouped in teams targeted to ensure diagnosis, follow-up, pharmacologic and non-pharmacologic treatment for patients and counseling for caregivers of patients with dementia. All the teams employed at the Units received common basic training to minimize inter-rater bias regarding evaluations and treatment plans; moreover, staff inter-rater reliability was periodically reviewed.

**Measurements and instruments**

All the subjects evaluated by the GPs were asked to make an appointment with the Dementia Assessment Unit, at least one month later. Patients and controls were assessed as follows in the Dementia Assessment Units:

(a) **Cognitive domain:** GPCOG-It (administered only to the subjects not previously tested by GPs), Mini-mental State Examination (MMSE) (Magni et al., 1996), Cambridge Cognitive Assessment (CAMCOG), the cognitive section of the Cambridge Examination for Mental Disorders of the Elderly – Revised (Giannelli et al., 2001), Alzheimer’s Disease Assessment Scale – Cognitive subscale (ADAS-Cog) (Lucca et al., 1994)

(b) **Affective domain:** 15-item Geriatric Depression Scale (GDS) (Sheikh and Yesavage, 1986)

(c) **Clinical staging:** The Clinical Dementia Rating score (Morris, 1993), namely Total (CDR-T) and CDR Sum of Boxes, was applied by two geriatricians (one for each Unit) who were formally trained to apply the procedure as certified by United BioSource Corporation (http://www.unitedbiosource.com). The CDR is a 5-point scale used to characterize six domains of cognitive and functional performance applicable to AD and related dementias: memory, orientation, judgment and problem solving, community affairs, home and hobbies, and personal care. The necessary information to make each rating is obtained only through a semi-structured interview of the patient and a reliable informant or collateral source (e.g. family member) (http://alzheimer.wustl.edu/cdr/default.htm)

(d) **Diagnoses of dementia:** Diagnoses were made for patients who fulfilled DSM-IV criteria as judged by an expert geriatrician or neurologist blind to the GPCOG-It score based on a semi-structured interview of each outpatient and his/her informant, a physical examination and scores on Italian versions of the MMSE, CAMCOG and ADAS-Cog (Dementia Assessment Unit of Modena) and MMSE and ADAS-Cog (Dementia Assessment Unit of Cento). Two thresholds for diagnoses of dementia on the MMSE were used: (i) the standard score of less than 25, and (ii) a score less than 27, as approved by Italian law for free supply of cholinesterase inhibitors (“Progetto Cronos”). As one of the goals of “Progetto Cronos” was to detect the very early stage of dementia, the MMSE cut-off score was set at a higher level (26/27) in order to improve sensitivity with reliance on further assessments to increase specificity (O’Bryant et al., 2008). The CAMCOG cut-off score for diagnoses of dementia was <80 (range 0–105) while for ADAS-Cog the cut-off was >10 (range 0–70).

**Validation**

The GPCOG-It was validated in two ways: (i) against the gold standard of DSM-IV dementia diagnoses (absence / presence) and (ii) assessing the functional stage with the CDR-T (absence / four worsening stages of presence). Dementia on the CDR-T was categorized as absent (CDR-T = 0), questionable or very mild dementia (CDR-T = 0.5), mild (CDR-T = 1) and moderate (CDR-T ≥ 2).

In order to simplify analyses, the 62 mild and 16 moderate cases with dementia were combined, i.e. 78 cases with CDR-T ≥ 1. The GPCOG-It – total score and its cognitive and informant sections – were
Results
Characteristics of the sample
Of the 260 subjects recruited, a total of 200 completed the study (see Figure 1). These comprised 84 males and 116 females with a mean age of 76.1 years (SD 7.2 years; range 56–94 years). Elementary school was completed by 143 subjects (71.5%), middle school by 25 (12.5%) and secondary or university education by 32 (16.0%). Controls were younger and more educated than patients (respectively t = -4.23 and t = 2.84, p < 0.05 for both), whereas no significant difference was found with respect to depression scores and gender distribution (Table 1).

Comparison of cognitive test scores and DSM diagnosis of dementia
Scores on rating instruments of cognition were highly significantly different between the two groups, even when corrected for age and education (Table 1).

The sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV) of the GPCOG-It total, 2-stage, patient and informant sections and MMSE using both cut-points against DSM diagnoses of dementia are shown in Table 2. The different indices were reported with both the misclassification rates and the improvement over chance criterion (IOCC; range 0–1). The MMSE Italian cut-off and the GPCOG-It total showed the highest values of IOCC, very close to 1, corresponding to a perfect classification rate. The informant scale alone did not perform as well as the cognitive tests (Table 2).

