Prevalence of Mild Cognitive Impairment in the Lothian Birth Cohort 1936

Miles Welstead, MSc,* Michelle Luciano, PhD,* Graciela Muniz-Terrera, PhD,† Adele M. Taylor, MA,* and Tom C. Russ, PhD, MRCPsych†‡

BACKGROUND: The Lothian Birth Cohort 1936 (LBC1936) is a highly phenotyped longitudinal study of cognitive and brain ageing. Given its substantial clinical importance, we derived an indicator of mild cognitive impairment (MCI) and amnestic and nonamnestic subtypes at 3 time points.

Methods: MCI status was derived at 3 waves of the LBC1936 at ages 76 (n = 567), 79 (n = 441), and 82 years (n = 341). A general MCI category was derived as well as amnestic MCI (aMCI) and non-amnestic MCI (naMCI). A comparison was made between MCI derivations using normative data from the LBC1936 cohort versus the general UK population.

Results: MCI rates showed a proportional increase at each wave between 76 and 82 years from 15% to 18%. Rates of MCI subtypes also showed a proportional increase over time: aMCI 4% to 6%; naMCI 12% to 16%. Higher rates of MCI were found when using the LBC1936 normative data to derive MCI classification rather than UK-wide norms.

Conclusions: We found that MCI and aMCI rates in the LBC1936 were consistent with previous research. However, naMCI rates were higher than expected. Future LBC1936 research should assess the predictive factors associated with MCI prevalence to validate previous findings and identify novel risk factors.

Key Words: MCI, cognitive aging, amnestic, nonamnestic, prevalence

Alzheimer Dis Assoc Disord 2021;35:230–236

BACKGROUND

In conjunction with advancements in health and social care in the past century, life expectancy has improved dramatically and contributed to a rapidly increasing older population.1 A consequence of this demographic shift is the challenge we now face to care for a larger number of older adults with susceptibility to cognitive deterioration.2 Understanding how cognitive decline affects older people is imperative in order to design interventions to slow or delay decline and ensure individuals are on the healthiest aging trajectory possible.3 Decline in memory is a key indicator of dementia: however, it is common in older age, and differences between normal age-related decline and the early stages of dementia can be difficult to differentiate.4

The concept of mild cognitive impairment (MCI) traces back many years but has gained particular traction over the past few decades.5 Petersen et al6 popularized the concept as a distinct clinical condition and established a set of criteria based on memory changes without loss of ability to undertake normal activities. These criteria heavily influenced the way in which MCI was, and continues to be, identified in research and clinical settings. However, other researchers such as Dubois and Albert7 disputed the notion of MCI as a distinct clinical entity, instead proposing it as a stage of severity for particular disorders. Accordingly, they proposed a “prodromal Alzheimer disease” based upon subjective memory complaints with progressive onset, preserved ability to undertake activities of daily living, neuroimaging, and biomarker testing. Disagreement on how MCI should be conceptualized has led to multiple attempts at an international consensus. Winblad et al8 reached consensus that MCI criteria should assess whether an individual has a dementia diagnosis, whether their cognition has shown subjective and/or objective decline over time, and whether their activities of daily living are significantly affected—and, indeed, how this latter criterion is judged.

