Lower practice effects as a marker of cognitive performance and dementia risk: A literature review

Roos J. Jutten 1 | Evan Grandoit 2 | Nancy S. Foldi 3 | Sietske A. M. Sikkes 1 | Richard N. Jones 4 | Seo-Eun Choi 5 | Melissa L. Lamar 6 | Diana K. N. Louden 7 | Joanne Rich 7 | Douglas Tommet 4 | Paul K. Crane 5 | Laura A. Rabin 8

1 Alzheimer Center Amsterdam, Amsterdam Neuroscience, Amsterdam UMC, Vrije Universiteit, Amsterdam, the Netherlands
2 Northwestern University, Evanston, Illinois, USA
3 Queens College and The Graduate Center of The City University of New York, Queens, New York, USA
4 Warren Alpert Medical School of Brown University, Providence, Rhode Island, USA
5 School of Medicine, University of Washington, Seattle, Washington, USA
6 Rush Alzheimer’s Disease Center, Rush University Medical Center, Chicago, Illinois, USA
7 University Libraries, University of Washington, Seattle, Washington, USA
8 Brooklyn College and The Graduate Center of The City University of New York, Brooklyn, New York, USA

Correspondence
Roos J. Jutten, Alzheimer Center Amsterdam, Amsterdam Neuroscience, Amsterdam UMC, location VUmc, P.O. Box 7057, 1007 MB Amsterdam, the Netherlands.
Email: r.jutten@amsterdamumc.nl

Abstract
Background: Practice effects (PEs) are improvements in performance after repeated exposure to test materials, and typically viewed as a source of bias in repeated cognitive assessments. We aimed to determine whether characterizing PEs could also provide a useful marker of early cognitive decline.

Methods: We conducted a systematic review of the literature, searching PsycInfo (Ebsco) and PubMed databases for articles studying PEs in aging and dementia populations. Articles published between 1920 and 2019 were included.

Result: We identified 259 articles, of which 27 studied PEs as markers of cognitive performance. These studies consistently showed that smaller, less-robust PEs were associated with current diagnostic status and/or future cognitive decline. In addition, lower PEs were associated with Alzheimer’s disease risk factors and neurodegeneration biomarkers.

Conclusion: PEs provide a potentially useful marker of cognitive decline, and could prove valuable as part of a cost-effective strategy to select individuals who are at-risk for dementia for future interventions.

KEYWORDS
Alzheimer’s disease, cognition, learning effects, practice effects, retest effects

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1 | BACKGROUND

Practice effects (PEs) are expected improvements in cognitive performance seen on repeated exposure to test material in the absence of intervention.1 PEs, also referred to as retest or learning effects, are typically viewed as a source of bias or error when analyzing data from repeated cognitive assessments,2,3 particularly in the study of Alzheimer’s disease (AD) and other neurodegenerative disorders leading to dementia where cognitive decline is a key marker of clinical change.4,5 PEs can hinder our understanding of the disease course of AD and other neurodegenerative diseases, as well as the evaluation of interventions that aim to slow or halt cognitive decline.6 That is, by masking cognitive decline due to an underlying neurodegenerative process or by inflating cognitive gain in the absence of treatment-induced brain changes, PEs may lead to underestimating the severity of disease progression or overestimating the efficacy of treatment effects.7

The absence of PEs may also provide useful information in the context of AD and dementia. More specifically, one might expect attenuated PEs in disorders such as mild cognitive impairment (MCI) and AD, in which learning is compromised. This was, for example, supported by Duff et al.,8 who showed that lower PEs were predictive of cognitive decline 1 year later in individuals with MCI. Another study found that cognitively healthy older adults who later progressed to AD dementia had substantially lower PEs on episodic memory tasks compared to those who remained cognitively healthy.9 Together, these findings suggest that lower PEs may indicate a subtle cognitive impairment preceding overt reduction in cognitive performance, and may serve as an early marker to differentiate neurodegeneration from healthy cognitive aging. This would be of particular relevance in pre-dementia disease stages such as MCI or subjective cognitive decline, when objective cognitive decline is modest or not easily captured by traditional cognitive assessments.10,11 Therefore, the aim of the current review was to examine the role of PEs as a potential marker for cognitive decline in the study of cognitive aging.

