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Cognitive Ability in Late Life and Onset of Physical Frailty: The Lothian Birth Cohort 1936

Catharine R. Gale, PhD,*† Stuart J. Ritchie, PhD,* Cyrus Cooper, DM,‡ John M. Starr, PhD,*‡ and Ian J. Deary, PhD*

OBJECTIVES: To investigate whether poorer cognitive ability is a risk factor for development of physical frailty and whether this risk varies according to cognitive domain.

DESIGN: Prospective longitudinal study with 6-year follow-up.

SETTING: Edinburgh, Scotland.

PARTICIPANTS: Members of the Lothian Birth Cohort 1936 (N = 594).

MEASUREMENTS: Frailty was assessed at ages 70 and 76 using the Fried criteria. Cognitive function was assessed at age 70, 73, and 76. Factor score estimates were derived for baseline level of and change in four cognitive domains: visuospatial ability, memory, processing speed, and crystallized cognitive ability.

RESULTS: Higher baseline levels of processing speed, memory, visuospatial ability and crystallized ability at age 70, and less decline in speed, memory, and crystallized ability were associated with less risk of becoming physically frail by age 76. When all cognitive domains were modelled together, processing speed was the only domain associated with frailty risk, for a standard deviation (SD) increment in initial level of processing speed, the risk of frailty was 47% less (0.53 95% confidence interval (CI) = 0.33–0.85) after adjustment for age, sex, baseline frailty status, social class, depressive symptoms, number of chronic physical diseases, levels of inflammatory biomarkers, and other cognitive factor score estimates; for a SD increment in processing speed change (less decline) risk of frailty was 74% less (RR = 0.26, 95% CI = 0.16–0.42). When additional analyses were conducted using a single test of processing speed that did not require fast motor responses (inspection time), results were similar.

CONCLUSIONS: The speed with which older adults process information and the rate at which this declines over time may be an important indicator of the risk of physical frailty. J Am Geriatr Soc 2017.

Key words: fried frailty phenotype; processing speed; memory; visuospatial ability; crystallized ability

Frailty is a clinical syndrome observed in older adults, the core feature of which is greater vulnerability to stressors due to impairments in multiple systems, lower physiological reserves, and a decline in the ability to maintain homeostasis.1 It increases the risk of adverse outcomes.1–3 The phenotype model—in which frailty is based on three or more of five components: poor grip strength, slow walking speed, low physical activity, exhaustion, and unintentional weight loss2—is one of the two principal models of frailty.1 The frailty index, or cumulative deficit model, defines frailty in terms of the accumulation of deficits (symptoms, signs, diseases, disabilities), whereby an individual's frailty index score reflects the proportion of potential deficits present.4 These models differ in the potential role that cognitive impairment plays in their definition of frailty. The Fried phenotype defines frailty in purely physical terms, whereas the cumulative deficit model permits cognitive impairment to be included as a deficit. A consensus conference agreed that this broader definition of frailty should be distinguished from the medical syndrome of physical frailty.5 Given the importance of cognitive function and physical robustness for quality of life and survival, it is crucial to understand the extent to which cognitive ability and physical frailty are associated and the reasons for this.

Physical frailty and poorer cognitive function often coexist.6–8 The direction of this relationship and the underlying mechanisms are uncertain. Some longitudinal studies suggest that physical frailty increases risk of cognitive decline9,10 or dementia.11–13 Poor cognitive function might be a risk factor for becoming physically frail, but evidence

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is sparse. Two longitudinal studies have found that lower Mini-Mental State Examination (MMSE) scores increase the risk of incident physical frailty, but it is unclear whether differences over the range of cognitive ability can predict the onset of physical frailty or whether some domains of cognitive ability are more important as risk factors than others. Results from a longitudinal study found that poorer executive function and greater decline in executive function were more strongly linked to physical frailty than level of or decline in psychomotor speed or memory. Further longitudinal investigations are needed to understand the role of specific cognitive domains in the development of physical frailty.