This groundwork informed the most recent guidelines proposed by the National Institute on Aging-Alzheimer’s Association (NIA-AA) workgroups on diagnostic guidelines for Alzheimer disease.9 These guidelines propose 4 criteria based on: (1) concern regarding a change in cognition, (2) impairment in 1 or more cognitive domains, (3) preservation of independence in functional abilities, and (4) no diagnosis of dementia. In addition to identifying general MCI, there has also been increased interest in identifying specific subtypes of MCI that may precede certain types of dementia. For instance, amnestic MCI (aMCI) focuses solely on memory-related cognitive impairment, whereas nonamnestic MCI (naMCI) focuses on cognitive impairment in other domains such as processing speed, attention, and executive functions.10 Associations found between these subtypes and the risk of converting to dementia depend heavily on how the measures are defined and the population in which they are implemented. With this in mind, some research has indicated that aMCI may be associated with an increased risk of converting to Alzheimer disease, while naMCI is linked with other types of dementia such as diffuse Lewy body dementia.11 Another caveat of using MCI and its subtypes as a measure of subclinical cognitive impairment is...
that there is great debate surrounding its clinical utility. It remains contested as to how useful MCI is in a clinical context, what it actually captures, and whether other measures such as other cognitive impairment no dementia, which does not factor in functionality, may in fact provide a better estimate of those at high risk of developing dementia.\textsuperscript{12} Despite this, MCI can be a useful tool to capture cognitive decline in research. By identifying MCI and its subtypes it will potentially allow for improved knowledge on how early prevention strategies can identify individuals who are at high risk of cognitive decline and subsequent dementia. Here we use the NIA-AA guidelines to derive an identification of MCI and its subtypes using data from the Lothian Birth Cohort 1936.\textsuperscript{13,14} We hypothesize that MCI rates will be similar to those found in other older adult cohorts and that prevalence of all types of MCI will be higher in later data waves.

\textbf{METHODS}

At Wave 1, the LBC1936 study consisted of 1091 participants, born in 1936 with a mean age of 69 (SD = 0.89) years, mostly surviving members of the Scottish Mental Survey 1947.\textsuperscript{15} Wave 1 took place between 2004 and 2007, with follow-up waves approximately every 3 years thereafter at ages: 73 (n = 866), 76 (n = 697), 79 (n = 550), and 82 years (n = 431). More details on recruitment and testing procedures have been published previously.\textsuperscript{13,14,16} The LBC1936 study was conducted according to the Declaration of Helsinki guidelines. Ethical permission for the LBC1936 study was conducted according to the Declaration of Helsinki guidelines. Ethical permission for the LBC1936 study protocol was obtained from the Multi-Centre Research Ethics Committee for Scotland (Wave 1: MREC/01/0/56), the Lothian Research Ethics Committee (Wave 1: LREC/2003/2/29), and the Scotland A Research Ethics Committee (Waves 2, 3, 4 and 5: 07/MREC00/58). Written consent was obtained from participants at each of the waves.

\textbf{Identification of MCI}

Using data previously collected in the LBC1936, an algorithm was created which identifies participants who fulfill the MCI criteria as outlined by the NIA-AA workgroups on diagnostic guidelines for Alzheimer’s disease.\textsuperscript{9} Variables necessary to conduct MCI coding were collected from Wave 3 (age 76) onwards. In order to be classified in the MCI category, participants must have shown met all 4 criteria reported below:

1. Concern regarding a change in cognition: self-reported memory problems that are interfering with their life, as recorded in a questionnaire at each wave.

2. Impairment in 1 or more cognitive domains: scores at least 1.5 SD below the mean on at least 1 cognitive domain (memory, executive function, attention, language, or visuospatial skills) and either shows a decline from the previous wave to below the 10th percentile on 1 test, a decline from wave 1 to below the 20th percentile on 1 test, or a decline from the previous wave to below the 20th percentile on 2 tests.

3. Preservation of independence in functional abilities: scores at least 1.5 SD below the mean on the Townsend Disability Scale overall score.\textsuperscript{17}

4. No diagnosis of dementia: does not self-report or have a formal diagnosis of dementia and scores at least 24 on the Mini-Mental State Examination (MMSE).\textsuperscript{18}

Cognitive domains were assessed using the following cognitive tests: Symbol Search, Digit Symbol Coding, Matrix Reasoning, Letter-Number Sequencing, and Block Design from the Wechsler Adult Intelligence Scale III (WAIS) and Logical Memory I \& II from the Wechsler Memory Scale III (WMS-III).\textsuperscript{19} A cut-off of \( \geq 1.5 \) SD below the mean or scoring below specific percentiles was used to indicate cognitive impairment. Consistent with previous research co-authored by the creators of the NIA-AA guidelines,\textsuperscript{20} cognitive decline was determined as a decline from the previous wave to below the 10th percentile on 1 test, a decline from wave 1 to below the 20th percentile on 1 test, or a decline from the previous wave to below the 20th percentile on 2 tests. Two versions of the cognitive impairment criterion were conducted using the means and SD of individual tests from (1) the LBC1936 sample at each wave and (2) a more representative UK sample provided by the WAIS-III-WMS-III technical manual.\textsuperscript{19} Preliminary comparisons showed that fewer participants were identified as having MCI using the general population norms, likely due to the higher rates of overall healthiness in the LBC1936.\textsuperscript{14} Therefore, the definition using UK normative data were used here as they were more reflective of the general population.