Previous summaries of the PE literature in aging populations have largely focused on PEs as a source of bias. An example is the meta-analysis by Calamia et al.,12 which examined the magnitude of PEs on several widely-applied cognitive tests (both memory and non-memory tests), and investigated the influence of age, test-retest interval, use of alternate forms, and clinical diagnosis on those effects.12 Of interest, they found that clinical groups (i.e., patients with neurological or psychiatric conditions) showed lower PE on average compared to cognitively healthy adults. The authors concluded that PEs should be accounted for in cognitively healthy populations to accurately assess group-level changes. Moreover, this finding also suggests that lower PEs in patient groups may reflect a cognitive (i.e., learning) deficit, which could serve as a clinical marker of interest. The potential value of PEs as an indicator or marker of dementia risk is further supported by other studies, which suggest that PEs could serve as a proxy of specific fluid or imaging AD biomarkers,13 or could be used in combination with those biomarkers to identify individuals at greatest risk for clinical progression.14

HIGHLIGHTS

- We reviewed published research that studied practice effects (PEs) as markers of cognitive performance
- Lower PEs may associate with current cognitive status and predict future decline
- Lower PEs may associate with specific biological risk factors for Alzheimer’s disease

RESEARCH IN CONTEXT

1. Systematic review: The authors conducted a systematic review of the literature, searching PsycInfo and PubMed databases for articles studying practice effects (PEs) on cognitive testing in aging and dementia populations. Of 259 identified articles, 27 studied PEs as a clinically useful marker of cognitive performance in older adults.

2. Interpretation: We found accumulating evidence that lower PEs may represent an early indicator of cognitive decline, and that lower PEs associate with specific Alzheimer’s disease (AD) biological risk factors. The combination of quantifying PEs and assessing AD biomarkers may yield an optimal approach to estimate AD risk.

3. Future directions: Future research should focus on identifying high-risk individuals from a combination of cognitive and clinical features, AD biomarkers, and lower than expected PEs. This could yield a cost-effective strategy to enrich samples in clinical trials and provide a valuable cognitive marker for subtle pharmacological or treatment responses.

Although PEs have been investigated and addressed in many studies for over a century, we sought to consolidate and compare studies by conducting a literature review of published research relevant to aging and dementia. More specifically, we aimed to summarize research that investigated whether the presence or absence, or magnitude of PEs could (1) be an indicator of current diagnostic status (i.e., cognitively normal, MCI, or dementia); (2) predict future cognitive decline or progression to dementia; and/or (3) relate to AD risk factors or biomarkers of neurodegeneration.

2 | METHODS

2.1 | Search strategy

The authors obtained published empirical studies through a systematic search of the PsycInfo (Ebsco) and PubMed databases. Based on
consultation with research medical librarians (DKNL and JR), our search strategy used the following key search terms: practice effects, learning effects, retest effects, repeat testing, serial testing, serial assessment, longitudinal testing, neuropsychological testing, cognitive change, reliability, and early detection. A full text and abstract were required for inclusion; there were no restrictions for date of publication or language. The authors also manually examined the references of relevant studies to identify additional articles. Searches were conducted on December 14, 2017, and on April 18, 2019, December 2, 2019, and January 31, 2020, to include published manuscripts of interest.

### 2.2 Screening and review process

We selected articles using the following steps:

1. **Identification.** Two authors (EG and LAR) reviewed titles and abstracts to identify peer-reviewed full-text articles focusing on PEs in aging samples or important analytic/statistical issues about modeling PEs, which were selected for subsequent evaluation.

2. **Initial screening.** Research team members (NSF, PKC, LAR, MLL, RJJ, SAMS, and RNJ) evaluated full texts of articles identified in Step 1. One investigator reviewed each article and introduced it to the group for discussion to classify the article in the subsequent step.

3. **Eligibility.** Based on the full-text evaluations, articles were categorized as (A) PEs as a nuisance variable and possible solutions; (B) measuring and understanding the construct of PEs (eg, underlying mechanisms, moderators); and/or (C) PEs as a potential measure of cognitive change. Categories were not mutually exclusive, meaning that articles could be assigned to multiple categories.