The Lothian Birth Cohort 1936 (LBC1936) was established to study cognitive aging. Three waves of data on processing speed, memory, visuospatial ability, and crystallized cognitive ability were used to examine how initial level of and change in cognitive function in these domains were related to risk of developing physical frailty or prefrailty.

**METHODS**

**Participants**

The LBC1936 was established to study cognitive aging in surviving members of the 1947 Scottish Mental Survey. Community-dwelling people approximately age 70 were recruited (N = 1,091). Wave 2 took place when participants were approximately age 73 (n = 866). Wave 3 took place when participants were approximately age 76 (n = 697). Ethical approval was obtained from the Multi-Centre Ethics Committee for Scotland and Lothian Research Ethics Committee.

**Measures**

**Physical Frailty**

Frailty status was assessed during Waves 1 and 3 using the Fried phenotype, which defines frailty as the presence of three or more of unintentional weight loss, weakness, self-reported exhaustion, slow walking speed, and low physical activity. Prefrailty is defined as the presence of one or two of these criteria. These criteria were operationalized using definitions similar to Fried’s (Appendix S1).

**Cognitive Ability**

Participants took a variety of cognitive tests in an identical fashion at each wave that were used as indicators of four domains of cognitive ability. Visuospatial ability was assessed according to scores on tests of matrix reasoning and block design from the Wechsler Adult Intelligence Scale (WAIS-IIIUK) and spatial span forward and backward from the Wechsler Memory Scale (WMS-IIIUK). Verbal-declarative memory (henceforth memory) was assessed according to scores on tests of digit-symbol substitution and symbol search from the WAIS-IIIUK and measures of four-choice reaction time and inspection time. Of these measures of speed, inspection time is the only test requiring no speeded responses, in other words, participants are not required to respond as fast as possible. Crystallized cognitive ability was measured using National Adult Reading Test and Wechsler Test of Adult Reading scores, The MMSE was used solely to identify participants with likely cognitive impairment or dementia. With the exception of three of the tests for processing speed that required fast motor responses, none of the tests relied on physical function.

**Covariates**

Age, socioeconomic status, smoking status, number of chronic physical diseases, depressive symptoms, and inflammatory biomarkers were chosen at Wave 1 as potential confounding variables. Assessment details are given in Appendix S2.

**Statistical Analysis**

The cognitive tests were organized into four domains: visuospatial ability, memory, speed, and crystallized ability. An intercept factor (baseline level of the ability) and a slope factor (change in the ability across the three waves) were estimated within each grouping using latent growth curve modelling in a factors-of-curves format. Latent-variable models reduce the influence of test-specific measurement error by using the shared variance between the baseline levels and changes in observed scores on multiple cognitive tests to estimate latent (unobserved) variables of cognitive ability baseline and change. Factor models and score estimates, which used full-information maximum likelihood estimation to use all the data in the full sample at each wave, were produced using Mplus v7.3 (Muthén & Muthén, Los Angeles, CA). Details of the factors-of-curves structural equation models and mean decline in the cognitive test scores over the three waves are given in Appendix S3 and Table S1.

Other analyses were performed in Stata version 13 (Stata Corp., College Station, TX). Multinomial logistic regression was used to calculate relative risks of prefrailty or frailty at age 76 according to a standard deviation (SD) increment in factor score estimates for baseline cognitive ability in each domain and change in cognitive ability in each domain from age 70 to 76, with adjustment for potential confounding factors. Relationships did not vary according to sex, so the data were pooled, and sex was adjusted for. To reduce potential bias due to attrition, all models included inverse probability weights that made the sample more representative of the cohort at baseline. Three of the speed factor tests required fast, accurate motor responses. The fourth, inspection time, required no speeded response. To test whether associations found with the speed factor were artefacts caused by overlap of components of the frailty phenotype measure—slow walking speed and exhaustion—with the motor aspects of three of these tests, models were estimated in which only inspection time baseline and slope were used as predictors. Finally, analyses were repeated excluding participants who scored less than 24 on the MMSE.
RESULTS