We also coded 2 subtypes of MCI: aMCI and naMCI. Creation of these subtypes followed the same procedure as for the general MCI; however, aMCI was only identified if the participant showed impairment in the memory domain. Similarly, classification for naMCI was met if the participant showed impairment in cognitive domains other than memory (executive function, attention, language, or visuospatial skills).

\textbf{Covariates}

We examined the association between a range of covariates and MCI status. Covariates included: age, sex, years of education, age 11 cognitive function, body mass index; calculated in the standard way of kg/m\(^2\), occupational social class (professional/manag/erial/skilled, non-manual/skilled manual or semiskilled/unskilled), APOE ε4 status (allele present/absent), self-reported history of cardiovascular disease, self-reported history of stroke, depression, and physical frailty level (not frail/prefrail/frail). Physical frailty was derived using the Fried Phenotype guidelines,\textsuperscript{21} for information on how this was calculated in LBC1936 see Welstead et al.\textsuperscript{22} Depression was measured using the Hospital Anxiety and Depression scale.\textsuperscript{23} Age 11 cognitive function was based on LBC1936 participant’s scores on the Moray House Test (MHT) at age 11\textsuperscript{24}; for more detail see Taylor et al.\textsuperscript{14} To adjust for age in days at time of testing, MHT11 scores were residualized for age at 11 years.

\textbf{Statistical Analysis}

Three participants had been diagnosed with dementia before age 76 (wave 3) by the LBC1936 study doctor and were excluded, leaving 694 participants at that wave. In addition, since a wide variety of variables were required in order to derive an MCI coding, missing data at each wave meant that some participants were excluded from analyses (wave 3: n = 127, wave 4; n = 106, wave 5; n = 87). Accordingly, MCI status was coded for 567 participants at wave 3 (age 76), 441 at wave 4 (age 79), and 341 at wave 5 (age 82). Descriptive analyses including number and percentages of people with MCI were used to characterize the study sample. Linear model analysis of variance and Pearson \( \chi^2 \) tests were used to assess characteristics associated with MCI and...
RESULTS

Figure 1 show the rates of MCI in the LBC1936. There was an increase in people with MCI over time with 15% at wave 3 (n = 87/567), 17% at wave 4 (n = 77/441), and 18% at wave 5 (n = 62/341) having MCI. As there were a substantial number of participants who withdrew from the study between baseline and final follow-up, we also looked at MCI rates for completers only, that is, those who completed waves 3, 4, and 5. Results showed an overall proportional increase over follow-up with 14% of completers identified as having MCI at wave 3 (n = 38/271) and wave 4 (n = 38/271), and then a rise to 21% at wave 5 (n = 57/271).

MCI rates did not differ significantly by sex at any of the waves. The only significant differences found indicated that higher rates of MCI were associated with APOE ε4 status at wave 3 (P < 0.001) and wave 5 (P < 0.05), and history of stroke at wave 3 (P < 0.01) and wave 5 (P < 0.05). Covariate differences according to MCI status are reported in Table 1.

MCI Subtypes

We also derived 2 subtypes of MCI: aMCI and naMCI. As reported in Figure 2, proportions of aMCI remained fairly low across follow-up from 4% at wave 3 (n = 24/604), to 4% at wave 4 (n = 21/484), and 6% at wave 5 (n = 24/376). Prevalence of naMCI was higher and showed a gradual proportional increase over follow-up from 12% at wave 3 (n = 73/609), to 14% at wave 4 (n = 63/466), and 16% at wave 5 (n = 56/361).