The ROC curves showed very similar slopes (Figure 2). The AUCs of the GPCOG-It total, two-stage and cognitive section and of the MMSE, standard and Italian cut-points, did not show any significant difference for any possible comparison when Z scores were applied corrected for Hanley and McNeil criteria (Hanley and McNeil, 1982).

Table 1. Comparison of demographics and cognitive test scores with DSM diagnosis of dementia (variable mean ± SD, unless otherwise stated)

<table>
<thead>
<tr>
<th>CHARACTERISTIC</th>
<th>CONTROLS (68)</th>
<th>PATIENTS (132)</th>
<th>STATISTICS UNCORRECTED FOR AGE AND EDUCATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>72.63 ± 9.03</td>
<td>77.39 ± 6.65</td>
<td>t = - 4.23, p &lt; 0.05 N/A</td>
</tr>
<tr>
<td>Gender</td>
<td>M = 28 F = 40</td>
<td>M = 56 F = 76</td>
<td>χ² = 0.29, n. s.</td>
</tr>
<tr>
<td>Education</td>
<td>7.09 ± 3.07</td>
<td>5.77 ± 3.12</td>
<td>t = 2.84, p &lt; 0.05</td>
</tr>
<tr>
<td>GDS</td>
<td>2.71 ± 3.11</td>
<td>3.12 ± 3.64</td>
<td>t = 1.02, n. s.</td>
</tr>
<tr>
<td>GPCOG-It cognitive sect.</td>
<td>7.65 ± 1.47</td>
<td>2.77 ± 2.21</td>
<td>t = 267.2, p &lt; 0.01 F = 91.36, p &lt; 0.01</td>
</tr>
<tr>
<td>Subjects scoring 5–8/9</td>
<td>38 (55.9%)</td>
<td>36 (27.3%)</td>
<td></td>
</tr>
<tr>
<td>Subjects scoring ≤3</td>
<td>7 (10.3%)</td>
<td>75 (56.8%)</td>
<td></td>
</tr>
<tr>
<td>GPCOG-It informant</td>
<td>5.19 ± 1.2</td>
<td>2.77 ± 1.87</td>
<td>t = 94.03, p &lt; 0.01 F = 21.27, p &lt; 0.01</td>
</tr>
<tr>
<td>Subjects scoring ≤3</td>
<td>7 (10.3%)</td>
<td>75 (56.8%)</td>
<td></td>
</tr>
<tr>
<td>GPCOG-It total score</td>
<td>12.96 ± 2.40</td>
<td>5.51 ± 3.41</td>
<td>t = 257.01, p &lt; 0.01 F = 90.60, p &lt; 0.01</td>
</tr>
<tr>
<td>MMSE</td>
<td>28.16 ± 1.41</td>
<td>20.57 ± 5.14</td>
<td>t = 142.01, p &lt; 0.01 F = 47.66, p &lt; 0.01</td>
</tr>
<tr>
<td>ADAS-Cog</td>
<td>8.45 ± 4.12</td>
<td>26.42 ± 13.02</td>
<td>t = 120.89, p &lt; 0.01 F = 42.06, p &lt; 0.01</td>
</tr>
<tr>
<td>CAMCOG</td>
<td>92.95 ± 5.02</td>
<td>66.98 ± 11.52</td>
<td>t = 176.60, p &lt; 0.01 F = 69.52, p &lt; 0.01</td>
</tr>
</tbody>
</table>

DSM-IV = Diagnostic and Statistical Manual of Mental Disorders, 4th edn; GDS = Geriatric Depression Scale; GPCOG = General Practitioner Cognitive Assessment of Cognition; MMSE = Mini-mental State Examination; ADAS-Cog = Alzheimer Disease Assessment Scale – Cognitive subscale; CAMCOG = Cambridge Cognitive Assessment.
Table 2. Sensitivity, specificity, and area under the curve (AUC) for GPCOG-It cognitive and informant sections, GPCOG–It total score, GPCOG-It “two stage” method and MMSE against DSM-IV dementia diagnosis

