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A systematic review of the validity and reliability of apathy scales in neurodegenerative conditions

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Abstract

Background: There are several scales used to detect apathy in disease populations. Since apathy is a prevalent symptom in many neurodegenerative diseases, this is an especially important context in which to identify and compare scales.

Aims: To provide an overview of apathy scales validated in generic and specific neurodegenerative disease populations, compare validation studies’ methodological quality and the psychometric properties of the validated apathy scales.

Methods: A systematic review of literature was conducted of articles published between 1980 and 2013. The final articles selected for review were rated on methodological quality and the psychometric properties of the scales used were interpreted.

Results: Sixteen articles validating apathy scales were included in the review, five in a generic neurodegenerative sample and eleven in specific neurodegenerative samples. The methodological quality of specific studies varied from Poor to Excellent. The highest quality, which had psychometrically favourable scales, were the Dementia Apathy Interview and Rating and the Apathy Evaluation Scale- Clinical version in Alzheimer’s Disease and the Lille Apathy Rating Scale in Parkinson’s disease. Generic neurodegenerative disease validation studies were of average methodological quality and yielded inconsistent psychometric properties.

Conclusions: Several instruments can be recommended for use in some specific neurodegenerative diseases. Other instruments should either be validated or developed to assess apathy in more generic populations.

Key words: apathy, rating scales, psychogeriatrics, dementia, motor disorders

Introduction
Historically, apathy is reported to have undergone a transformation in its definition. Originally, rooted in *apatheia*, meaning without (*a-*) passion (*-pathos*), it was regarded as a virtue in Greek Stoic philosophy, allowing individuals to be more objective. However, the modern meaning of apathy has come to be regarded as undesirable and, as a part of the psychiatric field, can be a syndrome or symptom relating to disease. Clinically, as a behavior, apathy is most often characterised as a deficit in motivation, which is displayed through reduced interest, emotions and goal directed behaviors (Marin, 1996). Apathy has been found to have a variable relationship with various symptoms and experiences, such as depression and boredom. As a symptom, apathy has a variable association with depression dependent on the neurodegenerative disease (Levy *et al.*, 1998). However, depression is expressed as negative and, in the case of bipolar depression, extremely positive affect (Levy *et al.*, 1998), whereas, apathy is observed as emotional neutrality, where neither positive nor negative emotions are observed. Boredom, on the other hand, is a widespread human experience that is linked, through motivational elements, to apathy. However, research has shown them to be statistically distinct when measured by scales (Goldberg *et al.*, 2011). This is due to apathy scales being designed to measure apathy as a symptom rather than a domain of normal motivational behavior.

Apathy has been frequently observed as a symptom of neurodegenerative diseases (for review see van Reekum, Stuss and Ostrander, 2005; Chase, 2011) and is assessed by a variety of methods. Both clinically and in research, these methods often range from administration of measurement scales to observational ratings of behaviors by an experienced or trained individual. While specific methods of assessing apathy have guidelines to ensure that it is administered in a standardised
fashion, the problem is that there are many different methods to measure apathy. This creates uncertainty of which method would be most appropriate to measure apathy in a given pathological population. The utility of these methods are often based on previous studies, which have validated the measure in a specific pathological population. However, these methods and the studies validating them are seldom compared within literature.

Since apathy is a common symptom in neurodegenerative disease, this is an important area in which to produce a comprehensive review of how apathy is measured both generally and for specific neurodegenerative diseases. It would also be beneficial to compare the validity and reliability of assessment methods, so that the highest quality, well-validated and reliable method can be chosen for a specific patient group.

**Apathy in neurodegenerative disease:**

Apathy is common in all types of dementia (Clarke et al., 2008) and most prominent in both Alzheimer’s disease (AD) and vascular dementia (VaD; Jonsson et al., 2010). In both small and large vessel VaD, apathy is a consistent symptom with negative prognostic and treatment implications (Staekenborg et al., 2010). In AD, apathy was found to occur in 61% to 92% of patients (e.g. Landes et al., 2005) with an almost equally high prevalence in frontotemporal dementia (FTD; e.g. Mendez et al., 2008). However, recent research has found differing apathy profiles which can distinguish between AD and FTD (Quaranta et al., 2012). This distinction between AD and FTD was also observed in functional impairments of activities of daily living, which could be associated with apathy level (Mioshi et al., 2009).
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Apathy is also a prominent symptom in amyotrophic lateral sclerosis (ALS; Grossman et al., 2007; Girardi et al., 2011), the most common form of motor neurone disease (Goldstein and Abrahams, 2013). A large cohort study found that apathy was the most common behavioral symptom in ALS, occurring in 31% of patients (Witgert et al., 2010). Recent neuroimaging research in ALS has shown evidence of neuroanatomical correlates relating to apathy of abnormalities in the anterior cingulate cortex (Woolley et al., 2011). In a recent review of social cognition in ALS apathy was observed as the most prevalent, overlapping behavioral change in both ALS and FTD (Abrahams, 2011).

In other motor disorders such as Parkinson’s disease (PD), over a third of patients have been found to exhibit apathy (Pedersen et al., 2009). Apathy may be predictive of cognitive decline and even dementia in PD (Dujardin et al., 2009) but may also have marked variability in its relation to the clinical presentation of PD (Dujardin et al., 2007).

Apathy is also a persistent and progressive symptom across all stages of Huntington’s disease (HD; Thompson et al., 2012) with a variable prevalence reported between 34% and 76% associated with stage of disease (van Duijn et al., 2007). A study comparing HD gene mutation carriers versus non-carriers found that the latter had 0% prevalence compared to carriers who had 32% prevalence (van Duijn et al., 2010). Functional decline was also found to be greater in HD patients with apathy, independent of motor or cognitive symptoms (Hamilton et al., 2003).
Due to this variable prevalence of apathy in these neurodegenerative diseases, this review focuses on apathy as a symptom of these specific diseases.

**Emerging concepts:**

Apathy is conceptualised by the measurement scales associated with it. It is originally defined by a substructure of cognition, behavior and emotion (Marin et al., 1991) but is measured as singular concept, with specified clinical cut-offs dependent on the scale.

A previous review by Clarke et al. (2011) looked at all the different measurement methods to assess apathy in research and clinical practice and found that most used rating scales or scale-based interviews. These scales were self-assessments, informant-based or clinical semi-structured interviews for patients or informants. The studies reviewed included a range of pathological populations with variable aetiologies (Clarke et al., 2011). However, Clarke et al.’s review did not directly examine apathy as a symptom of specific neurodegenerative diseases and was not a systematic review of literature. The present review follows the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA; Moher et al., 2009) criteria to methodologically identify the quality of existent apathy scales and define which scales have been validated in neurodegenerative diseases. As apathy occurs as a symptom in a majority of neurodegenerative diseases (for review see van Reekum, Stuss and Ostrander, 2005; Chase, 2011) it is important to examine methods of apathy assessment that can identify apathy as a symptom in any neurodegenerative disease. Furthermore, as previously mentioned, patients with specific neurodegenerative diseases, such as HD and AD, exhibit apathy in variable prevalence. So it is important
to classify scales that were specifically designed, and therefore only validated, in specific neurodegenerative disease so that they may be used as such in the future. This will allow for identification of apathy scales that are neurodegenerative disease generic and specific. This will result in recommendations for suitable apathy scale choices to be made in future research and clinical practice in a disease group where apathy is a prevalent symptom.

**Quality, Validity and Reliability:**

The methodology and the measures used within a research study are important determinants of article quality. Individual articles employ different patient groups, inclusion and exclusion criteria, index tests and reference or gold standards of diagnosis, all of which are described with different levels of detail. Due to this variation, we considered it important to use a standardized tool for quality assessment of each study.

