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## Pre-statistical Considerations for Harmonization of Cognitive Instruments: Harmonization of ARIC, CARDIA, CHS, FHS, MESA, and NOMAS

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## Abstract

**BACKGROUND:** Meta-analyses of individuals' cognitive data are increasing to investigate the biomedical, lifestyle, and sociocultural factors that influence cognitive decline and dementia risk. Pre-statistical harmonization of cognitive instruments is a critical methodological step for accurate cognitive data harmonization, yet specific approaches for this process are unclear.

**OBJECTIVE:** To describe pre-statistical harmonization of cognitive instruments for an individual-level meta-analysis in the blood pressure and cognition (BP COG) study.

**METHODS:** We identified cognitive instruments from six cohorts (the Atherosclerosis Risk in Communities Study, Cardiovascular Health Study, Coronary Artery Risk Development in Young Adults study, Framingham Offspring Study, Multi-Ethnic Study of Atherosclerosis, and Northern Manhattan Study) and conducted an extensive review of each item's administration and scoring procedures, and score distributions.

**RESULTS:** We included 153 cognitive instrument items from 34 instruments across the six cohorts. Of these items, 42% were common across 2 cohorts. 86% of common items showed differences across cohorts. We found administration, scoring, and coding differences for seemingly equivalent items. These differences corresponded to variability across cohorts in score distributions and ranges. We performed data augmentation to adjust for differences.

**CONCLUSION:** Cross-cohort administration, scoring, and procedural differences for cognitive instruments are frequent and need to be assessed to address potential impact on meta-analyses and cognitive data interpretation. Detecting and accounting for these differences is critical for accurate attributions of cognitive health across cohort studies.

## Keywords

cognition; dementia; epidemiology; methods

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## Introduction

Researchers are increasingly performing meta-analyses of individual participant cognitive data from existing large databases and cohorts to better understand cognitive decline and dementia risk [1,2]. Pooling cognitive data from longitudinal population-based cohorts facilitates improved power and novel capabilities to investigate biomedical, lifestyle, and sociocultural factors that may affect cognition across the lifespan [3,4]. Prior to pooling cognitive data, pre-statistical harmonization is necessary to ensure accurate and consistent inferences about cognitive health across studies.

Pre-statistical harmonization [5] of cognitive instruments is a complicated, qualitative process that involves a careful review of cohort characteristics (e.g., subject selection procedures, demographic factors such as race/ethnicity, socioeconomic status, language), and cognitive instruments to identify common and unique items across datasets, and candidate items that might be made comparable with minimal transformation. Specific procedures for pre-statistical harmonization of cognitive instruments across studies are not established [6].

Pre-statistical harmonization of cognitive instruments is a unique challenge because heterogeneity exists in the instruments used to measure cognition, and the procedures for administering them. Over 500 instruments are available for clinical and research use [7], and they differ by domain of assessment, level of measurement precision (e.g., ranging from a 10-minute, in-person or telephone-based cognitive screening assessment to a several-hour battery of neuropsychological tests), sensitivity to change, and other factors. Heterogeneity may also be present across seemingly parallel cognitive instruments, such as differences in test version, test adaptation for individual study purposes, and differences in administration, scoring, and coding procedures. These procedural details are crucial to uncover prior to the harmonization process. Data augmentation procedures are often required to address this variability across studies to create one cohesive meta-analysis dataset. This aspect of pre-statistical harmonization is rarely discussed or documented in harmonization research [6].

To promote reproducible research in cognitive decline and dementia risk, in this report we describe pre-statistical harmonization for meta-analysis of individual participant data from six American population-based cohorts for the Effect of Lower Blood Pressure over the Life Course on Late-life Cognition (BP-COG) study, quantifying the effect of blood pressure (BP) levels over the life course on cognitive trajectories in Black, Hispanic, and White individuals [1,2]. We aim to describe our approach to pre-statistical harmonization of cognitive data, including our detailed review of the administration, scoring procedures, and score ranges of cognitive instruments across cohorts to determine their degree of equivalence and suitability for pooling. Second, we summarize our findings from this detailed review and their implications for data interpretation. Finally, we offer recommendations for

pre-statistical harmonization procedures for cognitive instruments to inform future meta-analyses.

