Effects of Sex, APOE4, and Lifestyle Activities on Cognitive Reserve in Older Adults

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Abstract

Background and Objectives
Lifestyle activities, such as physical activity and cognitive stimulation, may mitigate age-associated cognitive decline, delay dementia onset, and increase cognitive reserve. Whether the association between lifestyle activities and cognitive reserve differs by sex and APOE4 status is an understudied yet critical component for informing targeted prevention strategies. The current study examined interactions between sex and physical or cognitive activities on cognitive reserve for speed and memory in older adults.

Methods
Research participants with unimpaired cognition, mild cognitive impairment, or dementia from the Washington Heights-Inwood Columbia Aging Cohort were included in this study. Cognitive reserve scores for speed and memory were calculated by regressing out hippocampal volume, total gray matter volume, and white matter hyperintensity volume from composite cognitive scores for speed and memory, respectively. Self-reported physical activity was assessed using the Godin Leisure Time Exercise Questionnaire, converted to metabolic equivalents (METS). Self-reported cognitive activity (COGACT) was calculated as the sum of 3 yes/no questions. Sex by activity interactions and sex-stratified analyses were conducted using multivariable linear regression models, including a secondary analysis with APOE4 as a moderating factor.

Results
Seven hundred fifty-eight participants (mean age = 76.11 ± 6.31 years, 62% women) were included in this study. Higher METS was associated with greater speed reserve in women (β = 0.04, CI 0.00–0.08) but not in men (β = 0.004, CI −0.04 to 0.05). METS was not associated with memory reserve in women or men. More COGACT was associated with greater speed reserve in the cohort (β = 0.13, CI 0.05–0.21). More COGACT had a trend for greater memory reserve in women (β = 0.06, CI −0.02 to 0.14) but not in men (β = −0.04, CI −0.16 to 0.08). Only among women, APOE4 carrier status attenuated relationships between METS and speed reserve (β = −0.09, CI −0.22 to 0.04) and between COGACT and both speed (β = −0.26, CI −0.63 to 0.11) and memory reserves (β = −0.20, CI −0.50 to 0.093).

Discussion
The associations of self-reported physical and cognitive activities with cognitive reserve are more pronounced in women, although APOE4 attenuates these associations. Future studies are needed to understand the causal relationship among sex, lifestyle activities, and genetic factors on cognitive reserve in older adults to best understand which lifestyle activities may be most beneficial and for whom.

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With a scarcity of disease-modifying treatments for Alzheimer disease (AD), prevention is critical. Enhancing cognitive reserve, operationalized as preserved cognitive health despite the presence of brain pathology,1,2 is a promising path toward dementia prevention. Lifestyle factors such as physical activity and cognitive stimulation are 2 modifiable behaviors that may enhance cognitive reserve and reduce dementia risk,3–8 yet it is unclear who benefits most from these activities. Sex differences associated with cognitive benefits of physical activity are evident but mixed. Although meta-analytic reviews aggregating retrospective data suggest a stronger relationship between exercise and cognition in women than men,9,10 recent prospective studies, including an exercise randomized controlled trial, show larger cognitive gains following exercise in men than in women.11 Even less is known about sex differences on the benefit of cognitive stimulation on cognitive reserve.5 Factors such as sex, APOE4, the major genetic risk factor for late-onset AD, and their interaction importantly affect cognitive trajectories, yet little is known on whether they moderate the beneficial effects of modifiable lifestyle activities on cognitive reserve. The current study investigated whether associations of physical and cognitive activities with cognitive reserve are modified by sex and APOE4 carrier status in multiethnic, community-dwelling older adults. This approach will enable a better understanding of which modifiable factors may influence cognitive reserve and in whom.