In addition, we considered the psychometric properties reported by each study of the specific apathy instrument, which are commonly presented in terms of validity and reliability. Validity is best described as accuracy of an instrument, and can be defined through content, concurrent, convergent, congruent and discriminant or divergent. Content validity relates to the design of a measure and whether the instrument is measuring what it is aiming to measure. Concurrent, convergent or congruent validity are often grouped together as measuring construct validity and measure whether the instrument correlates with other well-validated methods of detection. This method of detection is known as a gold or reference standard. In terms of apathy, a gold standard would be the diagnosis of apathy by one or two trained
psychiatrists via observation, a structured or semi structured clinical interview. Discriminant or divergent validity is also an important psychometric property and is where two components, or constructs, that are theoretically unrelated are actually shown to dissociate. With symptomatic apathy, this behavioral component is often depression (Levy et al., 1998), which is due the emotional nature of patients with depression and, conversely, the emotionally neutral nature of apathetic patients. This apathetic component also differentiates it from exhilarating happiness observed in elation, the uncomfortable uneasiness of anxiety and excessive agitation associated with irritability. These behavioral components can also be used to determine discriminant validity.

Reliability is best defined as the consistency of the measure. This relates to the replicability of findings independent of how accurate they are. This can be determined through between-item correlations of the specific measure, which is known as internal consistency reliability and usually measured by Cronbach’s alpha (Cronbach, 1951). Test-retest reliability is another form where the relationship between responses to the measure over several different time periods is recorded and correlated. Another form of reliability is inter-rater based, which examines whether two, or more informants or interviewers responses or scores agree.

Both validity and reliability are important determinants of an assessment scale’s utility and a good apathy measure should embody both of these characteristics measured in the same sample. Therefore, the focus of this review will be on studies that examine both the validity and reliability of scales. However, there are no gold
standards of interpretation regarding validity and reliability psychometric statistics, therefore previously used guidelines were adopted to categorise them.

**Study aims:**

Based on the above points and issues, the aims of this review are:

1. To determine apathy assessment scales that have been validated in neurodegenerative populations
2. To determine neurodegenerative generic and neurodegenerative specific apathy assessment scales
3. To quantify the methodological quality of the apathy assessment validation studies
4. To compare reliability and validity statistics of each apathy assessment scale as to establish suitable neurodegenerative generic and specific apathy assessment methods

**Methods**

This systematic review was conducted following the PRISMA guidelines (Moher *et al.*, 2009). The competed checklist can be found in the Supplementary materials (see figure S1).

**Eligibility criteria**

Only articles, abstracts and letters limited to English language publications between 1980 and 2013 containing human participants, aged 18 or older were screened.
Information Sources

The main search was conducted on 31 December 2013. The electronic databases used were EMBASE, PSYCHINFO, MEDLINE and PUBMED.

Search criteria

Through the use of Boolean operators, the database searches were restricted to an apathy term (‘Apathy’ or ‘Motivation’ or ‘Amotivation’), neurodegenerative disease term (‘Dementia’ or ‘Alzheimer Disease’ or ‘Alzheimer’s Disease’ or ‘Multiinfarct Dementia’ or ‘Vascular Dementia’ or ‘Frontotemporal Dementia’ or ‘Semantic Dementia’ or ‘Parkinson Disease’ or ‘Parkinson’s Disease’ or ‘Motor Neuron Disease’ or ‘Amyotrophic Lateral Sclerosis’ or ‘Huntingtons Disease’ or ‘Huntingtons Chorea’), a scale term (‘Rating Scale’ or ‘Rating Scales’ or ‘Summed Rating Scale’ or ‘Psychological Rating Scale’ or ‘Psychiatric Status Rating Scale’ or ‘Questionnaire’ or ‘Questionnaires’ or ‘Measurement’) and a validity or reliability term (‘Reliability’ or ‘Validity’ or ‘Test Reliability’ or ‘Interrater Reliability’ or ‘Statistical Reliability’ or ‘Test Validity’ or ‘Statistical Validity’).

Study selection

After applying the above mentioned search criteria and eligibility criteria, duplicate articles were removed. Study selection was conducted in two main stages following the electronic search. In the first stage, two authors (R.R. and C.H.) went through the titles and abstracts independently, selecting which abstracts to include in the next stage, exclude from the next stage and which were ambiguous or disagreed upon by the aforementioned two authors. Inclusion was based on the following: study
involved a neurodegenerative disease population, used scalar or quantifiable methods of assessing apathy levels, and recorded both validity and reliability statistics. The ambiguous or disagreed upon abstracts were then forwarded to the third author (J.S.) who made suggestions, of inclusion, exclusion or examination of whole articles.

In the second and final stage, the selected abstracts were then examined as full articles by authors R.R. and C.H., independently. These were classed as include in the review, exclude from the review and ambiguous or disagreed upon by authors R.R. and/or C.H. Articles that were included in the review had to meet the same criteria as in the first stage but authors were additionally asked to record the neurodegenerative disease assessed, apathy assessment methods used and validity and reliability statistics produced. In all cases, the authors had to provide reasons for exclusion or ambiguity. Advice from the third author, J.S., was sought once again for ambiguous or disagreed upon articles. A consensus had to be reached for articles that were to be used in the review.

Neurodegenerative disease generic and specific study grouping

The final articles were then grouped based on whether the study population assessed for apathy was neurodegenerative disease generic (e.g. containing a mixture of AD and PD patients) or specific (e.g. containing only AD patient) as to determine the utility in each group.

Quality Assessment

The final articles selected for review were subjected to the Quality Assessment of Diagnostic Accuracy Studies (QUADAS) checklist (Whiting et al., 2003, see Table
1). This was used to determine the quality of each article. The target condition was apathy, the index test was the method by which apathy was assessed and the reference standard was the method by which apathy was diagnosed and, therefore, the validation method for the index test.

Items 1 to 5, 7 to 9 and 13 to 14 from the QUADAS were found to be applicable for determining quality of methodology and used for this particular review (see Table 1). Some QUADAS items (6 and 10 to 12) were not included due to their lack of applicability for assessing bias in psychiatric symptom scale validation studies, such as apathy. Items 1 and 2 relate to variability of the studies in the sample assessed and which impacts on the generalizability of findings to the study population. Items 3 to 5, 7 and 14 examine methodological rigour and bias, which influences the validity of the study results and, therefore, the validity of the scale. Finally, items 8, 9 and 13 determined the quality of reporting of methodology. Items were answered ‘yes’, ‘no’ or ‘unsure’ after close examination of the articles by author R.R.
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Table 1. QUADAS items for quality review

<table>
<thead>
<tr>
<th>Items</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Was the spectrum of patients representative of the patients who will receive the test in practice?</td>
</tr>
<tr>
<td>2. Were selection criteria clearly described?</td>
</tr>
<tr>
<td>3. Is the reference standard likely to correctly classify the target condition?</td>
</tr>
<tr>
<td>4. Is the time period between reference standard and index test short enough to be reasonably sure that the target condition did not change between the two tests?</td>
</tr>
<tr>
<td>5. Did the whole sample or a random selection of the sample, receive verification using a reference standard of diagnosis?</td>
</tr>
<tr>
<td>6. Was the reference standard independent of the index test (i.e. the index test did not form part of the reference standard)?</td>
</tr>
<tr>
<td>7. Was the execution of the index test described in sufficient detail to permit replication of the test?</td>
</tr>
<tr>
<td>8. Was the execution of the reference standard described in sufficient detail to permit its replication?</td>
</tr>
<tr>
<td>9. Were uninterpretable/intermediate test results reported?</td>
</tr>
<tr>
<td>10. Were withdrawals from the study explained?</td>
</tr>
</tbody>
</table>

Target condition= Apathy; Index test= Apathy assessment method; Reference standard= Apathy diagnosis method

There is no standardized scoring system for the QUADAS. Therefore, we categorised each study based on the number of QUADAS items that were answered ‘yes’ to. The categorisation was as follows; 9 or 10 items was classed as Excellent, 7 or 8 was Good, 4 to 6 was Adequate, 2 to 3 items was Poor and 0 to 1 was considered Unacceptable.