Normative Data Comparisons

We compared whether MCI rates were sensitive to the use of different normative data. Comparisons were made between MCI rates when using normative data based on the LBC1936 and a UK-wide sample to derive the identification of MCI. As might be expected with a healthy cohort, at all waves there were higher proportions of MCI when using the LBC1936 norms compared with the UK based norms. Supplementary Figure 1 (Supplemental Digital Content 1, http://links.lww.com/WAD/A319) reports MCI rates at each wave according to the LBC1936 normative data.

DISCUSSION

We found MCI proportions in the LBC1936 study of 15%, 17%, and 18% at ages 76, 79, 82 years, respectively. Similar proportions were found when looking only at the individuals who attended all waves. MCI status at wave 3 and wave 5 (but not wave 4) was significantly associated with APOE ε4 status and history of stroke. Proportions of people with aMCI were 4% at ages 76 and 79 years and 6% at 82 years, whereas rates of naMCI were higher but still showed an increase in proportions from 12% at age 76 years to 14% and 16% at 79 and 82 years, respectively.

Comparison With Other Literature

We observed higher rates of MCI in men, albeit not at a statistically significant level, a finding that is consistent with some previous research, but not all. As discussed by Xue et al, sex differences in MCI research are inconsistent and may differ according to alternate methods of deriving MCI. Importantly, the assessment of day-to-day function in men and women presents different challenges, and perhaps surprisingly, there were minimal significant associations between groups of individuals defined by key features. At 2 of the time points APOE ε4 status was associated with having MCI, a finding which has been consistently found in previous MCI research and is also strongly linked to the risk of progression to dementia. The only other characteristic associated with MCI change was having a history of stroke, again somewhat unsurprising given the extensive evidence that stroke patients have higher risk for developing of MCI and dementia. The lack of significant association between these factors and MCI status at wave 4 is unexpected and not readily explained. However, it may be related to attrition or other factors leading to sample differences at wave 4; the proportion of participants with MCI who had an APOE ε4 allele...
As expected, findings also showed an increase in proportion of participants with MCI at wave 5 compared with wave 3. The rates of MCI we find are consistent with previous research using the same MCI coding guidelines which reports an average prevalence of 14.8% for 70 to 75 year olds.32 The rates of 2 subtypes of MCI—aMCI and naMCI—were in partial agreement with previous literature. Some previous research10 has found rates of around 3% to 4% of both aMCI and naMCI in older populations, while others have found 11% for aMCI and 5% prevalence for naMCI.28 Thus, while the aMCI results are expected, the rates of naMCI in the LBC1936 are higher than anticipated. Higher rates of naMCI than aMCI may indicate that participants of the LBC1936 are more prone to nonamnestic cognitive impairment in areas such as language, visual-spatial skills, attention, or executive functioning. Another possibility is that the salient memory problems associated with aMCI may make participants more likely to withdraw from the study, whereas the cognitive problems associated with naMCI (executive function, attention, language, or visuospatial skills) may more often go unnoticed by the participant. However, it is also important to note that making comparisons between our proportions of aMCI and naMCI cannot be done entirely accurately given that cases of missing data differed between them.