4. **Inclusion in the current study.** The goal of the current article was to determine whether PEs could provide clinically useful information in the context of dementia or dementia risk, and we therefore further evaluated articles within Category C (ie, reported practice effects as a valuable measure of cognitive change). Each Category C article was reviewed by two independent investigators (from among authors NSF, PKC, LAR, MLL, RJJ, SAMS, RNJ, and EG) and assigned to at least one theme: (1) PE variability defines cases; (2) PE variability predicts outcomes; and (3) PEs associate with AD risk factors (Table 1). Articles could be assigned both primary and secondary themes. The authors also rated each article’s methodological rigor as low, medium, or high. If an article was identified as having more than one theme, it was assigned a secondary theme. If two investigators disagreed on the theme(s) or quality of an article, it was reviewed by a third investigator.

### 2.3 Data extraction

The following data were extracted from each included article: study population (ie, normal cognition, MCI, and/or dementia), mean age, retest interval, cognitive domains, cognitive task, biomarkers (if available), study design (ie, cross-sectional, longitudinal, or mixed), method of quantifying PEs, research findings, and conclusions. With regard to study design, for the purpose of the current article, the term “cross-sectional” was applied to studies for which the PEs (ie, second testing to define a retest effect) were measured concurrently with the outcome of interest (eg, current diagnostic status); by contrast, the term “longitudinal” was applied to studies in which retesting to identify PEs preceded the outcome being measured (eg, progression to dementia, change in cognitive test scores). All data were extracted by one author (EG) and verified by a second author (LAR). Descriptive analyses on study population, age, retest interval, cognitive domains, cognitive task, and methods of quantifying PEs were performed by synthesizing the data across all of the relevant articles and presented in one overview table. Subsequently, data were summarized by theme to investigate whether PEs were associated with (1) current diagnostic status; (2) future cognitive decline or change in diagnostic status; and (3) AD risk factors.

### 3 RESULTS

Figure 1 shows the results from the screening and review processes. The search identified 259 articles published between October 1920 and December 2019. We determined that 107 of these investigated either PEs on cognitive tests in aging samples or methods on modeling PEs, and underwent full-text evaluation by research team members. Of those, 27 met eligibility for Category C and were included in the current study. We assigned six of the 27 articles to primary Theme 1 (22%), 10 to primary Theme 2 (37%), and 11 to primary Theme 3 (41%). We assigned secondary themes for five of the articles (three to Theme 3 and two to Theme 1).

Table 2 displays the key study characteristics of the 27 articles considered in this article. Studies in these reports included from n = 25 to n = 1390 participants, with mean ages ranging from 53.4 to 83 years, with an overall average of 73.6 years. Studies evaluated several different diagnostic groups, retest intervals, cognitive domains and tasks, and used different methods to calculate PEs (Table 2). Most studies
**TABLE 2**  Overview of the n = 27 included articles studying practice effects as a marker of cognitive performance in aging and dementia populations

| Research article and reference number | Retest interval | Baseline groups | Cognitive domains | Cognitive tasks | Biomarkers | Cross-sectional/Longitudinal/Mixed | How PEs calculated/quantified | Mean age/ and total n |
|--------------------------------------|----------------|-----------------|-------------------|----------------|------------|-----------------------------------|-------------------------------|---------------------|
| Cooper et al.15                      | 1-4 days       | AD vs. Normal Cognition | Memory (episodic) | AMT            | N/A        | Longitudinal                      | Change score                | ~70 years/ n = 123 |
| Cooper et al.16                      | 1 wk           | AD vs. MCI vs. Normal Cognition | Memory (episodic) | AMT            | N/A        | Longitudinal                      | Change score                | ~69 years/ n = 69  |
| Darby et al.17                       | Same day       | MCI vs. Normal Cognition | Memory (episodic, working); Attention; Psycho-motor speed | Non-standardized measures | N/A        | Longitudinal                      | Change score                | n/a / n = 60       |
| Dodge et al.18                       | Every 18 mos (> 14 yrs) | Normal Cognition | Learning; Memory; Language; Psycho-motor speed; Executive function | MMSE; TMT-A & B; CERAD; BNT | N/A        | Mixed                             | T-test of difference score | ~72 years/ n = 1,230 |
| Duff et al.19                        | 1 wk           | MCI vs. Normal Cognition | Visual processing speed; Psychomotor speed; Memory (verbal, visual) | HVLT-R; BVMT-R; SDMT; TMT-A & B | Hippocampal volume | Longitudinal                      | Z-score difference         | 77.5 years/ n = 25  |
| Duff et al.20                        | 1wk and 1 year | MCI vs. Normal Cognition | Memory; Non-memory | BVMT-R; HVLT-R; COWAT; Animal fluency; TMT-A & B; SDMT | N/A        | Longitudinal                      | Change score                | 78.7 years/ n = 127 |
| Duff et al.21                        | 2 wks, 3 mos., & 6 mos | MCI | Visual learning; Global cognitive function; Psycho-motor speed; Memory; Language | BVMT-R; MMSE; TMT-A & B; HVLT-R; WAIS-R; BNT; COWAT; SDMT | N/A        | Longitudinal                      | Correlation/ regression    | 72.4 years / 3 different samples (n = 80, n = 33, n = 170) |
| Duff et al.22                        | 1 wk           | MCI vs. Normal Cognition | Memory; Visuospatial constructive skills; Language; Attention | TMT-A & B; SDMT; BVMT-R; COWAT; HVLT-R; Animal fluency | N/A        | Longitudinal                      | Change score                | ~80 years/ n = 121 |
| Duff et al.23                        | Same day       | MCI | Global cognitive function | HVLT-R; MMSE | N/A        | Longitudinal                      | Change score; Correlation/ regression | 73.3 years/ n = 61 |
| Duff et al.24                        | 1 wk           | MCI vs. Normal Cognition | Visual memory | BVMT-R | Amyloid uptake | Longitudinal                      | Correlation/ regression    | 74.6 years / n = 25   |
| Duff et al.25                        | 1 wk           | MCI vs. Normal Cognition | Memory (verbal, visual); Attention & psychomotor speed; Visual scanning/processing speed; Premorbid intellect | HVLT-R; BVMT-R; SDMT; TMT-A & B; WRAT-4 | Amyloid uptake | Longitudinal                      | Correlation/ regression | 77.5 years / n = 27   |