Interpretation of psychometric properties

We have provided a table of guidelines for interpreting internal consistency and correlation coefficients in relation to validity and reliability statistics (see table S1). This was adapted based on Hermans, van der Pas and Evenhuis’ (2011) review paper of anxiety scales. However, we have updated the interpretation of internal
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consistency (George and Mallery, 2003) and Kappa statistic (Viera and Garrett, 2005).

Results

Systematic review

Figure 1 illustrates the review process in a flowchart. From the databases listed and using the search criteria specified earlier, a total of 168 studies were identified. When the eligibility criteria were applied prior to the first stage, 21 articles were removed, leaving 147 articles. After identifying and removing 31 duplicate articles, this left 116 studies eligible for the first stage of study selection; title and abstract screening. These included a variety of neurodegenerative populations that assessed apathy using various methods.
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16 hits PSYCHINFO
23 hits EMBASE database search
94 hits MEDLINE database search
35 hits PUBMED database search

168 hits from database searches

21 articles removed (non-English, non-human, participants age 19+ published before 1980)

147 articles remain

31 duplicate articles removed

116 study articles

28 articles retrieved for detailed evaluation

88 articles excluded based on Title/Abstracts:
- 47 not a reliability/validity studies
- 34 not relating to apathy measurement
- 4 not relating to apathy measurement
AND not a neurodegenerative population study
- 2 not a neurodegenerative population study
- 1 previous review paper in the field

116 study articles

28 articles retrieved for detailed evaluation

14 articles excluded:
- 5 only reported validity
- 5 only reported reliability
- 3 not apathy scales
- 1 insufficient patient information

Further 10 articles excluded:
- 6 only reported reliability
- 2 no validity and reliability
- 1 only reported validity
- 1 no neurodegenerative patients

12 extra articles identified:
- 11 from reference lists
- 1 from Google Scholar

16 included in review

Figure 1. Flowchart of review process
In the first stage of study selection, of the 116 studies, 88 were not relevant or excluded: 47 were not reliability/validity studies, 34 were not studies that related to apathy measurement, four were not studies that related to apathy measurement and were not conducted in a neurodegenerative population, two were not conducted in a neurodegenerative population and one was a previous review paper in the field (Clarke et al., 2008). This left 28 studies to be included for detailed evaluation. A further 12 articles were identified through Google Scholar and reference lists and included for detailed evaluation, resulting in a total of 40 studies for detailed evaluation.

The second stage of study selection, of the 40 studies, 24 articles were excluded: 11 only reported validity for the apathy measurement, six only reported reliability for the apathy measurement, three were not apathy scales, two did not report validity and reliability of the apathy measurement, one had insufficient patient information and one was not conducted in a neurodegenerative population.

Table 2 shows, in total, 16 studies were included in the review. Seven of these articles validated English versions of a scale (Starkstein et al. 1992, Cummings et al. 1994, Kaufer et al. 2000, Strauss et al. 2002, Clarke et al. 2007, Zahodne et al. 2009, de Medeiros et al. 2010).

The review of methodological quality of each study by the chosen QUADAS criteria is shown in Table 2. There was a range of Poor to Excellent quality studies with the modal quality rating being Adequate.
No studies stated explanations for patient withdrawals; a large proportion of which were unclear about their withdrawals and the remainder did not state any explanations. However, almost all of the studies used a reference standard that was independent of index test for their validation study (see figure S2). There were five studies that examined apathy scales or subscales in a generic neurodegenerative disease sample and 11 that examined them in specific neurodegenerative disease samples.
Table 2. Methodological quality ratings of review studies

<table>
<thead>
<tr>
<th>Author (Year)</th>
<th>Instrument</th>
<th>Generic/Specific</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>13</th>
<th>14</th>
<th>Overall Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clarke et al. (2007)</td>
<td>Self, Informant and Clinical Apathy Evaluation Scale (AES-S/I/C)</td>
<td>Generic</td>
<td>N</td>
<td>U</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>Adequate</td>
</tr>
<tr>
<td>Cummings et al. (1994)</td>
<td>Neuropsychiatric Inventory- Apathy (NPIa)</td>
<td>Generic</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>N</td>
<td>Adequate</td>
</tr>
<tr>
<td>de Medeiros et al. (2010)</td>
<td>Neuropsychiatric Inventory- Clinical Rating Scale- Apathy (NPIa-C)</td>
<td>Specific</td>
<td>N</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>Adequate</td>
</tr>
<tr>
<td>Dujardin et al. (2008)</td>
<td>Informant based Lille Apathy Rating Scale (LARS-i)</td>
<td>Specific</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>N</td>
<td>Adequate</td>
</tr>
<tr>
<td>Hseih et al. (2012)</td>
<td>Clinical Apathy Evaluation Scale (AES)</td>
<td>Specific</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>U</td>
<td>N</td>
<td>Good</td>
</tr>
<tr>
<td>Kaufer et al. (2000)</td>
<td>Brief Clinical Neuropsychiatric Inventory- Apathy (NPIa-Q)</td>
<td>Specific</td>
<td>U</td>
<td>U</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>Adequate</td>
</tr>
<tr>
<td>Leontjevas et al. (2012)</td>
<td>Abbreviated Apathy Evaluation Scale (AES-10)</td>
<td>General</td>
<td>U</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>Adequate</td>
</tr>
<tr>
<td>Lueken et al. (2007)</td>
<td>Abbreviated Apathy Evaluation Scale (AES-10)</td>
<td>Generic</td>
<td>N</td>
<td>U</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>Adequate</td>
</tr>
<tr>
<td>Malakouti et al. (2012)</td>
<td>Farsi Neuropsychiatric Inventory- Apathy (F-NPIa)</td>
<td>Specific</td>
<td>N</td>
<td>Y</td>
<td>N</td>
<td>U</td>
<td>U</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>Adequate</td>
</tr>
<tr>
<td>Pedersen et al. (2012)</td>
<td>Starkstein Apathy Scale (SAS)</td>
<td>Specific</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>U</td>
<td>N</td>
<td>U</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>U</td>
<td>Poor</td>
</tr>
<tr>
<td>Politis et al. (2004)</td>
<td>Hellenic Neuropsychiatric Inventory- Apathy (H-NPIa)</td>
<td>Specific</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>Good</td>
</tr>
<tr>
<td>Robert et al. (2002)</td>
<td>Apathy Inventory (AI)</td>
<td>Generic</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>N</td>
<td></td>
<td>Adequate</td>
</tr>
<tr>
<td>Sockeel et al. (2006)</td>
<td>Lille Apathy Rating Scale (LARS)</td>
<td>Specific</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>Y</td>
<td>U</td>
<td>Good</td>
</tr>
<tr>
<td>Starkstein et al. (1992)</td>
<td>Apathy Scale (SAS)</td>
<td>Specific</td>
<td>Y</td>
<td>N</td>
<td>U</td>
<td>N</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>U</td>
<td>Poor</td>
</tr>
<tr>
<td>Strauss et al. (2002)</td>
<td>Dementia Apathy Interview and Rating (DAIR)</td>
<td>Specific</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>Excellent</td>
</tr>
<tr>
<td>Zahodne et al. (2009)</td>
<td>Lille Apathy Rating Scale (LARS)</td>
<td>Specific</td>
<td>U</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>Adequate</td>
</tr>
</tbody>
</table>
## Generic neurodegenerative disease scales