Analyses were based on 594 participants with data on all variables of interest. People excluded because of attrition tended to be older; had poorer cognitive ability, more depressive symptoms, more chronic physical disease, and higher blood concentrations of C-reactive protein (CRP) and fibrinogen; were more likely to smoke and less likely to have professional or managerial socioeconomic status; and met more criteria for frailty at age 70. There were no significant differences between those in the sample and those excluded because of missing baseline data, except in level of the cognitive factor “speed,” which was lower in the missing-data group (Table S2).

By age 76, 47.0% of the participants were prefrail, and 14.3% were frail. (At age 70, equivalent figures were 45.5% and 4.9% respectively.) The increase in prevalence of frailty between these ages is similar to that found previously.29 Of those who were frail at age 76, the most common combination of frailty criteria was exhaustion with low activity (30.3%).

Table 1 shows participant characteristics according to frailty status at age 76. Greater frailty at age 76 was associated with older age, more depressive symptoms, more chronic physical disease, being a current smoker, having higher blood concentrations of CRP, and meeting more criteria for frailty at age 70. Greater frailty at age 76 was also associated with lower baseline level of visuospatial ability, memory, speed, and crystallized ability and greater decline in memory and speed between ages 70 and 76.

Table 2 shows the relative risk of incident prefrailty or frailty at age 76 according to a SD increment in factor score estimates for baseline level of cognitive ability in each domain. In models adjusted for age, sex, and number of frailty criteria at baseline, higher factor scores for speed were associated with lower risk of becoming prefrail. This association was attenuated and no longer significant after further adjustment for other covariates and for other cognitive factor score estimates. There were no significant associations between any of the other cognitive factor score estimate levels and risk of becoming prefrail. In initial models, having a higher level of speed or visuospatial ability (but not memory or crystallized ability) was associated with a significantly lower risk of becoming frail by age 76 (becoming frail per SD increment in cognitive factor score estimates: RR = 0.24, 95% CI = 0.17–0.35 for speed; RR = 0.63, 95% CI = 0.42–0.93 for visuospatial ability). Further adjustment in the models of frailty for the other potential confounding factors had only a small attenuating effect on these associations. In a final model with frailty as the outcome, all cognitive factor score estimates were examined simultaneously. In this model, processing speed was the only cognitive domain that was independently associated with risk of becoming frail (SD increment in speed: RR = 0.53, 95% CI = 0.33–0.85). When changes in depressive symptoms, in chronic physical illnesses, and in inflammatory markers between Waves 1 and 3 were adjusted for in place of these measures at Wave 1, results were similar (SD increment in speed: RR = 0.46, 95% CI = 0.28–0.77).

Table 3 shows RRs for incident prefrailty or frailty according to a SD increment in factor score estimates for the slope of the trajectory of cognitive ability in each domain between ages 70 and 76. Higher factor score estimates for change in speed and in visuospatial ability—indicating less decline—were associated with lower risk of becoming prefrail. No other cognitive domain was independently associated with prefraility. In initial models, for a SD increment in cognitive factor change—indicating less decline—the risk of pre-fraility was less by 56% in the case

Table 1. Characteristics of Study Sample at Age 70 According to Frailty at Age 76