(Continues)
| Research article and reference number | Retest interval | Baseline groups | Cognitive domains | Cognitive tasks | Biomarkers | Cross-sectional/Longitudinal/Mixed | How PEs calculated/quantified | Mean age and total n |
|--------------------------------------|----------------|-----------------|-------------------|----------------|------------|-----------------------------------|---------------------------|-------------------|
| Duff et al.13                        | ~1 wk          | MCI vs. Normal Cognition | Learning & memory; Attention & processing speed; Executive function; Premorbid intellect; Global cognitive function | HVLT-R; BVMT-R; TMT-A & B; SDMT; TMT-A; COWAT; WRAT-4; MMSE | Brain hypo-metabolism | Longitudinal Correlation/ regression | 74.6 years / n = 25 |
| Duff et al.8                         | 1 wk           | MCI vs. Normal Cognition | Memory (objective, immediate, delayed); Premorbid intellect; Executive function; Attention; Language; Visuospatial constructive skills | WRAT-3; 3MS; RBANS; BVMT-R; HVLT-R; COWAT; TMT-A & B; SDMT | N/A | Mixed Change score | ~80.6 years / n = 108 |
| Galvin et al.26                      | Yearly (x6)    | Normal Cognition | Memory (primary, working, episodic, verbal); Visuospatial constructive skills; Language | WMS; BVRT; Word fluency; WAIS; TMT-A; BNT | Amyloid deposition, Braak and Braak stage, Neurofibrillary scores, Lewy bodies, cortical infarcts & hemorrhages | Longitudinal Correlation/ regression | 80.7 years / n = 80 |
| Hanyu et al.27                       | 1 wk           | Amnestic MCI | Logical memory; Language; Global cognitive function | WMS-R; Category fluency; MMSE | N/A | Longitudinal T-test difference | ~76 years / n = 39 |
| Hassenstab et al.9                   | ~2 mos          | Normal Cognition | Memory (episodic, semantic); Executive function; Visuospatial function; Global cognitive function | FCSRT; WMS; WMS-R; WAIS; BNT; Animal Naming; TMT-A & B | APOE genotype | Longitudinal Correlation/ regression | 74.5 years / n = 263 |
| Howieson et al.28                    | Every 6 mos (x3) | Normal Cognition | Memory; Language; Visuospatial constructive skills | WMS; Category fluency; WAIS-R | N/A | Longitudinal Correlation/ regression | 83 years / n = 156 |
| Ihara et al.29                       | 3 yrs           | MCI and Normal Cognition | Memory (episodic); Global cognition; Executive function; Visuospatial function | MMSE; ADAS; WMS-R; WAIS-R; Category fluency; TMT; BNT; CDT; CCT | Amyloid uptake APOE | Longitudinal Correlation/ regression | ~68 years / n = 84 |
| Jonaitis et al.30                    | 4 yrs x1 then 2 yrs x10 | Normal cognition (AD family history vs. none) | Verbal learning & recall; Executive function; Visual learning & memory | RAVLT; BVMT; WMS | AD family history of AD; APOE genotype | Longitudinal Change score | ~54 years / n = 594 |
| Machulda et al.31                    | 15 mos (x3)    | Normal Cognition | Memory; Language; Visuospatial function; Attention/ Executive function | AVL1; WMS-R; BNT; Category fluency; WAIS-R; TMT-B | Hippocampal volume, brain hypometabolism; Amyloid status | Longitudinal Z-score difference | ~75 years / n = 190 |