## Methods

### Cohort studies

Six NIH-funded, longitudinal cohort studies were included in the present study: The Atherosclerosis Risk in Communities Study (ARIC [8]), the Coronary Artery Risk Development in Young Adults Study (CARDIA;[9]), the Cardiovascular Health Study (CHS; [10]), the Framingham Offspring Study (FOS; [11]), the Multi-Ethnic Study of Atherosclerosis (MESA; [12]), the Northern Manhattan Study (NOMAS; [13]). All participants provided informed consent. A description of each cohort study and the rationale for its inclusion in the larger study is available [1]. We selected these cohorts because all had repeated measures of cognition and BP using similar methods, high-quality data on dementia risk factors, overlapping cognitive instruments, physician-adjudicated dementia diagnosis, availability of brain magnetic resonance imaging, racial, ethnic, and geographic diversity, and included adults aged 18 and older. Four cohorts (NOMAS, ARIC, MESA, CARDIA) enrolled adults in early adulthood to mid-life, which is the time in the life course with the most consistent association between BP and later-life cognitive decline. The other two cohorts enrolled beginning in early life (FOS) and older adulthood (CHS).

### Procedure

First, study neuropsychologists (EB, BG) identified cognitive instruments in each of the six cohort studies from documentation provided by each cohort, available published papers, and scrutiny of cognitive datasets. We considered all cognitive instruments at all study visits for inclusion in the harmonization, with the exception of measures of literacy and premorbid intellectual functioning that are relatively less sensitive to age-related cognitive decline.

After identifying available cognitive instruments, we contacted cohort study investigators to request unpublished administration, scoring, and procedural details of cognitive test batteries. Documentation provided by each cohort study included test forms, data entry forms, and administration and scoring instructions. Procedural details extracted from this process included the published test version, administration and scoring details (e.g., stopping rules; acceptable responses for specific items), possible/theoretical score ranges (based upon the instrument structure and number of items), and metrics available for each instrument (e.g., individual item data, raw and standardized summary scores). We reviewed available raw data for each instrument for score ranges and distributions.

When available and appropriate, we reviewed individual test items comprising cognitive instruments rather than composite/summary test scores (e.g., individual test items for the Mini Mental State Examination [MMSE]; Telephone Interview for Cognitive Status [TICS], Modified Mini Mental State Examination [3MSE], or Montreal Cognitive Assessment [MoCA]). Decomposition of these instruments into their component items enabled us to identify additional common items between cohorts and to link telephone-administered test batteries with in-person test batteries.

Study neuropsychologists reviewed and categorized each cognitive instrument item into relevant cognitive domains (i.e., memory, executive functioning).

### **Comparability of items across cohort and language**

We identified cognitive test items that were comparable across cohorts based upon detailed documentation. For cohorts that offered test administration in Spanish, neuropsychologists considered the degree of linguistic and cultural equivalence based on the evidence base for individual cognitive instrument items. Items for which there was concern for non-equivalence across English and Spanish were considered separate items by language rather than comparable items.

### **Data augmentation**

We employed several data augmentation strategies to adjust for differences uncovered during the pre-statistical harmonization process, including alignment of coding procedures, winsorization, and equipercentile equating. The decision of how to address each disparate test item was made on a case-by-case basis by an experienced epidemiologist in consultation with study neuropsychologists, based on the distribution of scores across tests and assuming that we properly identified test items that were comparable across cohorts, except for version or administration differences. To make these decisions, we meticulously scrutinized exploratory analyses of test scores (e.g., dotplots and histograms, stem and leaf plots, tables of minima, maxima, medians, and means) by cohort for every test item presumed to be comparable across cohorts. When procedural or distributional differences across cohorts in common test items were identified, we reviewed each item for possible data augmentation.