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Not Frail, n = 230</th>
<th>Prefrail, n = 279</th>
<th>Frail, n = 85</th>
<th>P-Value for Difference*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean ± SD</td>
<td>69.4 ± 0.83</td>
<td>69.5 ± 0.81</td>
<td>69.7 ± 0.77</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Depressive symptom score, median (IQR)</td>
<td>1 (0–2)</td>
<td>1 (0–2)</td>
<td>2 (1–3)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Number of frailty criteria, median (IQR)</td>
<td>0 (0–1)</td>
<td>0 (0–1)</td>
<td>2 (1–2)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Number of chronic diseases, median (IQR)</td>
<td>0 (0–1)</td>
<td>1 (0–1)</td>
<td>1 (1–2)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Fibrinogen, g/L, median (IQR)</td>
<td>3.1 (2.7–3.5)</td>
<td>3.2 (2.8–3.5)</td>
<td>3.2 (2.9–3.7)</td>
<td>.14</td>
</tr>
<tr>
<td>C-reactive protein, mg/L, median (IQR)</td>
<td>1.5 (1.5–5)</td>
<td>3 (1.5–6)</td>
<td>4 (1.5–7)</td>
<td>.02</td>
</tr>
<tr>
<td>Female, n (%)</td>
<td>108 (47.0)</td>
<td>139 (49.8)</td>
<td>44 (51.7)</td>
<td>.70</td>
</tr>
<tr>
<td>Current smoker, n (%)</td>
<td>11 (4.78)</td>
<td>17 (6.09)</td>
<td>11 (12.9)</td>
<td>.03</td>
</tr>
<tr>
<td>Professional or managerial social class, n (%)</td>
<td>144 (62.6)</td>
<td>166 (59.5)</td>
<td>47 (55.3)</td>
<td>.48</td>
</tr>
<tr>
<td>Cognitive factor score estimates for baseline level, mean ± SD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visuospatial ability</td>
<td>0.32 ± 0.85</td>
<td>0.17 ± 0.84</td>
<td>−0.42 ± 0.92</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Memory</td>
<td>0.24 ± 0.79</td>
<td>0.12 ± 0.80</td>
<td>−0.29 ± 0.82</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Speed</td>
<td>0.46 ± 0.81</td>
<td>0.18 ± 0.76</td>
<td>−0.54 ± 1.01</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Crystallized ability</td>
<td>0.23 ± 0.94</td>
<td>0.12 ± 0.93</td>
<td>−0.29 ± 1.07</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Cognitive factor score estimates for slope, mean ± SD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visuospatial ability</td>
<td>−0.02 ± 0.50</td>
<td>−0.04 ± 0.51</td>
<td>−0.01 ± 0.56</td>
<td>.85</td>
</tr>
<tr>
<td>Memory</td>
<td>0.10 ± 0.68</td>
<td>−0.03 ± 0.75</td>
<td>−0.28 ± 0.82</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Speed</td>
<td>0.23 ± 0.54</td>
<td>−0.01 ± 0.69</td>
<td>−0.42 ± 0.81</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Crystallized ability</td>
<td>−0.02 ± 0.87</td>
<td>−0.03 ± 1.08</td>
<td>0.003 ± 0.06</td>
<td>.98</td>
</tr>
</tbody>
</table>

*From analysis of variance, Kruskal–Wallis, or chi-square tests as appropriate.
SD = standard deviation; IQR = interquartile range.
of speed (RR= 0.44, 95% CI 0.32 - 0.62), and by 24% in the case of visuospatial ability (RR=0.76, 95% CI = 0.53 - 0.98) were (RR = 0.44, 95% CI = 0.32, 0.62 for speed; RR = 0.76, 95% CI = 0.53–0.98 for visuospatial ability). The association between change in speed and risk of prefrailty changed little in subsequent models, but the association between change in visuospatial ability and risk of prefrailty ceased to be significant when adjusted for other cognitive factor score estimates. In initial models of frailty, higher factor score estimates for change in speed and memory—indicating less decline—were associated with lower risk (becoming frail per SD increment in cognitive factor change: RR = 0.20, 95% CI = 0.13–0.32 for speed; RR = 0.48, 95% CI = 0.33–0.70 for memory). Further adjustment for the other covariates had only a small attenuating effect. In the final model, higher estimate for change in speed was the only cognitive factor score estimate that remained significantly associated with lower risk of frailty (SD increment: RR = 0.26, 95% CI = 0.16–0.42). When changes in depressive symptoms, chronic physical illnesses, and inflammatory markers between Waves 1 and 3 were adjusted for in place of these measures at Wave 1, the association between change in speed and risk of frailty was very similar (SD increment in speed: RR = 0.28, 95% CI = 0.17–0.46).