(Continues)
| Research article and reference number | Retest interval | Baseline groups | Cognitive domains | Cognitive tasks | Biomarkers | Cross-sectional/Longitudinal/Mixed | How PEs calculated/quantified | Mean age/and total n |
|--------------------------------------|----------------|----------------|------------------|----------------|------------|----------------------------------|----------------------------|----------------------|
| Machulda et al.32                    | 15 mos (x2-x5) | Normal Cognition | Memory; Language; Visuospatial function; Attention/ Executive function | AVLT; WMS-R; BNT; Category fluency; WAIS-R; TMT-B | APOE genotype | Longitudinal | Z-score difference | 78.1 years / n = 1,390 |
| Oltra-Cucarella et al.14             | 6-72 mos       | MCI vs. Normal Cognition | Memory | AVLT | APOE genotype | Longitudinal | Z-score difference | ~74 years / n = 1,210 |
| Sanchez-Benavides et al.33          | 6 wks          | Normal Cognition (Family history of AD vs. APOE genotype) | Memory (short-term, working, visual); Processing speed; Visual perception; Coordination; Attention | WAIS-IV | APOE genotype Family history of AD | Longitudinal | T-test difference | 53.43 years / n = 400 |
| Schrijnemaekers et al.34            | 2-3 yrs        | Normal Cognition | Memory | HVLT; MMSE | N/A | Mixed | Correlation/ regression | ~76 years / n = 101 |
| Suchy et al.35                      | 17 mos         | MCI | General cognition status; Attention; Visual-construction ability | DRS-2 | N/A | Longitudinal | Correlation/ regression | ~70 years / n = 75 |
| Wilson et al.36                     | Yearly(x5)     | Normal Cognition group | Memory (episodic, semantic, working); Perceptual speed; Visuospatial ability | WMS; Category fluency; BNT; Word Reading Test; SDMT; Judgment Line Orientation; Standard Progressive Matrices | Tangles, beta-amyloid, and hippocampal volume | Longitudinal | Change point model change score | 78.7 years / n = 567 |
| Zehnder et al.37                    | 2.4 yrs        | AD vs. Normal Cognition | Global cognitive function | CERAD-NAB (German version: included the BNT, MMSE, Animal fluency, Word list I-III, Figures-copy, Word-list delayed recall, Word list recognition, Figures-delayed recall) | N/A | Longitudinal | Z-score difference | ~71.25 / n = 469 |
| Most Frequent                       | 1wk (8 total)  | Normal Cognition (23 total) | Visuospatial/ constructive ability (10 total) | TMT (14 total) | Higher amyloid uptake on amyloid PET scans (4 found an effect and 2 did not) | Longitudinal (24 total) | Correlation/regression (11 total) | Total average (73.57 years / n = 272) |

Abbreviations: 3MS, modified mini-mental state examination; AD, Alzheimer’s disease; ADAS, Alzheimer’s disease assessment scale; AVLT, auditory verbal learning test; BNT, Boston naming test; BVMT(-R), brief visuospatial memory test (revised); CCT = clock copying test; CDT = clock drawing test; CERAD, consortium to establish a registry for Alzheimer’s disease; COWAT, controlled oral word association test; DRS, dementia rating scale; FCSRT, free and cued selective reminding test; HVLT, Hopkins verbal learning test; MCI, mild cognitive impairment; MMSE, mini-mental state examination; RBANS, repeatable battery for the assessment of neuropsychological status; SDMT, symbol digit modalities test; TMT, trail making test; WAIS(-R), Wechsler adult intelligence scale (revised); WMS(-R), Wechsler memory scale (revised); WRAT-3, wide range achievement test.
(n = 23, 85%) recruited cognitively healthy older adults, whereas 15 studies (56%) recruited participants with MCI, and only 3 studies (11%) recruited individuals with AD dementia. Retest intervals ranged from same day to 4 years, where seven studies (26%) included additional multiple retest assessments (Figure 2). Ten articles included visuospatial functioning or construction, nine articles included language, nine articles included attention, eight articles included executive functioning, seven articles included episodic memory, four articles included psychosocial speed, and four articles included working memory. To calculate PEs, 12 (44%) of the studies used correlation or regression analyses, 8 (30%) used z-score or t-test difference scores, and 9 (33%) used a type of change score quantifier.