## **Results**

### **Summary of cognitive instrument items**

We identified 34 cognitive instruments, 13 of which were common across two or more cohorts. Animal naming spanned all six cohorts, letter fluency spanned five cohorts, and four instruments (Mini Mental State Examination, Boston naming, digit span, and digit symbol substitution) spanned four cohorts (Table 1). From these 34 cognitive instruments, we identified 153 items (Supplemental Tables 1 and 2), 64 of which were common across at least two cohorts. A detailed review of instrument administration and scoring procedures across cohorts revealed several sources of procedural differences across cohorts, despite seemingly common tests (Supplemental Tables 1 and 2). Frequent sources of such heterogeneity included differences in instrument version, instrument adaptation, administration procedures, data (i.e., type of score provided for a cognitive instrument), and component items.

### **Instrument Version**

Differences in instrument version affected 10 of 13 common instruments. For example, four studies (ARIC, CHS, CARDIA, MESA) implemented a speeded task requiring transcription of symbols that corresponded to specific digits, titled Digit Symbol Substitution Test for the Wechsler Adult Intelligence Scale-Revised (WAIS-R, [14]) published in 1981. Two studies (MESA and CARDIA) used an updated version, the Digit Symbol Coding Test, published

in the test's 1997 revision (Wechsler Adult Intelligence Scale- III; WAIS-III, [15]). Although these tests are nearly equivalent in structure, they had different times to complete (90 sec for WAIS-R vs. 120 sec for WAIS-III) and a different number of possible items (93 items for WAIS-R vs. 133 items for WAIS-III). Of note, a similarly-titled, but distinct, measure was administered in the NOMAS study (Symbol Digit Modalities Test; SDMT [16]), which allowed 90 sec to transcribe numbers corresponding to distinct symbols. Scores on this test tend to be lower than for the digit symbol versions, possibly due to subtle differences in task demands[17]. For each of these tests, performance is quantified as the number of correct items completed; as such, an equivalent raw score is associated with a different level of performance (i.e., speed) across these studies. For example, 50 items completed with a 90 sec time limit corresponds to 0.56 items/sec, whereas 50 items completed within a 120 sec time limit corresponds to 0.42 items/sec.

### **Instrument adaptation**

Adaptations from standard administration or scoring of tests impacted 8 of 13 common instruments. For example, the California Verbal Learning Test (CVLT) was administered in two cohorts (NOMAS, CHS Cognition Study). The CHS Cognition Study used the standard CVLT-1 administration procedures, with a list of 16 words and instructions to recall items on a shopping list. The NOMAS study, in contrast, used a list of 12 rather than 16 words; the words were presented via audio recording with a male or female voice, and respondents were required to recall the words and whether the word was presented in a male or female voice.

### **Administration procedures**

Differences in administration procedures, such as stopping rules for timed tests, were noted in 6 of 13 common instruments. Some administration differences led to different raw score ranges across common tests. For example, the Trail Making Test, Part B had a maximum raw score of 240 sec in the ARIC study due to their discontinuation rule of 240 sec and/or 5 errors. This same test had a maximum raw score of 572 sec in the CHS Cognition study and 600 sec in the FOS study, for which stopping rules were not available in study documentation. Studies also varied with procedures for the items evaluating working memory (e.g., serial subtractions or spelling backwards) on the MMSE: NOMAS administered serial subtraction only when full credit was not awarded for spelling backwards; CHS studies administered both items and scored the higher of the two responses, whereas ARIC and NOMAS administered only the spelling backwards item.

### **Score type**

Studies varied in the selection of summary scores provided for common instruments, and most instruments had several summary scores included in datasets. For 1 of 13 common instruments, there were no equivalent summary scores available for common instruments across studies. Specifically, for the digit span test, the FOS study provided the maximum string of digits correctly answered, whereas ARIC and MESA provided the total score for the measure, and CHS provided both metrics.