### TABLE 1. Covariate Descriptive Statistics for Participants With MCI Present Versus Absent

| Variables                                      | Wave 3 MCI Absent (N = 480) | Wave 3 MCI Present (N = 87) | Wave 4 MCI Absent (N = 364) | Wave 4 MCI Present (N = 77) | Wave 5 MCI Absent (N = 279) | Wave 5 MCI Present (N = 62) | P       |
|------------------------------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|---------|
| Age at wave 3, mean (SD)                        | 76.25 (0.68)                | 76.21 (0.66)                | 76.23 (0.68)                | 76.13 (0.69)                | 76.20 (0.69)                | 76.17 (0.72)                | 0.55*   |
| Sex, n (%)                                      |                             |                             |                             |                             |                             |                             | 0.07†   |
| Male                                           | 248 (52)                    | 54 (62)                     | 183 (50)                    | 43 (56)                     | 133 (48)                    | 38 (61)                     | 0.37†   |
| Female                                         | 232 (48)                    | 33 (38)                     | 181 (50)                    | 34 (44)                     | 146 (52)                    | 24 (39)                     | 0.05†   |
| Years of education, mean (SD)                  | 10.81 (1.13)                | 10.76 (1.16)                | 10.90 (1.19)                | 10.87 (1.14)                | 10.91 (1.17)                | 11.10 (1.17)                | 0.70*   |
| Age 11 cognitive function, mean (SD)           | 1.21 (11.70)                | 1.30 (11.28)                | 1.75 (11.41)                | 1.73 (12.01)                | 2.30 (11.24)                | 2.34 (10.91)                | 0.95*   |
| Missing data                                    | 29                          | 7                           | 25                          | 3                           | 16                          | 6                           | 0.23*   |
| Depressive symptoms, mean (SD)                 | 2.68 (2.20)                 | 3.00 (2.13)                 | 2.55 (2.13)                 | 3.04 (2.30)                 | 2.42 (1.95)                 | 2.97 (2.04)                 | 0.63†   |
| History of cardiovascular disease, n (%)       |                             |                             |                             |                             |                             |                             | 0.05*   |
| No                                             | 327 (68)                    | 50 (58)                     | 240 (67)                    | 49 (64)                     | 185 (67)                    | 37 (61)                     | 0.003†  |
| Yes                                            | 153 (32)                    | 37 (42)                     | 121 (33)                    | 28 (36)                     | 91 (33)                     | 24 (39)                     | 0.71†   |
| History of stroke, n (%)                       |                             |                             |                             |                             |                             |                             | 0.017†  |
| No                                             | 433 (90)                    | 69 (79)                     | 323 (89)                    | 70 (91)                     | 254 (92)                    | 50 (82)                     | 0.001†  |
| Yes                                            | 47 (10)                     | 18 (21)                     | 38 (11)                     | 7 (9)                       | 22 (8)                      | 11 (18)                     | 0.22†   |
| Social class, n (%)                            |                             |                             |                             |                             |                             |                             | 0.364†  |
| Professional                                    | 98 (21)                     | 19 (22)                     | 88 (24)                     | 16 (21)                     | 68 (25)                     | 16 (27)                     | 0.85†   |
| Managerial                                     | 189 (40)                    | 35 (41)                     | 136 (38)                    | 36 (47)                     | 106 (38)                    | 24 (41)                     | 0.36*   |
| Skilled                                        | 102 (21)                    | 11 (13)                     | 76 (21)                     | 10 (13)                     | 62 (23)                     | 12 (20)                     | 0.22†   |
| nonmanual                                      |                             |                             |                             |                             |                             |                             | 0.85†   |
| Skilled manual                                  | 70 (15)                     | 18 (21)                     | 50 (14)                     | 14 (19)                     | 32 (11)                     | 7 (12)                       | 0.22†   |
| Semiskilled/unskilled                          | 16 (3)                      | 3 (3)                       | 11 (3)                      | 0                           | 8 (3)                       | 0                           | 0.22†   |
| Missing data                                    | 5                            | 1                            | 5                            | 1                            | 5                            | 1                            | 0.36†   |
| APOE ε4 status, n (%)                           |                             |                             |                             |                             |                             |                             | 0.018†  |
| Absent                                         | 332 (74)                    | 47 (55)                     | 241 (70)                    | 46 (63)                     | 195 (74)                    | 34 (59)                     | 0.75†   |
| Present                                        | 118 (26)                    | 38 (45)                     | 102 (30)                    | 27 (37)                     | 68 (26)                     | 24 (41)                     | 0.36†   |
| Fried phenotype status, n (%)                  |                             |                             |                             |                             |                             |                             | 0.89†   |
| Not frail                                      | 197 (41)                    | 33 (38)                     | 160 (45)                    | 30 (39)                     | 135 (49)                    | 29 (47)                     | 0.75†   |
| Prefrail                                       | 224 (47)                    | 41 (47)                     | 164 (45)                    | 35 (45)                     | 119 (43)                    | 26 (43)                     | 0.36†   |
| Frail                                          | 59 (12)                     | 13 (15)                     | 37 (10)                     | 12 (16)                     | 22 (8)                      | 6 (10)                       | 0.36†   |
| Missing data                                    | 0                            | 0                            | 3                            | 0                            | 3                            | 1                            | 0.89†   |

* Linear model analysis of variance.
† Pearson χ² test.
MCI indicates mild cognitive impairment.