### Table 3. Apathy scale psychometric properties in generic neurodegenerative disease studies

<table>
<thead>
<tr>
<th>Apathy Measure</th>
<th>Author (Year)</th>
<th>Type of measure</th>
<th>Patient Group</th>
<th>Patient N</th>
<th>Internal Consistency</th>
<th>Validity Statistic</th>
<th>Reliability Statistic</th>
<th>Quality Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>AES-S/I/C</td>
<td>Clarke et al. (2007)</td>
<td>Self Report, Informant Report and Clinical Patient Interview</td>
<td>55.2% AD, 14.3% mixed dementia (possible AD and possible lewy body dementia), 9.5% lewy body dementia, 5.7% VaD, 5.7% mixed dementia (AD and VaD), 4.8% FTD, 4.8% other dementia</td>
<td>121 (diagnosis ascertained for 105 participants)</td>
<td>Crohnbach's alpha (of factors derived from FA): AES-S = 0.884 Other factor = 0.410</td>
<td>Convergent (NPIa): Frequency x Severity: AES-S r = 0.22 (p&lt;0.05) AES-I r = 0.49 (p&lt;0.01) AES-C r = 0.27 (p&lt;0.01)</td>
<td>-</td>
<td>Adequate</td>
</tr>
<tr>
<td>NPIa</td>
<td>Cummings et al. (1994)</td>
<td>Informant Interview</td>
<td>AD, Probable VaD and other dementias</td>
<td>Concurrent validity study = 40</td>
<td>Cronbach’s alpha (Frequency and severity):</td>
<td>Content (Delphi Panel: 10 national and international experts in geriatric psychiatry, behavioural neurology)</td>
<td>Between rater agreement (2 trained raters, from a pool of 4 behavioural)</td>
<td>Adequate</td>
</tr>
</tbody>
</table>
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<table>
<thead>
<tr>
<th>AES-10</th>
<th>Leontjevas et al. (2012)</th>
<th>Informant Interview</th>
<th>Nursing Home Residents:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>42% no dementia, 17% AD, 11% VaD, 8% mixed dementia, 4% other dementia (Korsakov syndrome, Lewy Body dementia, dementia with PD, FTD and HD), 18% non specified dementia</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>100</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Cronbach’s -alpha = 0.95, standardized alpha = 0.95</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Congruent (NPIa Frequency x Severity): Spearman’s rho = 0.62 (p&lt;0.001)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Discriminant (CSDD): Spearman’s rho = 0.27 (p&lt;0.05)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>AES-10</th>
<th>Lueken et al. (2007)</th>
<th>Informant Report</th>
<th>Nursing Home Residents:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>7% no dementia, 69% AD, 7% VaD, 4% mixed dementia, 1% FTD,</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>356 (Subsample A = 178, Subsample B = 178)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Subsample A+B Cronbach’s alpha = 0.92</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Subsample A+B Concurrent (AES-I): r = 0.97</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Subsample A Concurrent (NPIa): r = 0.61</td>
</tr>
</tbody>
</table>

Inter-rater reliability study = 45
Test-retest reliability study = 20
Within range of 0.87 to 0.88
and neuropsychology rating each subdomain between 1 and 4; well assessed and poorly assessed, respectively:
NPIa screening question =1.3
NPIa subquestions = 1.4

neurology fellows, a geriatric psychiatry fellow, premedical student and behavioural neurologist:
Frequency- 97.9%
Severity- 89.4%

Test-retest (“within 3 weeks”)
Frequency r = 0.74 (p<0.001)
Severity r = 0.68 (p<0.01)

Adequate

AES-10
Leontjevas et al. (2012)
Informant Interview
Nursing Home Residents:
42% no dementia, 17% AD, 11% VaD, 8% mixed dementia, 4% other dementia (Korsakov syndrome, Lewy Body dementia, dementia with PD, FTD and HD), 18% non specified dementia
100
Cronbach’s -alpha = 0.95, standardized alpha = 0.95
Congruent (NPIa Frequency x Severity): Spearman’s rho = 0.62 (p<0.001)
Discriminant (CSDD): Spearman’s rho = 0.27 (p<0.05)

AES-10
Lueken et al. (2007)
Informant Report
Nursing Home Residents:
7% no dementia, 69% AD, 7% VaD, 4% mixed dementia, 1% FTD,
356 (Subsample A = 178, Subsample B = 178)
Subsample A+B Cronbach’s alpha = 0.92
Subsample A+B Concurrent (AES-I): r = 0.97
Subsample A Concurrent (NPIa): r = 0.61

Adequate
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1% dementia with PD, 3% other nonclassified dementia, 8% MCI

Discriminant (NPId): $r = 0.07$

(GDS): $r = -0.51$

Subsample B
Concurrent (NPia): $r = 0.62$

Discriminant (NPId): $r = 0.09$

(GDS): $r = -0.58$

AI Robert et al. (2002)
Informant Interview AND Patient Interview
AD, PD, MCI
115 Informant Interview
Cronbach’s alpha = 0.92

Concurrent (NPia Frequency x Severity):

Informant interview
Emotional Blunting $r = 0.01$ (NS)
Lack of Initiative $r = 0.23$ (p<0.01)
Lack of Interest $r = 0.63$ (p<0.001)
Global score $r =$ not recorded

Patient Interview
Emotional Blunting $r = 0.04$ (NS)
Lack of Initiative $r = 0.01$ (NS)
Lack of Interest $r = 0.26$ (NS)

Between rater agreement (26 psychiatrists, psychologists and medical students rating one videotaped response):
Kappa coefficient for item scores and global score = 0.99

Test-retest (“same day”):
Kappa coefficient emotional blunting = 0.99
Kappa coefficient lack of initiative = 0.97

Adequate
## Apathy scales in neurodegenerative conditions: a review

### Global score $r = \text{not recorded}$

### Discriminant (NPId Frequency x Severity):

- **Informant interview**
  - Emotional Blunting $r =$ not recorded
  - Lack of Initiative $r = 0.32$ (p<0.05)
  - Lack of Interest $r = 0.40$ (p<0.001)
  - Global score $r = 0.37$ (p<0.01)

- **Patient Interview**
  - Emotional Blunting $r =$ not recorded
  - Lack of Initiative $r = 0.37$ (p<0.01)
  - Lack of Interest $r = 0.31$ (p<0.05)
  - Global score $r = 0.42$ (p<0.001)

### Kappa coefficient

- Lack of interest = 0.99
- Global score = 0.99

---

**NS** = Not Significant; **NPId** = NPI depression/dysphoria; **NPlanx** = NPI anxiety; **CSDD** = Cornell Scale for Depression in Dementia
Table 3. shows the descriptors and quality of apathy scale validation studies in generic neurodegenerative disease patient samples. All five studies were of Adequate methodological quality.

The Apathy Evaluation Scale (AES) was originally developed and validated in English by Marin et al. (1991). It is made up of 18 items that assess emotional, behavioral and cognitive aspects of apathy through a 4-point Likert scale. The responses are coded to give one apathy score. The AES is available in Self report (S), Informant report (I) or Clinical patient interview (C) forms. The validation study by Clarke et al. (2007) looks at all three forms of the AES in a generic neurodegenerative dementia sample group. Convergent validity was measured through correlations with the NPIa, all of which were positive ranging from little or no correlation for the AES-S and AES-C to a low correlation with the AES-I. Discriminant validity for AES-S, AES-I and AES-C when compared to the NPId showed little or no positive correlation and with the NPIanx, ranging from low positive correlation for the AES-S and C to a low correlation for the AES-I. This study does not provide internal consistency values for each form of the AES. However, it has subdivided each available form of the AES into two factors each, resulting in numerous internal consistency values. The two factors for the AES-I and AES-C were labelled Apathy and Interest. For the AES-I the Interest factor showed good internal consistency and the Apathy Factor showed excellent internal consistency. The AES-C showed a good internal consistency for the Apathy Factor and an excellent internal consistency for the Interest factor. The AES-C showed a differing factor structure labelled Apathy and Other, with internal consistencies that are good and unacceptable, respectively.
Lueken et al. (2007) developed the abbreviated Apathy Evaluation Scale (AES-10) in German as a shortened, 10-item version of the AES-I. The items were developed and, as stated in the study, translated in close cooperation with Dr Robert Marin, the original author of the AES. It is scored almost identically as the original AES, on a 4-point Likert scale. It was validated in a sample of Nursing home residents, of various neurodegenerative diseases, some of who did not have dementia. The internal consistency reliability was excellent. A high positive correlation between the AES-10 and NPIa in both subsample A and B showed good concurrent validity. This was further supported by a very high, positive correlation with the AES-C. The discriminant validity was show to be good through little or no positive correlation between the AES-10 and the NPId in both subsamples. Furthermore, a moderate, negative correlation was observed between the AES-10 and the GDS.