3.1 | Theme 1: PE variability defines cases

Studies assigned to Theme 1 addressed the hypothesis that individuals with MCI or AD had lower PEs than individuals with normal cognitive functioning. Most of these studies showed that cognitively healthy elderly showed significantly greater PEs on average than individuals with MCI or dementia on both memory and non-memory cognitive tasks. One study found that lower PEs in individuals with MCI could be detected after multiple repeated assessments on the same day. Conversely, Duff et al. reported that some individuals with amnestic MCI had PEs in delayed recall similar to those found in cognitively healthy older adults. However, when the amnestic MCI group was split into those who remained stable (“MCI-stable,” classified as MCI both at baseline and 1 week follow-up) and those who improved and subsequently appeared intact (“MCI-normal,” classified as MCI at baseline but as intact at 1 week follow-up), the latter showed significant PEs, whereas the former did not. Thus, it was the variability of PEs from initial to subsequent testing that served as a potential more reliable diagnostic indicator. Finally, it should be noted that results on whether PEs could differentiate clinical groups independently of baseline cognitive test performance were somewhat contradictory. For example, Zehnder et al. found that quantifying PEs did not add diagnostic accuracy to baseline cognitive test scores when discriminating cognitively healthy from AD participants. In contrast,
Duff et al. showed that the predictive value of PEs was additive to baseline cognitive performance after a 2-h retest interval.23

3.2 Theme 2: PE variability predicts future cognitive decline

All Theme 2 articles addressed the hypothesis that lower PEs observed over time may be an important early indicator of future cognitive decline. This was evidenced by associations between smaller or less robust PEs and (1) subsequent decline in cognitive test scores;8,20,21,35 (2) risk of progression to AD;9 (3) incidence of MCI or dementia14,32,34; and (4) terminal decline.18 Most studies showed that findings varied by cognitive domain, with abundant evidence that lower PEs on episodic memory measures predicted future cognitive decline,9,18,21,32,34 whereas others found that the predictive value of PEs was largely consistent across different cognitive domains.20 Furthermore, it should be noted that studies varied highly in terms of retest intervals, which ranged from single-time retesting at the same day,23 to multiple repeated assessments over several years (Figure 2).

3.3 Theme 3: PE variability associate with AD risk factors and biomarkers

The majority of articles we assigned to Theme 3 demonstrated that lower PEs were associated with AD risk factors and biomarkers indicative of neurodegeneration (Table 3). For example, several studies showed that lower PEs were associated with greater presence of AD risk factors such as apolipoprotein E gene (APOE) ε4 alleles or a positive family history of dementia.14,32,33 In addition, attenuated PEs have been found to be more common in groups with specific neuroimaging markers that are indicative of dementia, such as lower cerebral blood flow,27 brain hypometabolism,13 smaller hippocampal volumes,19 and higher levels of amyloid.24,25,29 More specifically, Machulda et al. showed that lower PEs were associated with lower hippocampal volume and brain hypometabolism regardless of amyloidosis, suggesting that PEs were more closely related to neurodegeneration than amyloid status.31 Studies linking PEs to neuropathology showed contrasting findings. Galvin et al. reported the relationship between PE differences and the presence of AD neuropathology among individuals who had died without a clinical diagnosis of dementia.26 Those with AD pathology had lower PEs on episodic and semantic memory tests than did those without AD pathology. In contrast, Wilson et al. did not identify a relationship between PEs over years and post-mortem neuropathological markers of AD in individuals without dementia.36