Copyright © 2021 The Author(s). Published by Wolters Kluwer Health, Inc. www.alzheimerjournal.com | 233
Limitations and Strengths

LBC1936’s rates of high physical health and cognitive ability is well documented,\textsuperscript{14,16} and highlights a limitation of this study: our sample is less representative of the general population who likely have higher rates of MCI. An additional limitation that affects the accuracy of our results was that there was a relatively small number of participants who were identified as having aMCI, which introduces an element of uncertainty into our results. For the participants who withdrew from the study, we did not have systematic information on their reason for dropping out. It is likely that at least some of these participants dropped out due to MCI or dementia, and accordingly we were unable to consider these cases in our analyses. Related to this, other than 3 cases in which we had confirmation from the LBC1936 study doctor, we relied primarily on the self-reporting of dementia diagnoses for part of the MCI criteria. This could have introduced bias if additional participants had a dementia diagnosis but did not report it. While self-reporting is used extensively in epidemiological studies and biases are usually insignificant,\textsuperscript{33} given the nature of dementia, using these measures may have introduced inaccuracies. Current work is being undertaken in the LBC1936 to ascertain dementia status for every participant and so future research will be able to revisit this.

The strengths of this study are our use of data collected at multiple time points over the course of ~6 years in a well-characterized longitudinal cohort study. Using more than 1 time point gives us better insight into how MCI proportions change over time in the LBC1936. An additional strength is that we derived and compared an MCI coding using normative cognitive data from the LBC1936 sample and the UK wide norms. By doing so, we were able to assess the extent to which the LBC1936 data are representative of the wider population. As anticipated, MCI rates were higher at all waves when using the LBC1936 norms, presumably due to an overestimation caused by the higher rates of healthiness found in the LBC1936 when compared with the general population. Deriving MCI using the cohort’s own normative data will cause the cognitive impairment cut-off points to be more lenient than using normative data from the UK population as we see in our results.

Implications

This study adds to the field by providing a picture of MCI at various time points in a cohort whereby all participants were born in the same year and same country/region, and thus have had similar life experiences.\textsuperscript{14} Our research contributes to the global effort to understand how subclinical cognitive impairment affects older adults. However, this study also highlights the imprecision of MCI, with factors such as the normative data used, or the types of cognitive tests used, significantly affecting MCI rates. This has major practical implications for the use of MCI in a
clinical setting. Future research should establish a more precise definition of subclinical cognitive impairment with more consistency in measurement approaches. By doing so, research may be able to provide evidence leading to improved clinical tools.

The identification of individuals with MCI in the LBC1936 and their comparison with findings in similar cohorts provides opportunities for future research to further explore MCI in this cohort. In particular, utilizing the wealth of longitudinal data in the LBC1936 could prove insightful. MCI has been shown to be relatively fluid over time with both declines and reversions being common.34–36 Accordingly, understanding this fluidity and the predictive factors associated with MCI change will be insightful for future interventions and prevention strategies that aim to lower the risk of MCI developing and progressing. However, our results do deviate in some ways from previous research. This inconsistency is not uncommon in the field of MCI and dementia research and highlights the issue with using diagnostic criteria which leave room for interpretation without clear cut-offs or specified measures.37 Indeed, definitions of dementia have continued to evolve over the past decade, causing a lack on effect on how MCI can be identified.37 Accordingly, comparisons between studies need to be made with caution. Recent research has proposed that future research may benefit from exploring data driven computer algorithms for identifying MCI which may subsequently provide greater validity and enable data synthesis to be more accurate.37,38 Furthermore, some criticism aimed at measures of MCI suggest that it is a somewhat restrictive perspective of subclinical cognitive impairment.12,39 As previously mentioned, there are other ways to identify these subpopulations, and accordingly, future LBC1936 research may benefit from considering other less restrictive measures such as other cognitive impairment no dementia which does not rely on functional impairment as a factor.12

CONCLUSION

This study is largely consistent with previous research, finding MCI rates of 15% to 18% in the LBC1936 at ages 76 to 82. When considering subtypes of MCI, nonamnestic MCI is more likely to affect participants than aMCI indicating that perhaps this population is more prone to cognitive decline in nonamnestic cognitive domains. These results help highlight the prevalence of MCI in the LBC1936 and allow for future studies to explore cognitive trajectories over time and the predictive factors which may increase the risk of developing MCI.