**Methods**

**Participants**

Study participants were selected from the Washington Heights/Inwood Columbia Aging Project (WHICAP). WHICAP recruitment was conducted in 3 waves, starting in 1992, 1999, and 2009. Participants were English or Spanish speaking, Medicare eligible, racially and ethnically diverse residents of Northern Manhattan.12 The WHICAP inclusion criteria included individuals aged 65 years and older and language competency in English or Spanish. Exclusion criteria included dementia diagnosis for the 1999 and 2009 waves only.12,13 No other health-related exclusions were included to maximize population representativeness. Participants completed assessments for health status, functional ability, neurologic status, and neuropsychological test performance.14 A subset of participants received high-resolution structural T1 MRI brain scans at 1.5T or 3T MRI field strength.15 Their first imaging visit was selected for the analysis. Participants met eligibility for inclusion in this study if they had the following available data no more than 12 months apart (when applicable): (1) MRI data, (2) lifestyle activities data, (3) neuropsychological data, (4) APOE genotype, and (5) demographic information including age at the time of scan, sex, education, and diagnosis. Because of differing sex physiologies and systematic social norms that cannot be differentiated in our current study design, the terms women and men used in the present study refer to sex differences that may have biological, physiologic, or social etiologies.

Diagnoses were assigned through diagnostic consensus conferences attended by a panel of neurologists, psychiatrists, and neuropsychologists using a combination of neuropsychological, functional, and neurologic assessments.13,16 Dementia diagnoses also included use of the DSM-III criteria for dementia. The flowchart shows the final sample size of 758 participants (eFigure 1, links.lww.com/WNL/C103).

**Standard Protocol Approvals, Registrations, and Patient Consents**

The project design and protocol were approved by the Institutional Review Boards of the Columbia Presbyterian Medical Center and the New York State Psychiatric Institute. Participants provided written informed consent at the enrollment.

**MRI**

1.5T and 3T MRI scans were collected at Columbia University Medical Center with the following parameters: 1.5T Phillips Intera MRI scanner (T1-weighted structural MRI: repetition time [TR] 20 ms, echo time [TE] 2.1 ms, FOV 240 × 256 × 160 matrix, and 1.3 mm slice thickness; T2-FLAIR: TR 11,000 ms, TE 144 ms, inversion time = 2,800, FOV 25 cm × 256 × 192 matrix, and 3 mm slice thickness; proton density: TR 2675 ms, TE 12 ms, FOV 220 × 165 × 140, and 4 mm slice thickness) and 3T Philips MRI scanner (T1-weighted structural MRI: TR 6.6 ms, TE 3 ms, FOV 256 × 256 × 165, and 1 mm slice thickness; T2-FLAIR: TR 8.0 seconds, TE 332 ms, FOV 240 × 240 × 180, and 0.43 mm slice thickness). Hippocampal volumes, total gray matter volumes, and white matter hyperintensity (WMH) volumes were calculated using FreeSurfer and in-house processing pipeline as previously described.15,17 Hippocampal volumes and total gray matter volume were regressed against total intracranial volume. WMHs were log transformed to normalize their distribution. Scanner field strength was covaried for in the analysis using the recruitment cohort categorical variable, as described below.
APOE Genotyping
APOE genotype was determined using restriction isotyping. Participants with at least one E4 allele were classified as APOE4 carriers. All other participants were classified as APOE4 noncarriers. Analyses were repeated excluding APOE E2/E2 (n = 7) and E2/E4 genotype (n = 20).

Physical and Cognitive Activity Measures
Physical activity measures of duration, intensity, and frequency were collected using the Godin leisure time exercise questionnaire. Participants’ typical weekly physical activity was evaluated in terms of metabolic equivalent of task (METS) and was constructed as number of minutes × number of times × coefficient (9 for vigorous, 5 for moderate, and 3 for light activities corresponding to the metabolic equivalent) based on previously published work. A natural logarithm of physical activity was taken, and participants with no physical activity were bottom coded and assigned the lower limit of the score distribution. Cognitive activities (COGACT) were defined as an aggregate score based on self-reported participation (yes/no) in the following activities in the preceding 13 months: reading magazines, newspapers, or books; going to classes; and playing cards, games, or bingo. These measures were collected within 12 months of the MRI scan date.

Domain-Specific Cognitive Reserve Measures
In previously published work, confirmatory factor analysis identified the Selective Reminding Test (total recall, immediate recall, and delayed recognition) and Color Trails Test A and B as domain-specific scores for memory and speed domains, respectively. We used a residual approach to quantify domain-specific cognitive reserve. Memory and speed factor scores were regressed on hippocampal volume, total gray matter volume, and WMH volume. Sex was not included as a factor in the regression models. The resulting residuals from these regressions were used as proxies to represent their respective domain-specific cognitive reserve according to prior approaches to quantify cognitive reserve. Higher residual values indicate better speed or memory function.