4 DISCUSSION

The literature tends to characterize PEs as a nuisance in estimating group-level characteristics such as normative decline over time in advanced age across different clinical and cognitive populations.2,38–40 In addition, in clinical neuropsychological evaluations, neuropsychologists are typically concerned with the question of whether individuals show statistical evidence of change beyond that expected based on average PEs.34–46 In the current systematic review, we focused on a different question, namely whether individual-level PEs in older adults could serve as a marker of clinical status, such that individuals...
TABLE 3 Overview of studies with evidence for associations between PEs and AD risk factors or biomarkers

| Finding                              | Number of articles showing an association | Number of articles not showing an association |
|--------------------------------------|------------------------------------------|---------------------------------------------|
| Presence of ≥ 1 APOE ε4 allele        | 3 [14, 27, 32]                           | 3 [9, 24, 36]                               |
| Higher amyloid uptake on amyloid PET scans | 4 [30-32, 34]                           | 2 [33, 35]                                 |
| Lower hippocampal volume             | 2 [29, 33]                               | 1 [35]                                     |
| Brain hypometabolism on FDG-PET      | 2 [13, 33]                               | 0                                           |
| Family history                       | 1 [36]                                   | 1 [27]                                     |
| Cortical infarcts and hemorrhages    | 1 [34]                                   | 0                                           |
| Lewy bodies                          | 1 [34]                                   | 0                                           |
| Braak stage                          | 1 [34]                                   | 0                                           |

Abbreviations: FDG, fluorodeoxyglucose; PET, positron-emission tomography.

who showed greater PEs would be at lower risk of future negative cognitive/clinical outcomes than those who had lower PEs. Overall, we found consistent evidence from a modest-sized published literature that smaller, less robust PEs on repeated cognitive testing may be an important early indicator of current diagnostic status and future cognitive decline. In addition, lower PEs were associated with risk factors and markers indicative for dementia, such as APOE genotype and biological markers of neurodegeneration.

Our review identified 27 articles evaluating evidence that characterized PEs in older adults can provide a marker for (future) cognitive performance and risk of dementia. Overall, 25 of those articles provided support for one or more of our defined hypotheses (Figure 3). These studies supported the hypotheses that the magnitude of PEs is associated with (1) current cognitive performance or clinical status (Theme 1) and/or (2) future cognitive decline and risk of progression to AD dementia (Theme 2). Most of these studies were cross-sectional or longitudinal comparisons between cognitively healthy older adults and those with MCI or AD.8,15–17 However, one study suggested that characterizing PEs in cognitively healthy individuals at baseline could aid in the prediction of who would develop AD in the future.9 Lower PEs were also found to be more common in groups with specific biological markers (Theme 3), including brain metabolism, hippocampal volume, and amyloid load.13,29,31 Together with Theme 2 findings, these results suggest that the assessment of PEs, in combination with biomarkers, could be used to detect preclinical AD. This would be of particular relevance in the context of current AD research and clinical trials, which are increasingly focusing on earlier, preclinical populations.11,47

**FIGURE 3** Overview of the 25 articles providing evidence for one or more of the a priori defined hypotheses (Theme 1, 2, and/or 3)
Although we identified accumulating evidence of the clinical value of PEs in the study of AD, it should be noted that some findings remain inconclusive. For instance, it is yet unclear whether PEs can predict future cognitive decline beyond baseline cognitive performance. In addition, some studies showed consistent results for PEs across multiple cognitive domains, whereas other studies indicated that the predictive value of PEs was domain-specific. It is difficult to determine whether differences across studies reflected disparate assessment protocols or the selection of only statistically significant findings to present in publications. Findings regarding APOE ε4 status were also contradictory; whereas three studies found that APOE ε4 carriers had lower PEs for memory, three other studies found no such association between APOE ε4 alleles and PEs. Unfortunately, it was difficult to formally compare studies with contrasting findings, as the studies reported varied widely with respect to retest intervals, diagnostic groups, cognitive domains, biomarkers, and methods used to calculate and model PEs (see Table 2). Particularly the latter should be taken into account when integrating the various findings. For example, an empirical change score between the first- and second-time testing captures PEs differently than, for example, a regression-based slope of performance across multiple years. The method that reflects PEs most directly likely depends on multiple factors, such as the number and timing of the repeated assessments. Although a full review of methods used to quantify PEs is beyond the scope of the current article, this is an important topic to address in future research.