Table 2 shows RRs for incident prefrailty or frailty according to SD increments in baseline level and change in inspection time. Results were similar to those obtained using the speed factor estimates.

The analyses were repeated excluding those who scored less than 24 on the MMSE at all three waves (n = 27). Results were almost unchanged (data not shown).

A sensitivity analysis was performed with those who were physically robust at age 70 (n = 295). Effect sizes were very similar to those presented in Tables 2 and 3; speed was the only cognitive domain associated with frailty risk in the fully adjusted models (SD increment in baseline level of speed: fully adjusted RR = 0.78, 95% CI = 0.53–1.14 for prefrailty; RR = 0.24, 95% CI = 0.09–0.61 for frailty; SD increment in change in speed: RR = 0.49, 95% CI = 0.31–0.79 for prefrailty; RR = 0.23, 95% CI = 0.10–0.57 for frailty).

DISCUSSION
To the knowledge of the authors, only one study has examined the relationship between different cognitive abilities and onset of physical frailty. In 331 women from the Women's Health and Aging Study, higher initial level of

Table 3. Risk of Incident Physical Prefrailty and Frailty at Age 76 According to Change in Cognitive Function (Slope) Between Age 70 and 76

<table>
<thead>
<tr>
<th>Cognitive Factor Score Estimates for Slope, per Standard Deviation</th>
<th>Adjusted for Age, Sex, and Components of Frailty Present at Age 70</th>
<th>Further Adjusted for Depressive Symptoms, Chronic Physical Diseases, Social Class, Inflammatory Biomarkers, and Smoking Status at Age 70</th>
<th>Further Adjusted for Other Cognitive Factor Score Estimates</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Prefrail</td>
<td>Frail</td>
<td>Prefrail</td>
</tr>
<tr>
<td>------------------------------------------------------------------</td>
<td>----------</td>
<td>-------</td>
<td>----------</td>
</tr>
<tr>
<td>Relative Risk (95% Confidence Interval)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visuospatial ability</td>
<td>0.72 (0.53–0.98)</td>
<td>0.65 (0.40–1.06)</td>
<td>0.71 (0.52–0.98)</td>
</tr>
<tr>
<td>Memory</td>
<td>0.80 (0.62–1.03)</td>
<td>0.48 (0.33–0.70)</td>
<td>0.79 (0.61–1.02)</td>
</tr>
<tr>
<td>Speed</td>
<td>0.44 (0.32–0.62)</td>
<td>0.29 (0.19–0.39)</td>
<td>0.47 (0.34–0.65)</td>
</tr>
<tr>
<td>Crystallized ability</td>
<td>0.93 (0.77–1.12)</td>
<td>0.91 (0.69–1.19)</td>
<td>0.92 (0.776 1.11)</td>
</tr>
</tbody>
</table>

All estimates are weighted to adjust for attrition since baseline.
and slower decline in executive function—assessed using a single test—were associated with lower risk of physical frailty. Participants were also assessed for psychomotor speed and immediate and delayed verbal memory—again using single tests. Higher scores for speed, delayed verbal memory only, and general cognitive performance were associated with lower risk, but there were no significant associations between rate of decline on any cognitive test other than the test of executive function and physical frailty risk. The measure used to assess executive function in that study (the Trail-Making Test) may also reflect processing speed, conforming to the findings in the current cohort.