## Scoring and coding procedures

Review of item scoring and coding procedures revealed several sources of heterogeneity across cohorts and affected items in 2 of 13 common instruments (MMSE and TICS), in addition to common items identified from other instruments (3MSE, CASI, MoCA). Studies varied with regard to the degree of precision required by the respondent to award credit for an item. For example, for the orientation to season item, NOMAS awarded credit if respondents made an error during the first or last day of the month, whereas CHS, FOS, and ARIC required precise responses. Common items were also identified between the MMSE and other cognitive screening instruments (i.e., 3MSE, Cognitive Abilities Screening Instrument (CASI), MoCA). Across these measures, differences with regard to scoring and coding were noted frequently across equivalent items. For example, when orientation to month was administered in the CASI (MESA), coding was as follows: 2= accurate within 5 days; 1= missed by 1 month, and 0 = missed by 2+ months; whereas coding of this item for MMSE and TICS (ARIC, CHS, NOMAS) was 1=correct, 0=incorrect (Supplemental Table 2).

## Cultural and linguistic equivalence

We reviewed candidate comparable items that were administered across English and Spanish. We noted evidence of differential difficulty across language for some test items. For example, items from the Boston Naming Test have shown differential difficulty across English and Spanish [18]. Because of concern for possible non-equivalence in data interpretation across English and Spanish, we considered these test items as separate rather than comparable items.

## Data augmentation

Supplemental Tables 1 and 2 describe the data augmentation strategies used for each item included in the harmonization. Of the 64 comparable items identified across cohorts, 55 required data augmentation. For some instruments, such as the letter fluency test and the Trail Making Test, we transformed scores to curtail outliers via winsorization (i.e., pulling in extreme values) (Supplemental Table 1). We coarsened more finely measured test scores in some cohorts in order to map them to other available test items in other cohorts (e.g., creating summary scores comprised of individual test items); for example, this was done for test items such as delayed word recall for 3MSE from the CHS study (Supplemental Table 2). For tests with completion time scores, such that higher scores are indicative of worse performance (e.g., Trail Making Test, Stroop Color and Word Test, Grooved Pegboard Test), we adjusted the direction of the scores to reflect higher scores indicative of better performance (Supplemental Table 1). We used equipercentile equating [19] to adjust the distributions of like test items across cohorts to be on a common scale for tests such as the CVLT and Digit Symbol Substitution Test/Digit Symbol Coding Test (Supplemental Table 1). To address differences in coding procedures for correct items (e.g., CASI items, MESA cohort), we recoded items to align across studies (Supplemental Table 2). To address differences in age distributions across cohorts, the age range was restricted to 53–80 years to develop the equating algorithm. The equating algorithm was then applied to the full sample.



To illustrate the impact of the sources of heterogeneity on cognitive data and data augmentation procedures, we selected data from the DSST/DSC instrument (Figure 1). As illustrated in Figure 1 (panel B), instrument test version differences were associated with markedly different raw score distributions across cohorts. To address these differences in raw scores (and given equivalence of construct assessed), we used equipercentile equating to align distributions (Figure 1, Panel C).

## Discussion

In the pre-statistical harmonization of cognitive instruments for meta-analyses of individual participant data from six longitudinal cohort studies, we found numerous differences in cognitive instrument version, administration, scoring, coding, and other procedures across cohorts. We found differences even in seemingly comparable cognitive instruments across cohorts that made them potentially nonequivalent. We found procedural differences even for widely used, standardized cognitive tests, such as the MMSE. Many of these sources of cognitive instrument variability were associated with clear differences in cognitive data distributions across cohorts. Of the 64 comparable items identified across cohorts, 55 (86%) required data augmentation to facilitate comparability across cohorts.

Although studies have examined statistical harmonization of cognitive data across studies [20–22], no studies have examined pre-statistical harmonization of cognitive data across studies[6]. Our results provide evidence that careful pre-statistical harmonization, including detecting and accounting for cognitive instrument procedural differences across cohorts, is a critical analytic step of the harmonization process. Assuming equivalence could lead to misattribution of procedural sources of variance to systematic differences between groups of people across cohorts. In addition, transparency in the pre-statistical harmonization process is critical for reproducible research and to ensure scientific scrutiny of this aspect of harmonization methodology.

The present study highlights the critical need for performing specific, detailed and comprehensive pre-statistical harmonization steps prior to statistical harmonization, as we summarized in a checklist (Table 2). These procedures are warranted for all approaches to cognitive data harmonization. At each level of pre-statistical harmonization, expert content reviewers (e.g., neuropsychologists) are needed to ensure that decisions made about the equivalence and utility of test items are appropriate. This expert content review process is particularly necessary prior to harmonization of cognitive instruments across populations that have diverse cultural and linguistic characteristics, as expert reviewers may identify cultural and linguistic sources of non-equivalence across items that could lead to misattributions of cognitive test score differences if not detected.