A more recent study also looked at the Dutch version of the AES-10 in Nursing home residents, almost half of which were non demented (Leontjevas et al., 2012). As with the previous study, the internal consistency was shown to be excellent. For congruent validity, the AES-10 and NPIa showed a moderate positive correlation and when compared to a depression measure (CSDD), there was a little or no positive correlation, showing good discriminant validity.

The Neuropsychiatric Inventory (NPI; Cummings et al., 1994) is a 12 domain Informant interview that assesses behavioral disturbances in various diseases. Originally developed in English, it measures apathy via the NPI apathy/indifference (NPIa) domain firstly with a screening question to determine the presence of these behaviors (Yes/No/NA answers). This is followed by eight subquestions to determine
the characteristics of apathy (Yes/No answers). The frequency of the NPIa domain is rated on a 4-point Likert scale along with a 3-point Likert scale for severity. Frequency and severity are then multiplied to give a final NPIa domain score. The internal consistency was difficult to determine specifically for the NPIa due to lack of detailed reporting in the Cummings et al. (1994) study. However, based on the range of values provided in the article, the internal consistency of the NPIa was good. A Delphi panel determined content validity by rating whether screening question and sub-questions captured the essential elements of behavior. For the NPIa, they rated both as close to being ‘assessed well’. Inter-rater agreement and test-retest reliability for frequency was also found to be high with a further moderate test retest for severity.

The Apathy Inventory (AI; Robert et al., 2002) was developed and validated in French to assess global and subdomain apathy (emotional blunting, lack of initiative and lack of interest). It is a three item scale (one item for each domain) and is has been validated as a Self report and Informant interview. The Self report AI firstly has the patient assess their own behavior for each item (Yes/No) and then asks them to bisect a line in relation to the severity of their behavior that spans from mild to severe on a covert 12 point scale only know to the scorer. These three item severities are then added to give a composite apathy score. The Informant report is similar to the NPIa, recording whether the behaviors are displayed (Yes/No), frequency of behavior on a 4-point Likert scale along and severity of behavior on a 3-point Likert scale. Again, frequency and severity are multiplied to give a numeric apathy level for each subdomain, which are then added to give a composite apathy level. The internal consistency for the informant interview was excellent. The NPIa
generally held little or no to low positive correlations with AI subdomains, with only positive, moderate correlation with the informant Lack of Interest subdomain. There were no record of the correlation between the NPIa and the AI global scores. Discriminant validity determined by the relationship the NPId and the AI (global and subdomains), which showed low to moderate positive correlations. The between rater and test retest reliability was almost perfect for the global and subdomain scores.
Specific neurodegenerative disease scales

Table 4. Apathy scales psychometric properties in specific neurodegenerative disease patient populations

<table>
<thead>
<tr>
<th>Apathy Measure</th>
<th>Study</th>
<th>Type of measure</th>
<th>Patient Group</th>
<th>Patient N</th>
<th>Internal Consistency</th>
<th>Validity Statistic</th>
<th>Reliability Statistic</th>
<th>Quality Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPIa-C</td>
<td>de Medeiros et al. (2010)</td>
<td>Informant Interview</td>
<td>Mild to severe Probable AD</td>
<td>128</td>
<td>-</td>
<td>Convergent validity (AES-I): Frequency x Severity Pearson's r = 0.22</td>
<td>Inter-rater (ICC; 2 independently trained raters): NPIa-C subquestions range from 0.74 to 0.89</td>
<td>Adequate</td>
</tr>
<tr>
<td>LARS-i</td>
<td>Dujardin et al. (2008)</td>
<td>Informant Interview</td>
<td>Probable PD</td>
<td>60</td>
<td>Cronbach's between-item -alpha = 0.872 -standardized alpha = 0.877</td>
<td>Concurrent (AES-I): Global score r = 0.850 (p&lt;0.001) Concurrent (AES-C): Global score r = 0.811 (p&lt;0.001) Concurrent (LARS): Global score r = 0.814 (p&lt;0.001)</td>
<td>Inter-rater (ICC, 2 clinician interviewers): Global score r = 0.966 (p&lt;0.001)</td>
<td>Adequate</td>
</tr>
<tr>
<td>AES-C</td>
<td>Hseih et al. (2012)</td>
<td>Informant Interview</td>
<td>Mild to moderate AD</td>
<td>144</td>
<td>Cronbach's alpha = 0.85</td>
<td>Convergent (NPIa Frequency x Severity): Pearson's r = 0.61 (p&lt;0.001) Discriminant (NPId and NPIe, both Frequency x Severity): NPId Pearson's r = 0.16 (NS) NPIe Pearson's r = -0.46 (p&lt;0.001) Convergent (NPIa): Spearman's rho = 0.85 (p&lt;0.0001)</td>
<td>Test-retest (&quot;3 day period&quot;): Pearson's r = 0.89</td>
<td>Good</td>
</tr>
<tr>
<td>NPIa-Q</td>
<td>Kaufer et al. (2000)</td>
<td>Informant Report</td>
<td>Possible and probable AD</td>
<td>60</td>
<td>-</td>
<td>Convergent (NPIa): Spearman's rho = 0.85 (p&lt;0.0001)</td>
<td>Test-retest (&quot;same day&quot;): Pearson's r = 0.80 (p&lt;0.0001)</td>
<td>Adequate</td>
</tr>
</tbody>
</table>
| Scale        | Reference                                    | Domain          | Sample size | Cronbach’s alpha | Concurrent (PANSS): Frequency x Severity Pearson’s r | Inter-rater (ICC, 2 trained psychiatrist raters): 
<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>F-NPIa</td>
<td>Malakouti et al. (2012)</td>
<td>Informant Interview</td>
<td>Dementia 100</td>
<td>0.82</td>
<td>0.6-0.7 (p&lt;0.001)</td>
<td>Frequency Pearson’s r = 0.85 (p&lt;0.001) Severity Pearson’s r = 0.83 (p&lt;0.001)</td>
</tr>
<tr>
<td>SAS</td>
<td>Pedersen et al. (2012)</td>
<td>Self Report PD</td>
<td>194</td>
<td>0.69</td>
<td>Discriminant (MADRS): Spearman’s rho = 0.25 (p&lt;0.01)</td>
<td>-</td>
</tr>
<tr>
<td>H-NPIa</td>
<td>Politis et al. (2004)</td>
<td>Informant Interview</td>
<td>AD 29</td>
<td>Cronbach’s alpha (Frequency x Severity): Within range of 0.69 to 0.72</td>
<td>Concurrent (BPRS): Frequency x Severity Pearson’s r = 0.48 (p&lt;0.001)</td>
<td>-</td>
</tr>
<tr>
<td>LARS</td>
<td>Sockeel et al. (2006)</td>
<td>Patient Interview</td>
<td>Probable PD 159</td>
<td>Cronbach’s standardized alpha</td>
<td>Concurrent (AES): Global score r = 0.87</td>
<td>Inter-rater (ICC, 2 clinician raters): Global score r = 0.95</td>
</tr>
</tbody>
</table>
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<table>
<thead>
<tr>
<th>Scale</th>
<th>Reference</th>
<th>Type</th>
<th>Condition</th>
<th>Cronbach’s Alpha</th>
<th>Convergent</th>
<th>Discriminant</th>
<th>Inter-method</th>
<th>Test-retest</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SAS</strong></td>
<td>Starkstein et al. (1992)</td>
<td>Self Report</td>
<td>Idiopathic PD</td>
<td>0.76</td>
<td>Neurologist blind rating: Rated 6 apathetic and 6 apathetic with apathetic patients having significantly higher SAS score (apathetic SAS mean=14.8, SD=5.7; non-apathetic SAS mean=5.5, SD=2.2; t(10)=3.70, p&lt;0.001)</td>
<td></td>
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<tr>
<td><strong>DAIR</strong></td>
<td>Strauss et al. (2002)</td>
<td>Informant Interview</td>
<td>Possible and probable AD</td>
<td>0.89</td>
<td>Clinical ratings; Nurse, Physician and Neuropsychologist technician): r = 0.40, 0.31 and 0.46, respectively</td>
<td></td>
<td></td>
<td>Inter-rater (2 trained raters, one interviews one rates audiotape of interview): 100% agreement</td>
</tr>
<tr>
<td><strong>LARS</strong></td>
<td>Zahodne et al. (2009)</td>
<td>Patient Interview</td>
<td>Probable idiopathic PD</td>
<td>0.82</td>
<td>Convergent (SAS): Global score r = 0.62 (p&lt;0.001) Intellectual curiosity subscale r = 0.61 (p&lt;0.001) Action Initiation subscale r = 0.42 (p&lt;0.001) Emotion subscale r = 0.33 (p&lt;0.01) Self awareness subscale r = 0.229 (NS)</td>
<td></td>
<td></td>
<td>Test-retest (“an average of 56 days”): r = 0.85 (p&lt;0.001)</td>
</tr>
</tbody>
</table>