In the present study, initial level of and decline in memory and speed were associated with frailty risk. Speed seemed to be the more-powerful predictor of physical frailty because it was associated with risk independent of covariates and other cognitive domains; for a SD increment in initial level of speed or change in speed (less decline), risk of frailty was 47% or 74% less, respectively. To check whether overlap between the speed of motor response required by some tests of processing speed and the slow walking speed or exhaustion components of the frailty phenotype might produce these associations, the analyses were repeated using the psychophysical inspection time test as the sole measure of processing speed; this test of speed of visual discrimination does not rely on physical reactions. Effect sizes using this single test were smaller than those obtained using the speed factor—for a SD increment in baseline level of or change in inspection time, risk of frailty was 40% or 35% lower, respectively, after full adjustment—but these results demonstrate that the link between processing speed and risk of frailty is not arifactual. Processing speed may be an early signal of impending limitations in a number of physical–mental domains, with some underlying shared causes. There is evidence that greater decline in processing speed is associated with greater decline in walking speed, and in the current cohort, decline in processing speed, as measured according to inspection time, was strongly correlated with decline in general cognitive ability.

The mechanisms underlying associations between domains of cognitive ability, in particular speed, and risk of physical frailty remain unclear. Adjustment for covariates had modest attenuating effects. Neuropathology that has an adverse effect on cognitive function may also influence risk of physical frailty. Support for this comes from findings that rates of change in physical frailty and cognitive function were strongly correlated and that Alzheimer’s disease pathology, macroinfarcts, and nigral neuronal loss were associated with prior rates of change in physical frailty and cognitive ability. Disruption of connectivity in white matter affects processing speed and walking speed. Further investigation in this cohort could test whether this is the mechanism underlying these findings. Another explanation might be that some common biological process of cellular senescence underlies the associations. Cellular senescence is a stress response that occurs when cells are exposed to potentially oncogenic stimuli. Senescent cells appear with increasing frequency in older tissues. The secretion of proinflammatory cytokines, growth factors, and proteases that accompanies cellular senescence may be implicated in cognitive decline and physical frailty.

Strengths of this study include the characterization of each domain of cognitive function over three waves, enabling how initial level and change were related to onset of physical frailty or prefrailty to be examined. Other strengths are the narrow age range, data on a range of potential confounding factors, and the fact that the sample was of both sexes. One limitation is that, for some individuals, decline in cognitive ability and onset of physical frailty will have begun before age 70, making it uncertain whether poorer cognitive ability predates later frailty or whether cognitive and physical health are declining together. The finding that slower processing speed was as predictive of frailty in the subset of participants who were physically robust as in the whole sample suggests that poorer cognitive ability may increase the risk of frailty. A second limitation is that, largely because of attrition, the analyses were based on 54% of participants in the baseline survey. Attrition can result in biased estimates if there are differences in likelihood of follow-up related to exposure and outcome. In the analytical sample, higher baseline levels of processing speed were associated with lower risk of becoming physically frail. The risk of becoming physically frail was likely to have been higher in those lost to follow-up because they tended to be in poorer health and were frailer at baseline (Table S2). Those lost to follow-up (and those excluded because of missing data) also differ from the analytical sample in having lower levels of processing speed. The models were weighted to reduce potential bias due to attrition, but the
results may underestimate the predictive power of processing speed regarding risk of physical frailty.

The speed with which older people process information and the rate at which this declines may be important indicators of the risk of becoming physically frail. More research into cognitive domain-specific associations and risk of physical frailty is needed to confirm the importance of different domains for predicting onset of frailty and elucidate the underlying mechanisms.

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Conflict of Interest: None.

Author Contributions: CRG, IJD: Study concept. IJD, JMS: Participant recruitment, collection of data. SJR, CRG: Data analysis. CRG: Drafting of first version of manuscript. All authors: Data interpretation, final version of manuscript.

Sponsor’s Role: None.

REFERENCES


SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Table S1. Slope means for each cognitive test across the three waves (age 70 to age 76).

Table S2. Characteristics at wave 1 of participants included and excluded from the analytical sample.

Appendix S1. Operationalising the Fried phenotype of frailty criteria.

Appendix S2. Assessment of covariates.

Appendix S3. ‘Factors of curves’ structural equation models of the cognitive ability test scores.

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