Statistical Analyses
Analyses were conducted in SPSS Statistics 25 (IBM Corp., Released 2017; IBM SPSS Statistics for Windows, Version 25.0; Armonk, NY: IBM Corp.) and R. Sex differences of participants’ demographic variables were compared using t tests for continuous variables and χ² tests for categorical variables. Analysis of variance tests with interaction terms for physical or cognitive activities by sex were conducted to assess group differences in speed and memory reserves independently. Main effects were considered significant if type I error rate (α) was less than or equal to 0.05. Power analysis for one-way Analysis of variance F tests was used to determine significance levels for interactions. Interactions with p < 0.20 significance were subsequently probed with sex-stratified analyses (power = 0.6). Further analyses explored the moderating role of APOE4 in activity-reserve relationships by testing activity by sex by APOE4 3-way interactions. Activities by APOE4 interactions with a corresponding p < 0.05 were probed with APOE4-stratified analyses. Covariates for the analyses included age, sex (when applicable), race/ethnicity, education, recruitment cohort, diagnosis (cognitively unimpaired, MCI, or dementia), and APOE4 status (when applicable). APOE4 noncarriers, men, non-Hispanic White participants, and dementia diagnosis were used as the referent groups in the analyses. Partial R²-s were used to calculate the coefficient of determination of METS and COGACT.

Sensitivity analyses assessed whether imaging subsamples from wave 2 and wave 3 should be combined or considered separately by adjusting the reserve measures to reflect differences in association patterns and conducting the Student t test. In addition, participants were grouped by cognitive impairment status (cognitively normal [CN] vs cognitively impaired [MCI/AD]), and the modifying effect of the group on activity—domain reserve relationship was tested to assess whether the relationships were driven by group. The modifying effect of race/ethnicity was also evaluated in the sensitivity analyses.

Data Availability
Data can be requested and approved from the WHICAP Publication Committee (reference): cumc.co1..qualtrics.com/jfe/form/SV_6x5rRy14B6vpoqN.

Results
Demographic variables, physical and COGACT measures, and domain reserve scores in the cohort are summarized in Table 1. Men and women did not differ significantly on education, APOE4 carrier status, race/ethnicity, diagnosis, self-reported cognitive activities, and speed reserve. Women participants were older, reported less physical activity, and had larger memory reserve. For cognitive activities, women and men had similar reading and card-playing habits, but more women than men attended classes.

Physical Activity (METS) vs Speed and Memory Reserves
METS was associated with speed reserve among all participants (β = 0.05, 95% Confidence Interval [CI] 0.01–0.08, p = 0.01, model R² = 0.33, METS R² = 0.05); however, a METS by sex interaction (β = −0.05, CI −0.1 to 0.08, p = 0.09, interaction R² = 0.04) indicated the relationship differed by sex. Sex-stratified analyses revealed a significant relationship among women (β = 0.04, CI 0–0.08, p = 0.05, model R² = 0.36, METS R² = 0.08) that was not observed in men (β = 0.004, CI −0.04 to 0.05, p = 0.85, model R² = 0.26, METS R² < 0.01) (see Figure 1A, eTable 1, links.lww.com/WNL/C103, for model estimates).

METS was not associated with memory reserve (β = 0.004, CI −0.04 to 0.05, p = 0.82, model R² = 0.32, METS R² = 0.02), and
A METS by sex interaction was not observed ($\beta = -0.01$, CI $-0.06$ to $0.03$, $p = 0.60$, interaction $R^2 < 0.01$) (Figure 1B, eTable 1, links.lww.com/WNL/C103).

**COGACT vs Speed and Memory Reserves**

COGACT was associated with speed reserve among all participants ($\beta = 0.13$, CI $0.05$–$0.21$, $p = 0.003$, model $R^2 = 0.33$, COGACT $R^2 < 0.01$), and COGACT by sex interaction was not observed ($\beta = -0.01$, CI $-0.17$ to $0.16$, $p = 0.93$, interaction $R^2 < 0.01$) (Figure 1C, eTable 2, links.lww.com/WNL/C103).