REFERENCES

1. Buckinx F, Rolland Y, Regnier J-Y, et al. Burden of frailty in the elderly population: perspectives for a public health challenge. Arch Public Health. 2015;73:19–26.
2. Pankratz VS, Roberts RO, Mielke MM, et al. Predicting the risk of mild cognitive impairment in the Mayo Clinic Study of Aging. Neurology. 2015;84:1433–1442.
3. Robertson DA, Savva GM, Kenny RA. Frailty and cognitive impairment—a review of the evidence and causal mechanisms. Ageing Res Rev. 2013;12:840–851.
4. Lo RY. The borderland between normal aging and dementia. Tzu-Chi Med J. 2017;29:65–71.
5. Heinik J, VA Kral and the origins of benign senescent forgetfulness and mild cognitive impairment. Int Psychogeriatr. 2010;22:395–402.
6. Petersen RC, Doyle R, Kurz A, et al. Current concepts in mild cognitive impairment. Arch Neurol. 2001;58:1985–1992.
7. Dubois B, Albert ML. Amnestic MCI or prodromal Alzheimer’s disease. Lancet Neurol. 2004;3:246–248.
8. Winblad B, Palmer K, Kivipelto M, et al. Mild cognitive impairment—beyond controversies, towards a consensus: report of the International Working Group on Mild Cognitive Impairment. J Int Med. 2004;255:240–246.
9. Albert MS, DeKosky ST, Dickson D, et al. The diagnosis of mild cognitive impairment due to Alzheimer’s disease: recommendations from the National Institute on Aging-Alzheimer’s Association workgroups on diagnostic guidelines for Alzheimer’s disease. Alzheimers Dement. 2011;7:270–279.
10. Katz MJ, Lipton RB, Hall CB, et al. Age and sex specific prevalence and incidence of mild cognitive impairment, dementia and Alzheimer’s dementia in blacks and whites: a report from the Einstein Aging Study. Alzheimer Dis Assoc Disord. 2012;26:335–343.
11. Csukly G, Siraly E, Fodor Z, et al. The differentiation of amnestic type MCI from the non-amnestic types by structural MRI. Front Aging Neurosci. 2015;7:154.
12. Richardson C, Stephan BC, Robinson L, et al. Two-decade change in prevalence of cognitive impairment in the UK. Eur J Epidemiol. 2019;34:1085–1092.
13. Deary IJ, Gow AJ, Taylor MD, et al. The Lothian Birth Cohort 1936: a study to examine influences on cognitive ageing from age 11 to age 70 and beyond. BMC Geriatr. 2007;7:28–40.
14. Taylor AM, Pattie A, Deary IJ. Cohort profile update: the Lothian Birth Cohorts of 1921 and 1936. Int J Epidemiol. 2018;47:1042–1042r.
15. Deary IJ, Whalley LJ, Starr JM. A Lifetime of Intelligence: Follow-Up Studies of the Scottish Mental Surveys of 1932 and 1947. Washington, DC: American Psychological Association; 2009.
16. Deary IJ, Gow AJ, Pattie A, et al. Cohort profile: the Lothian Birth Cohorts of 1921 and 1936. Int J Epidemiol. 2012;41:1576–1584.
17. Townsend P. Poverty in the United Kingdom: A Survey of Household Resources and Standards of Living. Oakland, California: University of California Press; 1979.
18. Folstein MF, Folstein SE, McHugh PR. “Mini-mental state”: a practical method for grading the cognitive state of patients for the clinician. J Psychiatr Res. 1975;12:189–198.
19. Psychological Corporation. WAIS-III/WMS-III Technical Manual. San Diego, California: Harcourt Brace & Company; 1997.
20. Knopman DS, Gottesman RF, Sharrett AR, et al. Mild cognitive impairment and dementia prevalence: the atherosclerosis risk in communities neurocognitive study. Alzheimers Dement. 2016;2:1–11.
21. Fried LP, Tangen CM, Walston J, et al. Frailty in older adults: evidence for a phenotype. J Gerontol A: Biol Sci Med Sci. 2001;56:M146–M157.
22. Welstead M, Muniz-Terrera G, Russ TC, et al. Inflammation as a risk factor for the development of frailty in the Lothian Birth Cohort 1936. Exp Gerontol. 2020;139:111055.
23. Zigmond AS, Snaith RP. The hospital anxiety and depression scale. Acta Psychiatr Scand. 1983;67:361–370.
24. Education SCfRi. The Intelligence of Scottish Children: A National Survey of an Age-Group (Vol 5). London: University of London Press; 1933.
25. R Core Team. R: A Language and Environment for Statistical Computing. 2015.
26. Juarez-Cedillo T, Sanchez-Arenas R, Sanchez-Garcia S, et al. Prevalence of mild cognitive impairment and its subtypes in the Mexican population. Dement Geriatr Cogn Disord. 2012;34:271–281.
27. Hämmen T, Hallikainen M, Tuomainen S, et al. Prevalence of mild cognitive impairment: a population-based study in elderly subjects. Acta Neurol Scand. 2002;106:148–154.
28. Petersen RC, Roberts RO, Knopman DS, et al. Prevalence of mild cognitive impairment is higher in men: The Mayo Clinic Study of Aging. Neurology. 2010;75:889–897.
29. Xue J, Li J, Liang J, et al. The prevalence of mild cognitive impairment in China: a systematic review. *Aging Dis*. 2018;9:706–715.