Our findings have important implications for future studies that aim to harmonize data from cognitive instruments. We summarize our recommended procedures for pre-statistical harmonization of cognitive instruments in Table 2 and Figure 2. We outline pre-statistical harmonization steps, critical information to document, relevant examples from the BP COG study, and offer possible solutions when differences are uncovered. We offer the following conclusions and recommendations for future cognitive data

harmonization studies based upon this work. First, use of parallel, standardized cognitive tests does not guarantee identical implementation of these tests across studies [23]; we recommend careful scrutiny and documentation of procedural differences prior to data pooling. This process is resource-intensive, should involve a content expert in neuropsychological assessment, and should be budgeted for when planning studies requiring cognitive instrument harmonization. Availability of source documents pertaining to cognitive instrument administration and scoring procedures are critical for comprehensive and thorough pre-statistical harmonization. We only know what we know: undocumented deviations in standard administration are difficult to detect post-hoc for investigators who were not involved during original data collection. Second, heterogeneity in implementation of common cognitive instruments can lead to important differences in score distributions across studies. Failure to detect and account for these differences during pre-statistical harmonization could lead to erroneous attributions regarding cognitive health across cohort studies. Finally, detecting these differences facilitates use of statistical procedures, in IRT-based models, to account for these differences and allow for equating scores.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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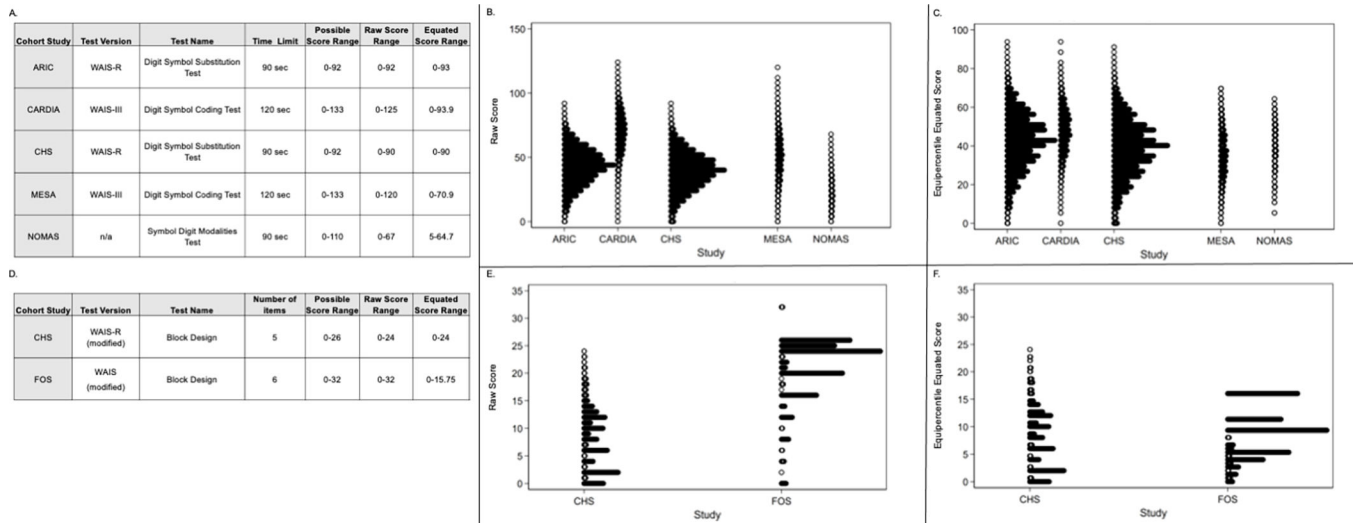
#### Conflicts of Interest/Disclosure Statement