- Between item = 0.80
- Between subscale = 0.74

Intelectual curiosity subscale r = 0.84
Action Initiation subscale r = 0.65
Emotion subscale r = 0.44
Self awareness subscale r = 0.15

Convergent (Neurologist blind rating): Rated 6 apathetic and 6 apathetic with apathetic patients having significantly higher SAS score (apathetic SAS mean=14.8, SD=5.7; non-apathetic SAS mean=5.5, SD=2.2; t(10)=3.70, p<0.001)

Inter-method agreement (ICC; SAS): Global score = 0.75

Divergent (BDI-II):
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Global score $r = 0.45\ (p<0.001)$

Inter-method agreement (ICC; BDI-II):
Global score $r = 0.62\ (p<0.001)$

NS= Not Significant; ICC= Intra-class correlation NPlde= NPI depression/dysphoria; NPlanx= NPI anxiety; NPlle= NPI elation/euphoria; PANSS= Positive and Negative Symptoms Scale; MADRS= Montgomery-Aasberg Depression Rating Scale; BPRS= Brief Psychiatric Rating Scale; BRSD-d= Behavior Rating Scale for Dementia- Depression subscale; BDI-II= Beck Depression Inventory II
Table 4 shows the different apathy scale validation studies in specific neurodegenerative disease patient samples, with descriptors and quality rating for each. The 11 studies ranged from Poor to Excellent quality.

The NPI has been translated into Greek (Hellenic; Politis et al., 2004) and a Persian language (Farsi; Malakouti et al., 2012) and validated in AD and generic dementia populations. Both are administered and scored in the same way as the original NPIa (Cummings et al., 1994). The Hellenic NPIa (H-NPIa) was translated by two bilingual psychiatrists from English to Greek and then independently translated back to English by one other psychiatrist. The quality of the study was classed as Good. The H-NPIa frequency x severity internal consistency fell within a range of questionable and acceptable with no further reliability reported. The concurrent validity between the apathy factor derived from the BPRS and the H-NPIa showed a low positive correlation.

The Farsi NPIa (F-NPIa) was also translated by two bilingual psychiatrists from English to Farsi and then back-translated to English independently by two other bilingual psychiatrists. It was then validated in an unspecified dementia patient sample. The methodological quality of the study was classed as Adequate. However, the internal consistency was reported of the F-NPIa was good. The concurrent was determined against the PANSS, which was found to be of a high correlation. The reliability of the scale was good, with an excellent inter-rater intra-class correlation coefficient (ICC) and the test retest correlation coefficient was good.
In addition to the NPI translations above, several revised versions have been developed. The Brief Clinical NPI (NPI-Q: Kaufer et al., 2000), developed in English, is scored identically to the NPI but differs in administration method in that it is an informant report measure. The informants are asked to fill in the NPI-Q about the patient. The original validation study by Kaufer et al. (2000) recruited AD patients and was found to be of Adequate quality. While there was no report of the internal consistency for the NPI-Q apathy (NPIa-Q), the test-retest reliability correlation coefficient and convergent validity against the NPIa were both high positive correlations.

The NPI Clinical rating scale (NPI-C; de Medeiros et al., 2010) is a revised version of the NPI with expanded domains and questions. It recognises the importance of the domain subquestions and rates frequency and severity through an item-by-item basis. It was validated in mild to severe probable AD patients by de Medeiros et al. (2010). This was a large-scale, cross-sectional validation study with 8 participating countries (Argentina, Brazil, Canada, France, Greece, Hungary, Italy and America) and the NPI-C was translated to 7 languages (English, French, Greek, Hungarian, Italian, Portuguese and Spanish). The quality of this study was Adequate. It does not report internal consistency NPI-C apathy (NPIa-C) and the convergent validity against the AES-I showed little or no positive correlation. The inter-rater ICC for each of the subquestions of the NPIa-C ranged from good to excellent.

The Starkstein Apathy Scale (SAS) was developed in English to assess apathy and validated in idiopathic PD patients (Starkstein et al., 1992). It is a condensed version of the AES, with only 14 items, scored by same 4-point Likert scale. It
defines itself as less comprehensive than the AES as to be less demanding for PD patients to complete. This study was of Poor quality with only an acceptable internal consistency for the SAS. The convergent validity was determined through a blind neurologist rating participants as apathetic or non-apathetic and then examining if their SAS scores are significantly different. They were found to be different, based on a sample size of 12 (six rated as apathetic, six rated as non-apathetic by the neurologist). The inter-rater reliability correlation coefficient was high and the test-retest reliability correlation was very high. In a later study, the SAS was back-translated in to Norwegian, the translation critically appraised by four of the authors and used in a Norwegian PD patient sample study (Pedersen et al., 2012). This study was also of Poor quality. This study reported a questionable internal consistency for the SAS with good discriminant validity due to little or no positive correlation with the MADRS.

The Lille Apathy Rating Scale was originally developed in French and validated in patients with probable PD by Sockeel et al. (2006). It is a 33-item scale scored by a mixture of simple (Yes/No/NA) and Likert-type responses. It is composed of factor analytically derived subscales (Intellectual curiosity, Action initiation, Emotional and Self awareness), which are also summed to provide a global apathy score. The original validation study (Sockeel et al., 2006) was of Good quality. It showed an acceptable (between subscale) to good (between item) internal consistency. Concurrent validity between the AES and LARS global score was high positive correlation, with LARS subscales being positive and ranging from little or no to high correlations. Inter-rater ICC and test retest reliability was found to be excellent. A study by Zahodne et al. (2009) of a slightly less Adequate quality further
validated LARS in English in probable idiopathic PD patients. It was showed to have a good internal consistency. For convergent validity, the SAS and the LARS global score correlated positively and moderately, with the subscales positively correlating from little to none to moderately. Additionally, the inter-method agreement (ICC) for the SAS and LARS global score was excellent. Divergent validity between the BDI-II and LARS global score was moderate and the ICC between the two was fair.

Similarly to the LARS, the informant version of the LARS (LARS-i; Dujardin et al., 2008) was also developed in French and for validation in a PD population. Some items have been reworded to be suitable as a question for a patient’s informant. The administration and scoring methods are the same as the LARS. This particular study was of Adequate quality. The internal consistency for the LARS-i was shown to be good. The concurrent validity was good due to high positive correlations with the AES-I, AES-C and LARS. Inter-rater ICC and test retest reliability was excellent.