30. Qian J, Wolters FJ, Beiser A, et al. APOE-related risk of mild cognitive impairment and dementia for prevention trials: an analysis of four cohorts. *PLoS Med*. 2017;14:e1002254.

31. Al-Qazzaz NK, Ali SH, Ahmad SA, et al. Cognitive impairment and memory dysfunction after a stroke diagnosis: a post-stroke memory assessment. *Neuropsychiatr Dis Treat*. 2014;10:1677–1691.

32. Petersen RC, Lopez O, Armstrong MJ, et al. Practice guideline update summary: mild cognitive impairment: Report of the Guideline Development, Dissemination, and Implementation Subcommittee of the American Academy of Neurology. *Neurology*. 2018;90:126–135.

33. Kriegsman DM, Penninx BW, Van Eijk JTM, et al. Self-reports and general practitioner information on the presence of chronic diseases in community dwelling elderly: a study on the accuracy of patients’ self-reports and on determinants of inaccuracy. *J Clin Epidemiol*. 1996;49:1407–1417.

34. Overton M, Philsgård M, Elmnåhl S. Diagnostic stability of mild cognitive impairment, and predictors of reversion to normal cognitive functioning. *Dement Geriatr Cogn Disord*. 2019;48:317–329.

35. Pandya SY, Clem MA, Silva LM, et al. Does mild cognitive impairment always lead to dementia? A review. *J Neurol Sci*. 2016;369:57–62.

36. Pandya SY, Lacritz LH, Weiner MF, et al. Predictors of reversion from mild cognitive impairment to normal cognition. *Dement Geriatr Cogn Disord*. 2017;43:204–214.

37. Grasset L, Matthews FE, Pérès K, et al. Evolution of dementia diagnosis over time (1988–2013): evidence from French and English cohorts. Implication for secular trends analyses. *Alzheimers Dement*. 2018;10:490–497.

38. Matthews FE, Stephan B, Robinson L, et al. A two decade dementia incidence comparison from the Cognitive Function and Ageing Studies I and II. *Nat Commun*. 2016;7:1405–1412.

39. Ritchie K, Artero S, Touchon J. Classification criteria for mild cognitive impairment: a population-based validation study. *Neurology*. 2001;56:37–42.