<table>
<thead>
<tr>
<th></th>
<th>GPCOG-IT COGNITIVE SECTION</th>
<th>GPCOG-IT INFORMANT SECTION</th>
<th>GPCOG-IT TOTAL SCORE</th>
<th>GPCOG-IT TWO-STAGE METHOD</th>
<th>MMSE STANDARD CUT-OFF</th>
<th>MMSE ITALIAN CUT-OFF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cut-point</td>
<td>7/8</td>
<td>4/5</td>
<td>10/11</td>
<td>24/25</td>
<td>26/27</td>
<td></td>
</tr>
<tr>
<td>Max score</td>
<td>9</td>
<td>6</td>
<td>15</td>
<td>30</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>Number</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>98%</td>
<td>83%</td>
<td>92%</td>
<td>82%</td>
<td>78%</td>
<td>93%</td>
</tr>
<tr>
<td>Specificity</td>
<td>54%</td>
<td>75%</td>
<td>88%</td>
<td>92%</td>
<td>98%</td>
<td>91%</td>
</tr>
<tr>
<td>Positive Predictive Value</td>
<td>0.81</td>
<td>0.86</td>
<td>0.92</td>
<td>0.95</td>
<td>0.99</td>
<td>0.95</td>
</tr>
<tr>
<td>Negative Predictive Value</td>
<td>0.94</td>
<td>0.61</td>
<td>0.88</td>
<td>0.70</td>
<td>0.72</td>
<td>0.87</td>
</tr>
<tr>
<td>Misclassification rate</td>
<td>16.5%</td>
<td>20.0%</td>
<td>9.0%</td>
<td>17.4%</td>
<td>13.5%</td>
<td>7.5%</td>
</tr>
<tr>
<td>Improvement over chance criterion</td>
<td>0.74</td>
<td>0.71</td>
<td>0.83</td>
<td>0.73</td>
<td>0.69</td>
<td>0.81</td>
</tr>
<tr>
<td>AUC</td>
<td>0.96</td>
<td>0.86</td>
<td>0.96</td>
<td>0.96</td>
<td>0.96</td>
<td>0.96</td>
</tr>
<tr>
<td>AUC 95% confidence interval</td>
<td>0.93–0.98</td>
<td>0.81–0.92</td>
<td>0.93–0.98</td>
<td>0.94–0.98</td>
<td>0.93–0.98</td>
<td>0.94–0.98</td>
</tr>
<tr>
<td>Standard error of AUCs</td>
<td>0.013</td>
<td>0.028</td>
<td>0.013</td>
<td>0.011</td>
<td>0.013</td>
<td>0.012</td>
</tr>
</tbody>
</table>

GPCOG = General Practitioner Cognitive Assessment of Cognition; MMSE = Mini-mental State Examination; AUC = area under the curve.

Figure 2. The Receiver Operating Characteristics (ROC) curves for the GPCOG-IT and MMSE.

Comparison of cognitive test scores and clinical staging (CDR)

Of the 132 patients referred to the Dementia Assessment Units who received DSM-IV diagnoses of dementia, 54 were rated as having very mild dementia on the CDR-T, and 78 mild-moderate dementia.

None of the 68 controls received DSM IV diagnoses of dementia but eight had CDR-T scores of 0.5 as some limited functional decline was noted.
Validation of the Italian version of the GPCOG

Despite normal neuropsychological assessment and no memory impairment as confirmed by the informants. Analyses were conducted with all 68 controls and repeated after reclassifying these eight subjects as CDR-T = 0.5 as they were considered a possible confound.

In order to examine the influence of severity of dementia on the performance of the cognitive tests applying an ANOVA, we compared those with CDR-T = 0, 0.5 and ≥1 performing a two-step analysis. At first, the 68 controls and 54 with very mild dementia (CDR-T = 0.5) original groups were considered and then the potentially confounding eight controls with CDR-T = 0.5 were moved into the very mild dementia group [i.e. CDR 0 (68 → 60 Ss.) to CDR 0.5 (54 → 62 Ss.).] The difference between groups for all possible comparisons was tested by means of an LSD-post-hoc analysis, taking into account a 5% level of significance.

In the first step, each CDR-T group proved to differ significantly from the others with respect to the GPCOG-It indices, MMSE, ADAS and CAMCOG. These differences persisted even when corrected for age and education with a MANCOVA. The second step analyses, performed with the confounding factor moved from the CDR 0 group to CDR 0.5 (Table 3), showed the same pattern of results.