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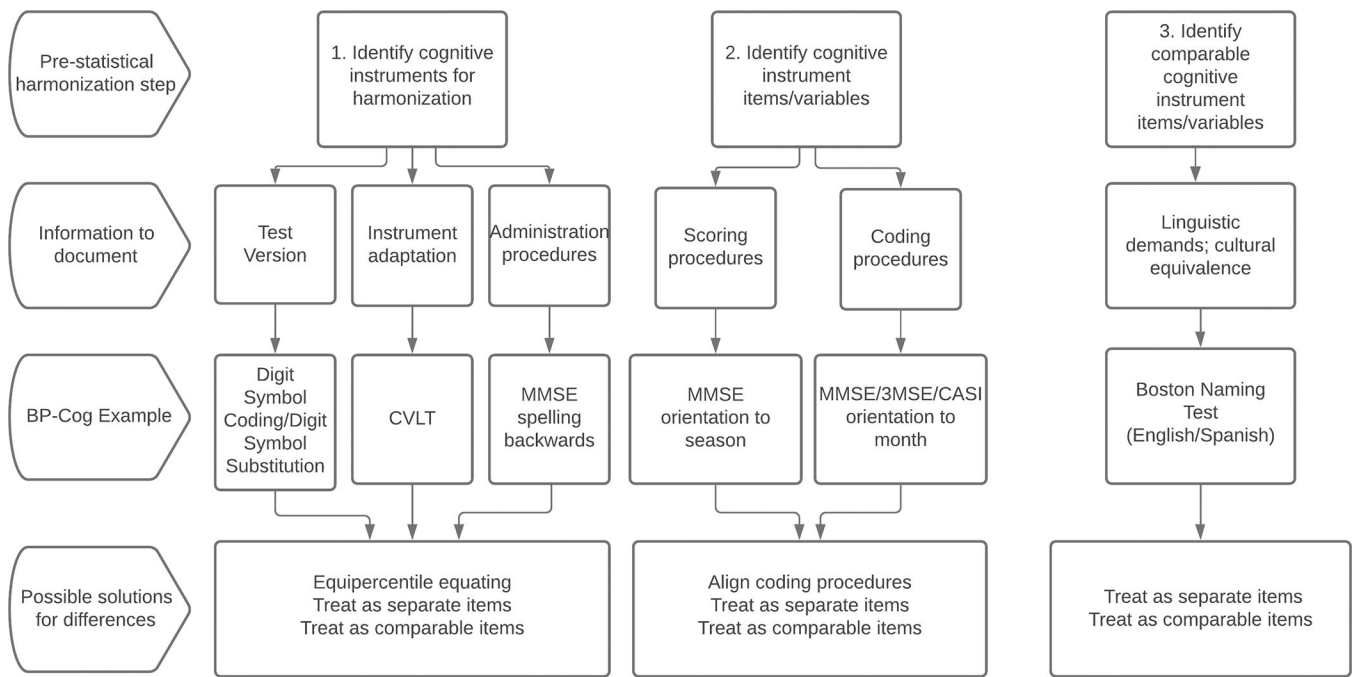
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**Figure 1.** Impact of procedural differences on digit symbol substitution test data across cohorts. Panel A summarizes differences in test version, procedure, and raw score ranges across cohorts for the digit symbol substitution test. Panel B displays the raw score distributions across cohorts for the digit symbol substitution test. Panel C displays the equipercentile-equated distributions for the digit symbol substitution test. All scores are scaled on a T score metric (mean = 50, standard deviation = 10).



**Figure 2.** Summary of recommended pre-statistical steps for harmonization of cognitive instruments. Figure outlines recommended pre-statistical procedures for cognitive instrument harmonization, examples of sources of heterogeneity across cohorts, and possible solutions for addressing sources of heterogeneity.