COGACT was not associated with memory reserve among all participants ($\beta = 0.04$, CI $-0.03$ to $0.11$), $p = 0.26$, model $R^2 = 0.33$, COGACT $R^2 = 0.07$); however, a COGACT by sex interaction was observed ($\beta = -0.11$, CI $-0.24$ to $0.02$, $p = 0.11$, interaction $R^2 = 0.01$), revealing a trend among women ($\beta = 0.06$, CI $-0.02$ to $0.14$, $p = 0.15$, model $R^2 = 0.37$, COGACT $R^2 = 0.07$) and no association among men ($\beta = -0.04$, CI $-0.16$ to $0.08$, $p = 0.52$, model $R^2 = 0.28$, interaction $R^2 = 0.02$) (Figure 1D, eTable 2, links.lww.com/WNL/C103).

**APOE4 Attenuates the Relationship Between Lifestyle Activity and Cognitive Reserve in Women**

Planned secondary analyses assessed the association of activity (METS and COGACT), sex, and APOE4 status on reserve (speed and memory). Among women, the recruitment cohort, diagnoses, and age differed between APOE4 carriers and noncarriers ($p < 0.01$), such that the frequency of APOE4 carriers was lower in wave 2, and APOE4 carriers were more likely to be CN and younger. There was a difference in education between APOE4 carriers and noncarriers among men ($p = 0.04$).

There was a significant METS by sex by APOE4 interaction on speed reserve ($\beta = -0.09$, CI $-0.22$ to $0.04$, $p = 0.17$, model $R^2 = 0.33$, interaction $R^2 = 0.02$). Sex-stratified analyses indicated that the presence of APOE4 allele moderated the relationships between METS and speed reserve among women ($\beta = 0.08$, CI $0.00$–$0.15$, $p = 0.05$, model $R^2 = 0.37$, interaction $R^2 = 0.08$), such that APOE4 noncarrier women had a stronger association between METS and speed reserve ($\beta = 0.06$, CI $0.02$–$0.11$, $p < 0.01$, model $R^2 = 0.42$, METS $R^2 = 0.03$) (Figure 2A) compared with APOE4 carrier women ($\beta = -0.03$, CI $-0.10$ to $0.05$, $p = 0.52$, model $R^2 = 0.36$, METS $R^2 = 0.02$). No significant METS effect ($\beta = 0.02$, CI $-0.08$ to $0.11$, $p = 0.71$, model $R^2 = 0.27$, METS $R^2 < 0.01$) or APOE4 allele moderation was observed among men ($\beta = -0.02$, CI $-0.12$ to $0.09$, $p = 0.74$, interaction $R^2 < 0.01$) (Figure 2B). The METS by sex by APOE4 interaction for memory reserve was not significant ($\beta = -0.04$, CI $-0.14$ to $0.07$, $p = 0.48$, model $R^2 = 0.32$, interaction $R^2 < 0.01$) (Figure 2, C and D).

The COGACT by sex by APOE4 interaction effect on speed reserve was significant ($\beta = -0.26$, CI $-0.63$ to $0.11$, $p = 0.16$, model $R^2 = 0.34$, interaction $R^2 = 0.03$). Sex-stratified analyses found that APOE4 allele moderated the COGACT-speed reserve relationship among women ($\beta = 0.24$, CI $0.03$–$0.44$, $p = 0.02$, model $R^2 = 0.37$, interaction $R^2 = 0.01$), such that increased engagement in COGACT was associated with higher speed reserve for APOE4 noncarrier women ($\beta = 0.20$, CI $0.07$–$0.32$, $p < 0.01$, model $R^2 = 0.37$, COGACT $R^2 = 0.03$). This association was not observed among women APOE4 carriers ($\beta = -0.11$, CI $-0.31$ to $0.09$, $p = 0.27$, model $R^2 = 0.43$, COGACT $R^2 = 0.01$) (Figure 3A). Among men, no significant APOE4 moderation was found for COGACT-