Relations between GPCOG-It indices and psychometric and staging validated measures

The levels of correlation were computed between the three sections of GPCOG-It and cognitive and functional measures. The GPCOG-It total score was highly and significantly correlated with all the cognitive (Pearson’s correlations: MMSE = 0.73, CAMCOG = 0.84, ADAS-Cog = −0.70) and functional measures (CDR-Sum of Boxes = −0.75). Also, the GPCOG-It subscales were highly and significantly correlated with all the cognitive (cognitive subscale: MMSE = 0.68, CAMCOG = 0.84, ADAS-Cog = −0.71; informant subscale: MMSE = 0.63, CAMCOG = 0.47, ADAS-Cog = −0.55) and functional measures (cognitive subscale = CDR-Sum of Boxes: −0.75; informant subscale = CDR-Sum of Boxes: −0.68).

Time for administration

The time required for completion of the GPCOG-It was significantly lower than for the other cognitive scales: GPCOG-It Cognitive = 2.97 ± 0.20 minutes; GPCOG-It Informant = 1.53 ± 0.22 minutes; GPCOG-It Total Score = 4.45 ± 0.42 minutes; MMSE = 8.9 ± 1.1 minutes; ADAS-COG = 18.4 ± 7.2 minutes.

**Table 3.** Comparison of demographics and cognitive test scores with clinical staging of dementia

<table>
<thead>
<tr>
<th>CHARACTERISTICS</th>
<th>CDR-T 0 (N = 64) (§)</th>
<th>CDR-T 0.5 (N = 54) (§)</th>
<th>CDR-T ≥ 1 (N = 60) (§)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>76.35 ± 7.71 (c)</td>
<td>74.63 ± 12.30 (c)</td>
<td>73.21 ± 13.20 (c)</td>
</tr>
<tr>
<td>Gender</td>
<td>M 53 (85.4%)</td>
<td>M 37 (68.9%)</td>
<td>M 36 (60.0%)</td>
</tr>
<tr>
<td>Affect (GDS)</td>
<td>7.0 ± 3.08 (c)</td>
<td>7.05 ± 3.08 (c)</td>
<td>7.22 ± 3.08 (c)</td>
</tr>
<tr>
<td>Education</td>
<td>7.05 ± 3.08 (c)</td>
<td>5.67 ± 3.54 (c)</td>
<td>2.92 ± 1.89 (c)</td>
</tr>
<tr>
<td>GPCOG-It cognitive section</td>
<td>7.80 ± 1.43 (c)</td>
<td>5.42 ± 1.15 (c)</td>
<td>3.68 ± 1.33 (c)</td>
</tr>
<tr>
<td>GPCOG-It informant section</td>
<td>5.42 ± 1.15 (c)</td>
<td>3.68 ± 1.33 (c)</td>
<td>2.08 ± 1.89 (c)</td>
</tr>
<tr>
<td>GPCOG-It total score</td>
<td>13.47 ± 2.17 (c)</td>
<td>7.94 ± 1.33 (c)</td>
<td>5.20 ± 1.66 (c)</td>
</tr>
<tr>
<td>MMSE</td>
<td>8.38 ± 2.46 (c)</td>
<td>5.20 ± 1.66 (c)</td>
<td>2.96 ± 1.33 (c)</td>
</tr>
<tr>
<td>ADAS-Cog</td>
<td>3.84 ± 5.07 (c)</td>
<td>2.08 ± 1.89 (c)</td>
<td>1.22 ± 0.88 (c)</td>
</tr>
<tr>
<td>CAMCOG</td>
<td>93.4 ± 5.07 (c)</td>
<td>93.4 ± 5.07 (c)</td>
<td>93.4 ± 5.07 (c)</td>
</tr>
<tr>
<td>CDR-Sum of Boxes</td>
<td>0.21 ± 0.41 (c)</td>
<td>0.21 ± 0.41 (c)</td>
<td>0.21 ± 0.41 (c)</td>
</tr>
</tbody>
</table>

Note: CDR-T 0.5 group combines both the 54 patients affected from “very mild dementia” and the 8 normal ones reporting functional but not memory impairment according to the CDR-T scoring rules (http://alzheimer.wustl.edu/cdr/default.htm).