**Table 1.**

Common and unique cognitive instruments administered in each cohort study

	Domain	ARIC	CARDIA	CHS	FOS	MESA	NOMAS	
Global cognitive performance (GCP)	Executive function/ processing speed	DSST	DSC	DSST		DSC		
							SDMT	
		Animal Naming	Animal Naming	Animal Naming	Animal Naming	Animal Naming	Animal Naming	Animal Naming
				Semantic Gen.				
		Letter Fluency	Letter Fluency	Letter Fluency	Letter Fluency	Letter Fluency		Letter Fluency
		Trail Making		Trail Making	Trail Making			
			Stroop Test	Stroop NST				
								Color Trails; Odd Man Out
				Baddeley-Papagno				
						WAIS Similarities		
			WAIS-R Digit Span		WAIS-R Digit Span	WAIS Digit Span	WAIS-III Digit Span	
							Digit ordering; WAIS-III LNS	
		Learning/Memory		RAVLT	CVLT			CVLT-II
			DWR					
						WMS Paired Assoc.		
			WMS-R LM			WMS LM		
					Rey O CFT Recall			
						WMS-R VR		
		General Mental Status	MMSE		MMSE	MMSE		MMSE
				MoCA				
			TICS		TICS			TICS
					3MSE			
							CASI	
		Language	Boston Naming (30-item)		Boston Naming (30-item)	Boston Naming (30-item)		Boston Naming (15-item)
		Motor			Grooved Pegboard			Grooved Pegboard
						Finger Tapping		

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Visuospatial	Clock Time Perception					
			WAIS-R Block Design	WAIS Block Design		
			Ravens			
			Rey O CFT Copy			
				Hooper VOT		
						VMI

*Note:* Cognitive instruments were administered to different participant samples and at different time intervals across the 6 cohorts. Instruments identified as candidate comparable items are listed in the same row.

Abbreviations:

Cohorts: CARDIA is the Coronary Artery Risk Development in Young Adults study. NOMAS is the Northern Manhattan Study. ARIC is the Atherosclerosis Risk in Communities Study. CHS is the Cardiovascular Health Study. MESA is the Multi-Ethnic Study of Atherosclerosis study. FOS is the Framingham Offspring Study.

Cognitive Instruments: 3MSE is the Modified Mini Mental State Examination. ANT is Animal Naming Test. BNT is Boston Naming Test. CASI is Cognitive Abilities Screening Instrument. CVLT is California Verbal Learning Test. DSC is WAIS-III Digit Symbol Coding Test. DSST is the WAIS-R Digit Symbol Substitution Test. DWR is the Delayed Word Recall test. Hooper VOT is Hooper Visual Organization Test. LFT is Letter Fluency Test. LM is Logical Memory. LNS is Letter-Number Sequencing. MMSE is Mini-Mental State Examination. MoCA is Montreal Cognitive Assessment. RAVLT is Rey Auditory Verbal Learning Test (RAVLT). Rey O CFT is Rey-Osterrieth Complex Figure Test. SDMT is Symbol Digit Modalities Test. Semantic Gen. is semantic word generation (birds, dogs). Similarities is the WAIS Similarities test. Stroop NST is the Stroop Neuropsychological Screening Test. TICS is the Telephone Interview for Cognitive Status. VMI is Developmental Test of Visuomotor Integration. VR is Visual Reproduction. WAIS is Wechsler Adult Intelligence Scale. WAIS-R is Wechsler Adult Intelligence Scale-Revised. WAIS-III is Wechsler Adult Intelligence Scale-3rd Edition. WMS is Wechsler Memory Scale. WMS-R is WMS-Revised. WMS-III is WMS-3rd Edition.

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**Table 2.**

## Checklist of procedures for pre-statistical harmonization of cognitive data

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Define and state the study question
Identify cohorts based on the study question
Identify cognitive instruments within each cohort
Collect all available documentation about cognitive instruments
Test versions
Test adaptations
Administration procedures
Stopping rules for timed tests
Items administered conditional on other items
Scoring and coding procedures
Theoretical and empirical score ranges
Language of administration
Cultural and linguistic comparability
Mode of administration (phone vs in-person vs internet)
Consider test items from cognitive instruments, rather than summary scores, when available and reasonable
Expert neuropsychologists to categorize cognitive test items into domains of interest, informed by the study question
Identify items common across cohorts based on available documentation
Consider data transformation strategies to derive common items across cohorts if items are conceptually comparable but different in distribution or with different response values
Exploratory data analysis
Winsorization
equipercentile equating

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