The AES-C was further validated in mild to moderate severity AD patient sample (Hseih et al., 2012). It was back-translated to Taiwanese by two bilingual individuals. This was done until an agreement on the translation was reached between the two. In terms of quality, this study was classed as Good. The internal consistency for the AES-C was found to be good, with a moderate positive convergent validity against the NPIa. Discriminant validity against the NPId showed little to no correlation while against the NPIe it showed a low negative correlation. Test retest reliability correlation coefficient for the AES-C was good.
The Dementia Apathy Interview and Rating (DAIR) is an informant based interview, developed and validated in English to assess apathy in probable and possible cases on AD (Stuss et al., 2002). The assessment is a 16-question interview of the primary caregiver to assess the patient’s initiation behavior, interest and engagement in the environment through a 4-point Likert scale. This was the only study with Excellent methodological quality. The internal consistency of the DAIR was good with a generally low positive correlation in relation to clinician’s ratings, indicative of average convergent validity. It had little or no positive correlation with the BDRS, showing good discriminant validity against depression. It had a prefect inter-rater agreement with a high positive test-retest reliability correlation coefficient.

Discussion

This review systematically examines and provides an overview of the quality and psychometric properties of apathy scales, which have been used within neurodegenerative disease populations. Generally, most of the validation studies in this review were found to be of Adequate quality. Studies did not reach higher quality scores due to under described selection criteria, poor reporting of withdrawals from studies and a lacking of a good reference standard for diagnosing apathy. Nevertheless, their merits are that both the reference standards and the index tests for apathy were described in sufficient detail and, mostly, were independent of each other.

Generic neurodegenerative scales

The methodological quality of all the validation studies for the neurodegenerative generic scales was Adequate. This could have been improved with
better, defined selection criteria and comprehensive sampling to form more generic neurodegenerative patient groups. Therefore, it is important to examine the psychometric properties of each as to determine their practicality.

The widely used NPI, developed and validated in a generic dementia sample by Cummings et al. (1994), is often used as a reference standard for other apathy measures. In the original validation study by Cummings et al. (1994), the design of the NPI was done using a Delphi panel of experts as to also determine content validity. They reported apathy as ‘well assessed’ but did not examine any other statistical validity criteria for the NPIa. Specific NPIa internal consistency was not reported but it was deduced as good, with a high between-rater agreement and good test-retest reliability. Most of the NPI subdomains were concurrently validated against existing instruments, whereas the NPIa domain was not. This lack of concurrent validity for the NPIa and a methodological rating of Adequate for this study create uncertainty of the robustness of this subscale’s ability to assess apathy. It is observed that many studies in this review have used the NPIa as a reference standard to validate their own methods of apathy assessment. As a consequence, perhaps these and any future validation studies using the NPIa as a reference standard should be considered and interpreted with caution.

Similarly to the NPI, the AES is also considered as a validation reference standard. The original Marin et al. (1991) AES development and validation study was excluded in the earlier stage of this review due to a small sample size of the AD patients (N = 21 our of 123/ less than 20%) and included a very mixed patient sample of AD, stroke (left and right hemisphere), major depression and healthy controls.
Additionally, there was no specific AD reliability or validity information reported for the AES. However, a later study by Clarke et al. (2007) validated all forms of the AES against a generic dementia sample. The Clarke et al. (2007) study does not report reliability statistics and only factor analytically derived internal consistencies, making it difficult to interpret the scales as a whole. The convergent and discriminant validity was variable for all forms of the AES, resulting in unclear psychometric properties and, once again, therefore caution should be taken in considering its application as a reference standard for a neurodegenerative group.

Two studies validating the AES-10 (Lueken et al., 2007, Leontjevans et al., 2012), which is a shortened form of the AES, showed excellent internal consistency reliability and improved validity. Similar to the Clarke et al. study, the AES-10 validation studies documented no other reliability statistics. It should, however, be noted that both studies recruited Nursing Home residents some of whom did not have a diagnosis of a neurodegenerative disease. The Lueken et al. (2007) was more representative of a neurodegenerative generic dementia population. However, the Leontjevans et al. (2012) study was not very representative of a neurodegenerative population because almost half of the participants had no signs of dementia. So while the AES-10 seems psychometrically strong, it could still benefit from further validation in a generic neurodegenerative disease population.

Robert et al.’s (2002) AI validation study utilised a neurodegenerative generic patient sample containing AD, PD and mild cognitive impairment patients. While the AI was generally reliable, the inter-rater agreement, internal consistency, test-retest reliability being excellent, the validity was unclear. The concurrent validity was
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underreported for the global score and showed an irregular relationship for the subscale scores. The association between the AI global and subscale scores with depression were stronger than that of the apathy measure. This stronger relationship with depression could indicate that the AI is not measuring the intended apathy, but rather some component associated with both apathy and depression, resulting in a lack of scale validity.

Based on the results of this review, it is difficult to establish which scale is best for assessing apathy in a generic neurodegenerative disease population. There are various methodological flaws with the validation studies that affect the validity and reliability of the each scale. The main issue being that a generic neurodegenerative sample should involve all forms of neurodegenerative disease, not just various dementias (Cummings et al., 1994, Clarke et al., 2007) or a mixture healthy individuals and neurodegenerative patients (Lueken et al., 2007, Leontjевans et al., 2012). Psychometrically, the AES-10 seems to be the most valid, but has not been applied in a purely neurodegenerative population.

**Specific neurodegenerative scales**

With regard to neurodegenerative specific apathy scales, the DAIR (Strauss et al., 2002) validation study has Excellent methodological quality for AD patients, due to lack of bias, a superior quality of methodological reporting and high generalizability of the results. This was closely followed by Good quality validation studies of the AES-C (Hseih et al., 2012) and the H-NPIa (Politis et al., 2004). Both had a representative sample and high generalizability of the results. However, the H-NPIa was validated against a combination of BPRS subscales that were labelled as an
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Apathy factor, which is not a direct or established measure of apathy but rather a new construct. Additionally, when compared psychometrically, the H-NPIa reliability and validity was inferior to both the DAIR and the AES-C, partly due to insufficient reliability criteria reporting. When comparing the DAIR and the AES-C, both are well constructed, informant interviews completed by trained assessor with matched psychometric properties. Both validation studies note good internal consistency reliability but vary in the reporting of reliability and validity. The AES-C reports more detailed convergent and discriminant validity that are marginally superior to the DAIR. Conversely, the DAIR has better inter-rater agreement and test-retest reliability in comparison to the AES-C. Based on the psychometric properties of both these scales and the quality of their validation studies, either of these methods would be recommended for assessing apathy in AD patient populations.

In relation to PD patients, both the SAS validation studies (Starkstein et al., 1992, Pedersen et al., 2012) scored Poor on quality mostly due to the lack of methodological reporting and selection bias and also showed unfavourable psychometric validity and reliability. Only the Sockeel et al. (2006) LARS validation study was methodologically classed as Good quality due to minimised bias, detailed methodology reporting and generalizability of results. The internal consistency reliability was found to be acceptable which was supported by very good validity and reliability properties of the LARS global score. The high convergent validity and excellent reliability make this a good measure for apathy in PD. However, it does not examine discriminant or divergent validity against depression, which has been shown to occur with apathy in PD (Lieberman, 2006). A later validation study showed average discriminant validity against depression (Zahodne et al., 2009).
In ALS, apathy is continually assessed using the Frontal Systems Behavior Scale (Grossman et al., 2007; Witgert et al., 2010) despite this scale not being validated in this neurodegenerative disease population. Additionally, research in HD has been conducted using scales that assess uncorroborated apathy that is a derivation of factor analysis (Thompson et al., 2012) or use scales that have not been validated in an HD sample (Hamilton et al., 2003, van Duijn et al., 2010). Therefore, no well-validated scales to assess apathy within these two neurodegenerative disorders have been found.

Strengths and limitations

This review has been undertaken using a systematic methodology (PRISMA), ensuring that eligibility criteria, inclusion and exclusion criteria were carefully outlined and implemented. Secondly, all studies were included that met the criteria, which could be seen to reduce publication bias. Thirdly, through rating the validation studies for the scales on methodological quality using the QUADAS, allowed us to determine the quality of the validation procedure for the apathy scale in question. Finally, interpreting the psychometric statistics for reliability and validity of the scales (adapted from Hermans et al., 2011) allowed for a more standardized assessment. This resulted in more objectively driven recommendations for the most robust scale to be used in research or clinical practice to be made.