Another important issue was the methodological complications of the included studies, such as, for example, small-sample sizes (eg, n < 50), questionable definitions and classification criteria used for diagnostic groups (eg, etiology of MCI unknown, no confirmation of underlying neurodegeneration), combining different clinical groups (eg, cognitively healthy and MCI subjects), potential confounders affecting statistical analyses that were not accounted for, and varied definitions and methods to calculate PE. Due to these overall differences as well as the heterogeneity of study design, participants, and measures, we were not able to conduct a formal meta-analysis or include a funnel plot. However, it should be noted that the potential for publication bias, with respect to statistically significant results for the themes identified in this study, is likely to be high. That is, if we assume that a lack of PE is associated with cognitive decline, and that on average the 27 studies were just adequately powered (ie, had an 80% probability) to detect an association between lack of PE and cognitive decline, we would have only a 7% probability of observing ≥25 of 27 studies returning a significant effect. Moreover, the probability of observing >25 of 27 studies with a positive effect would not reach about 50% until the average power of the 27 studies exceeds 90%. We do not know the power of each of the 27 published studies, but an average of 90% seems implausible. Thus, it would be more plausible that the number of studies performed is higher than 27, with all of the other (non-published) studies not submitted because they had “negative” findings. In this instance, the published literature may over-represent statistically significant findings.

It should also be considered that, as our primary focus was on PEs as a potential marker for cognitive decline, we included only articles that were assigned to Category C. Therefore, we excluded several articles that focused on understanding mechanisms underlying PEs, for example, by determining whether PEs can be attributed to repeated context rather than context effects, or by investigating different aspects of memory (eg, encoding vs. retrieval) as underlying mechanisms of PEs. We note that those and other articles were not overlooked in the review process, but rather fell outside the scope of the current review. On the other hand, novel, potentially eligible, Category C articles may have been published while the current article was in preparation, but were not included as they were not available online before our final search date (January 2020). Although these two articles were not included in this review, they provide further support for the hypothesis that the absence of PEs is a potential marker of cognitive decline.

This review provides an important stepping stone for future work on PE as a useful marker in AD research. To our knowledge, this is the first attempt to summarize the PE literature with a focus on PEs as a potential indicator of subsequent risk rather than as a source of bias in estimating group-level mean cognitive trajectories or as a confounding variable in determining whether an individual patient’s cognitive function has changed on a subsequent testing occasion. We performed a comprehensive, systematic literature search followed by a thorough review process benefiting from a multidisciplinary team with expertise in cognitive aging, psychometrics, neuropsychology, medicine, and library science. This study thus offers a novel valuable perspective on the concept and implications of PEs in the study of cognitive aging and dementia. Most of the included articles supported the idea that lower PEs could reflect subtle learning deficits and thereby represent an early clinical symptom of AD.

The idea that an absence of PEs could indicate (subtle) cognitive impairment, implies that researchers should consider characterizing PEs as a marker or risk factor of cognitive decline in studies of cognitive aging and AD. For example, identifying individuals with lower than expected PEs could potentially serve as a cost-effective strategy to enrich enrollment in longitudinal studies, and predict who might be at higher risk for developing AD. Furthermore, identifying high-risk individuals may be an effective strategy to enroll high-risk individuals in randomized-controlled trials of disease-modifying therapeutics. To develop evidence-based recommendations for enrollment in clinical trials, additional research is needed to determine the possible role of lower PEs for estimating AD risk, and potentially in combination with other approaches such as AD biomarkers. We did not find any evaluations of PEs with respect to longitudinal imaging or fluid biomarker data, so it will be important to relate the magnitude of PEs to longitudinal neuroimaging and biomarker data in future research. Furthermore, methods for calculating PEs also warrant consideration in future work, as we found that those methods varied widely, which may have accounted, in part, for contrasting findings across studies. Using more sophisticated characterization of PEs across specific cognitive domains and applying modern psychometric techniques to develop reliable estimates of learning and practice could improve identification of early stage AD. This, in turn, will help to clarify whether individuals with lower PEs are at higher risk for conversion to AD.
In conclusion, we found accumulating evidence that a lack of PEs may represent an early indicator of future cognitive decline and that lower PEs are associated with specific AD biomarkers. The combination of PEs and these biomarkers may yield an optimal approach to estimate AD risk. Future research could then focus on identifying high-risk cohorts from a combination of cognitive and clinical features, and lower than expected PEs. This cost-effective strategy may be able to enrich samples in clinical trials and provide a valuable cognitive marker for subtle pharmacological or treatment responses.

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

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