| Participant characteristics | All | Women | Men | $p$ Value | Effect size |
|----------------------------|-----|-------|-----|-----------|-------------|
| N                          | 758 | 474   | 284 | <0.01     | —           |
| Age, y                     | 76.11 (6.31) | 76.63 (6.54) | 75.26 (5.82) | <0.01     | 0.22        |
| Education, y               | 12.14 (4.49) | 12.11 (4.44) | 12.18 (4.58) | 0.83      | 0.02        |
| Frequency of APOE4 carriers | 28.1% | 28.5% | 27.5% | 0.83      | 0.01        |
| Cohort (wave 2, 1999/wave 3, 2009) | 410/348 | 273/201 | 137/147 | 0.02      | 0.09        |
| Race/ethnicity, n (Black/Hispanic/White) | 292/224/242 | 195/142/137 | 97/82/105 | 0.05      | 0.09        |
| Diagnosis n (normal/MCI/dementia) | 449/242/67 | 279/147/48 | 170/95/19 | 0.26      | 0.06        |
| Physical activity (METS)   | 6.12 (2.03) | 5.94 (2.06) | 6.41 (1.95) | <0.01     | 0.23        |
| Cognitive activities (COGACT) | 1.42 (0.70) | 1.45 (0.72) | 1.39 (0.67) | 0.25      | 0.09        |
| Speed reserve              | 0.06 (0.92) | 0.07 (0.97) | 0.04 (0.83) | 0.71      | 0.03        |
| Memory reserve             | 0.01 (0.73) | 0.09 (0.73) | -0.13 (0.73) | <0.01     | 0.30        |

Means and SDs are reported unless otherwise noted. Sex differences were tested by the Student t test for continuous variables and with the χ² test for categorical variables. $p$ Value for APOE4 carrier frequency was determined using absolute counts and the χ² test. Effect size was determined using Cohen D for continuous variable and φ for categorical variables. METS variable represents the natural log of METS.
speed relationships ($\beta = -0.04$, CI $-0.34$ to $0.25$, $p = 0.77$, model $R^2 = 0.28$, interaction $R^2 < 0.01$) (Figure 3B).

COGACT by sex by APOE4 interaction on memory reserve was significant ($\beta = -0.20$, CI $-0.50$ to $0.093$, $p = 0.18$, model $R^2 = 0.33$, interaction $R^2 = 0.02$). Sex-stratified analyses found an APOE4 moderation among women ($\beta = 0.24$, CI $0.08$–$0.40$, $p < 0.01$, model $R^2 = 0.34$, COGACT $R^2 = 0.02$) (Figure 3C). There was a significant association among APOE4 noncarrier women ($\beta = 0.15$, CI $0.05$–$0.25$, $p < 0.01$, model $R^2 = 0.34$, COGACT $R^2 = 0.03$), whereas a non-significant COGACT-memory reserve association was observed among APOE4 carrier women ($\beta = -0.10$, CI $-0.25$ to $0.05$, $p = 0.19$, model $R^2 = 0.36$, COGACT $R^2 = 0.01$) (Figure 3C). APOE4 moderation of COGACT-memory reserve was not significant among men ($\beta = 0.05$, CI $-0.21$ to $0.31$, $p = 0.70$, model $R^2 = 0.28$, interaction $R^2 < 0.01$) (Figure 3D). Analyses were repeated excluding those with an APOE E2/E2 or E2/E4 genotype, and the findings did not change.

**Sensitivity Analyses by Group (CN vs MCI/AD and Race/Ethnicity)**

Demographic variables, physical and COGACT measures, and domain reserve scores stratified by recruitment cohorts to account for different scanners and different MRI sequences
are summarized in eTable 3, links.lww.com/WNL/C103. After adjusting for key demographic variables (i.e., age, sex, APOE4, education, and diagnosis), there were no significant differences in the memory reserve between the cohorts ($p = 0.44$) and in the speed reserve between the cohorts ($p = 0.39$). Because there were no significant differences in the reserve measurements based on the MRI scanner, no further cohort-stratified analyses were conducted.

The results did not change for sensitivity analyses stratified by impairment group. Stratified analyses revealed that the relationship between METS and speed reserve was primarily driven by CN participants (eTable 4, links.lww.com/WNL/C103). All reported interactions were observed for CN participants. Although there was a significant interaction with impairment, the relationship between METS and memory was not significant in either subgroup (eTable 4, links.lww.com/WNL/C103).