CDR-T scoring rules: (a) CDR-T = Clinical Dementia Rating – Total Score; (b) CDR-T = Alzheimer Disease Assessment Scale – Cognitive subscale; (c) GPCOG = Geriatric Depression Scale; (d) GPCOG = General Practitioner Cognitive Assessment of Cognitions; MMSE = Mini-mental State Examination; ADAS-Cog = Alzheimer Disease Assessment Scale – Cognitive subscale; CAMCOG = Cambridge Cognitive Assessment.
Discussion

The data obtained in this study strongly indicate that the GPCOG-It maintains the original psychometric characteristics of the English-language version. The translation work has led to a culturally competent tool, able to provide data that may be compared to those in different cultural settings, such as English-speaking countries and other countries where GPCOG validated translations are available (http://www.gpcog.com.au). Like the original, the GPCOG-It proved to be a valid, efficient and rapid instrument for dementia screening. Moreover, the gain in the levels of both sensitivity and specificity obtained with the original GPCOG two-stage mechanism, was evident in the results of the Italian version as well.

The performance of the GPCOG-It was independent of the patient’s gender and affect, and relatively independent of age and educational level. Indeed, the cognitive section score differentiated dementia diagnosis and stage when controlled for age and education. This characteristic makes the instrument relatively free from the pitfall of a higher educational level being able to mask the clinical expression of a higher degree of neurodegeneration (Garibotto et al., 2008). In order to reduce misclassification rates, it has been proposed that scores adjusted for educational level be used for cognitive testing; such scores have been reported to perform worse than unadjusted ones by some (Teresi, 2007) but not others (Anderson et al., 2007). Among the strengths of the test, we emphasize that GPs require very little training to administer it. Moreover, the test is simple for testers to remember and repeat. The GPCOG-It requires only 3–5 minutes in an office or home setting where the printed form or just a pencil and paper can be used if other forms are unavailable. Further, the two-stage procedure (Denny et al., 2000) optimizes accuracy and efficiency as needed in the screening process. We concede that, as participants in this study were chosen because of their status as having dementia symptoms and diagnoses or not, the psychometric properties of the instrument including PPV and PNV are likely to differ if tested on consecutive general practice attendees where the prevalence of dementia is lower.

The GPCOG is one of a number of brief screening tests for dementia (Lorentz et al., 2002). Its uniqueness lies in its combination of both cognitive testing and informant reports as sources of information in one scale, its design specifically for primary care and its two-stage administration to enhance efficiency. The addition of informant data to cognitive testing has been reported to enhance accuracy of dementia screening by several authors (Mackinnon and Mulligan, 1998; Jorm, 2003; Kemp et al., 2002) though not by all (Knafelc et al., 2003) and has the advantage of being race, gender and education neutral, unlike most performance-based screening tests. The GPCOG-It strategy is aligned with the two-stage mechanism recommended for screening in the way that it is based on two different sources of information: the patient and the informant.

The expertise of GPs in testing procedures may not be particularly high if we consider the figures of incidence and prevalence of dementia in Italy, together with the characteristics of the patients in the practice of an Italian GP. As the maximum number of patients in the practice of an Italian GP is set at 1500, the GP does not have occasion to diagnose many cases of dementia (prevalence: 28 cases – 1.8% of GP patients: females 57%, males 43%; incidence: 4–5 new diagnoses/year – males 1%; females: 1.3%) (De Ronchi et al., 2005). However, owing to their long relationships with patients and their families, GPs are in a favorable and unique position to collect information to provide an evaluation of a patient’s cognitive and functional profile. This information is considered pivotal for both the diagnostic process and the analysis of treatment effects (Nygård, 2003). The GPCOG-It fully performs the information feature, making it very attractive as a psychometric tool for GP daily practice. Even though our data indicate similar diagnostic properties for the GPCOG-It total score when compared with MMSE, using either the standard or Italian cut-off score, the efficiency of the GPCOG-It makes it appealing for Italian GPs to use.

We conclude that the GPCOG-It is a suitable tool for GPs to use to screen for dementia. Our results are in accordance with a recent review (Milne et al., 2008) that found GPCOG to be one of the best tools among those proposed for screening for dementia in primary care. Regardless of the controversy about “case-finding” detection versus “cognitive screening of elderly outpatients” (Ashford et al., 2007), the simplicity and rapidity of GPCOG-It testing can improve the ability of GPs to evaluate their patients during their routine practice activity.

Conflict of interest

None.

Description of authors’ roles

Alessandro Pirani, Henry Brodaty and Mireco Neri designed the study and wrote the paper. Alessandro
Pirani and Mirco Neri supervised Emilio Martini, Davide Zaccherini and Francesca Neviani in collecting the data. Mirco Neri and Henry Brodaty developed the statistical design of the study, and Emilio Martini carried out the statistical analysis.

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