A limitation for this study could be that we chose to exclude studies that reported just validity or reliability of apathy scales. However, a diagnostic instrument should be both reliable and valid at detecting a symptom (Kimberlin and Winetrstein,
and both of these need to be demonstrated in the same sample (i.e. there is little point in having a scale that is reliable but not valid in one sample, and valid but not reliable in another). Therefore, by including validation studies that examine both of these criteria, we attempted to avoid partially validated scales that might not be suitable for assessment. Additionally, a systematic review using the PRISMA method requires “a clearly formulated question that uses systematic and explicit methods…” of identification, selection and critical appraisal of “relevant research” (Moher et al., 2009). So by clearly defining those studies reporting complete results (i.e. reliability and validity statistics of a scale) improve the quality of this systematic review, making it possible for more credible, scale-utility recommendations to be made.

Finally, it should be noted that apathy as a lack of motivation may manifest somewhat differently depending on the cultural context. Hence, caution is required when considering applying a scale validated in one country in another, especially if cultural norms are very dissimilar.

**Conclusion**

In conclusion, there are several recommended instruments to assess apathy for some specific neurodegenerative disease populations. The DAIR is a well-validated instrument in AD patients, shown to have very good psychometric characteristics, which is further supported by Excellent methodological quality of the validation study (Strauss et al., 2002). In relation to PD patient, the LARS validation study (Sockeel et al., 2006) showed that the global scoring of apathy was psychometrically strong, with further research needed to determine its subscalar structure and relationship with
depression. Finally, scales should be developed and validated in diseases such as HD and ALS, which make apathy difficult to assess due to marked motor symptoms.

This review did not find any methodologically strong studies that validated a psychometrically robust, generic neurodegenerative disease apathy scale. This was partly due to the unrepresentative samples, lack of standardizing selection criteria in the validation studies for generic neurodegenerative disease. Many of the studies validated scales in samples that were biased to dementia patients, with no studies including ALS patients. It may be difficult to design or validate such a generic scale due to practical difficulties of forming a broad and mixed neurodegenerative disease patient sample group. However, further well-constructed validation studies or newly developed scales are needed to explore if it is possible to assess apathy in generic neurodegenerative disease.

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Conflict of interest

None.

Description of authors’ roles

R.R. and J.S. devised the review. R.R. selected and condensed the material and wrote the paper with support of J.S. C.H. was the second reviewer for the paper, checked
material and co-rated papers. J.S. was the third reviewer and supervised R.R. S.A. supervised R.R. and provided intellectual input into the manuscript.

Acknowledgements
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Supplementary material
Supplementary Table S1. Guidelines for interpreting psychometric statistics (based on Hermans et al., 2011)
Supplementary Figure S1. PRISMA Checklist
Supplementary Figure S2. Stacked bar chart of methodological quality for all the apathy scale validation studies by QUADAS item

References
Neurodegenerative Disease Management, 1(5), 397-405.


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Supplementary Table S1. Guidelines for interpreting psychometric statistics (based on Hermans et al., 2011)

<table>
<thead>
<tr>
<th>Psychometric Statistic</th>
<th>Interpretation</th>
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<tr>
<td>Internal Consistency</td>
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<tr>
<td>&lt;0.50</td>
<td>Unacceptable</td>
</tr>
<tr>
<td>0.50-0.59</td>
<td>Poor</td>
</tr>
<tr>
<td>0.60-0.69</td>
<td>Questionable</td>
</tr>
<tr>
<td>0.70-0.79</td>
<td>Acceptable</td>
</tr>
<tr>
<td>0.80-0.89</td>
<td>Good</td>
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<tr>
<td>≥0.90</td>
<td>Excellent</td>
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<tr>
<td>Correlation Coefficients (Pearson’s product-moment and Spearman rank)</td>
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<tr>
<td>&lt;0.29</td>
<td>Little or no correlation</td>
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<tr>
<td>0.30-0.49</td>
<td>Low correlation</td>
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<td>0.70-0.89</td>
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<tr>
<td>≥0.90</td>
<td>Very high correlation</td>
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<tr>
<td>Intra-class Correlation Coefficient</td>
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<tr>
<td>&lt;0.40</td>
<td>Poor</td>
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<tr>
<td>0.40-0.59</td>
<td>Fair</td>
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<tr>
<td>0.60-0.74</td>
<td>Good</td>
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<tr>
<td>≥0.75</td>
<td>Excellent</td>
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<td>Kappa Statistic</td>
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<td>0.21-0.40</td>
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<td>0.41-0.60</td>
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<td>0.61-0.80</td>
<td>Substantial</td>
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<td>0.81-1.00</td>
<td>Almost prefect</td>
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## Supplementary Figure S1. PRISMA Checklist

<table>
<thead>
<tr>
<th>Section/topic</th>
<th>#</th>
<th>Checklist item</th>
<th>Reported on page #</th>
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<tr>
<td><strong>TITLE</strong></td>
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<tr>
<td>Title</td>
<td>1</td>
<td>Identify the report as a systematic review, meta-analysis, or both.</td>
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<tr>
<td><strong>ABSTRACT</strong></td>
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<tr>
<td>Structured summary</td>
<td>2</td>
<td>Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.</td>
<td>2</td>
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<tr>
<td><strong>INTRODUCTION</strong></td>
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<tr>
<td>Rationale</td>
<td>3</td>
<td>Describe the rationale for the review in the context of what is already known.</td>
<td>3 to 7</td>
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<tr>
<td>Objectives</td>
<td>4</td>
<td>Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).</td>
<td>7 and 8</td>
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<tr>
<td><strong>METHODS</strong></td>
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<tr>
<td>Protocol and registration</td>
<td>5</td>
<td>Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.</td>
<td>N/A</td>
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<tr>
<td>Eligibility criteria</td>
<td>6</td>
<td>Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.</td>
<td>8</td>
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<tr>
<td>Information sources</td>
<td>7</td>
<td>Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.</td>
<td>8</td>
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<tr>
<td>Search</td>
<td>8</td>
<td>Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.</td>
<td>8 and 9</td>
</tr>
<tr>
<td>Study selection</td>
<td>9</td>
<td>State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).</td>
<td>9 and 10</td>
</tr>
<tr>
<td>Data collection process</td>
<td>10</td>
<td>Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.</td>
<td>9 and 10</td>
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<tr>
<td>Data items</td>
<td>11</td>
<td>List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.</td>
<td>8 to 10</td>
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<tr>
<td>Risk of bias in individual studies</td>
<td>12</td>
<td>Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.</td>
<td>10 and 11</td>
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<tr>
<td>Summary</td>
<td>13</td>
<td>State the principal summary measures (e.g., risk ratio, difference in means).</td>
<td>10 and 11</td>
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<tr>
<td>measures</td>
<td>14</td>
<td>Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I²) for each meta-analysis.</td>
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<td>Risk of bias across studies</td>
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<td>Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).</td>
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<tr>
<td>Additional analyses</td>
<td>16</td>
<td>Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.</td>
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<tr>
<td>RESULTS</td>
<td>17</td>
<td>Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.</td>
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<td>18</td>
<td>For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.</td>
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<td>19</td>
<td>Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).</td>
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<td>20</td>
<td>For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.</td>
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<td>21</td>
<td>Present results of each meta-analysis done, including confidence intervals and measures of consistency.</td>
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<td>22</td>
<td>Present results of any assessment of risk of bias across studies (see Item 15).</td>
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<td>23</td>
<td>Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).</td>
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<tr>
<td>DISCUSSION</td>
<td>24</td>
<td>Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).</td>
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<td>25</td>
<td>Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias).</td>
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<td>26</td>
<td>Provide a general interpretation of the results in the context of other evidence, and implications for future research.</td>
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<tr>
<td>FUNDING</td>
<td>27</td>
<td>Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.</td>
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Supplementary Figure S2. Stacked bar chart of methodological quality for all the apathy scale validation studies by QUADAS item

Target condition = Apathy; Index test = Apathy assessment method; Reference standard = Apathy diagnosis method