Impairment status significantly moderated COGACT-speed reserve relationship ($\beta = 0.17$, CI 0.01–0.33, $p = 0.04$, model $R^2 = 0.32$, interaction $R^2 < 0.01$, eTable 5, links.lww.com/WNL/C103). The relationship between COGACT and speed reserve was primarily driven by participants with MCI and dementia. Impairment was not a modifier in COGACT-memory reserve relationship ($\beta = 0.05$, CI −0.08 to 0.18, $p = 0.45$, model $R^2 = 0.31$, COGACT $R^2 < 0.01$, eTable 5, links.lww.com/WNL/C103).
Impairment was not a significant moderator in reserve—activity by sex by APOE4 relationships. Race/ethnicity was not a significant moderator of any of the relationships of interest (eTables 6–7, links.lww.com/WNL/C103).

**Discussion**

Understanding how the relationship between lifestyle activities and cognitive reserve is modified by sex and APOE4 status is needed to inform strategies for Alzheimer prevention and clinical trials. In the present study, physical activity was associated with speed reserve in women but not men. Based on the effect sizes observed for physical activity (METS) and age, a 2-fold increase in physical activity would be equivalent to an estimated 2.75 fewer years of processing speed aging in women. Physical activity was not associated with memory reserve in women or men. While cognitive activities were positively associated with speed reserve in both women and men; they were positively associated with memory reserve in women only. Each additional COGACT corresponded to 13 fewer years of processing speed aging (10 years among women and 17 years among men). Furthermore, the associations observed for women were attenuated by APOE4 carrier status. The plots demonstrate the association between COGACT and speed and memory reserves among women (A and C) and men (B and D) stratified by APOE4 carrier status by regressing out the covariates from both the dependent and independent variables in each panel.
status, such that APOE4 had a negative effect on the association between lifestyle activities and cognitive reserve in women only. Overall, our findings suggest that sex and APOE4 carrier status are important factors to consider in the association between the beneficial effects of lifestyle activities on cognitive reserve.

It is widely believed that physical activity confers brain benefit by maintaining or enhancing brain integrity through neuronal growth, synaptic plasticity, or dendritic spine growth based on animal studies.28–30 In the current study, physical activity mapped onto speed reserve, despite controlling for hippocampal volume, total gray matter volume, and white matter hyper-intensities. These observations suggest that in addition to physical activity preserving brain volume, physical activity may also maintain other aspects of brain health not captured by structural markers, such as functional brain networks and brain perfusion, important for processing speed.31,32 This relationship is consistent with the literature that reports that aerobic exercise more often affects frontally mediated processes, such as executive function and processing speed, than hippocampal-mediated processes, such as episodic memory.33–35 However, the effects of aerobic exercise on cognitive domains are not always consistent likely due to differences in population studied, type, duration, and intensity.36

Contrary to physical activity, which was only associated with speed reserve, cognitively stimulating activities such as reading or group classes were associated with both speed and memory reserve. Cognitive activities have been postulated to help maintain the integrity of the brain4 in a way that reflects the cognitive demands of the specific activity. For example, if playing a card game uses executive skills, playing card games more frequently would confer longer-term benefits in the cognitive domain of executive functioning.37 Based on the nature of the cognitive activities reported in the present study, such as card games and reading, these activities could conceivably involve both processing speed and memory functions. However, reverse causality cannot be ruled out, such that those with greater speed or memory reserve are more likely to engage in these types of cognitively stimulating activities, and these relationships may differ in those with normal cognition compared with those with impaired cognition.

The sex-specific associations of physical activity—speed (observed in both) and COGACT-memory (observed only in women) may be related to the types of activities women vs men engaged in. Although no differences were observed for card-playing and reading behaviors by sex, women did report higher levels of group-based classes than men. Contrary to card play and reading activities, group-based classes inherently encompass a social component that may differentially engage cognitive abilities.

These sex effects may be further modified38 by interactions with APOE4, the major genetic risk for late-onset AD.39 Compared with noncarrier women or APOE4 carrier men, women with an APOE4 allele have increased lifetime risk for AD,40,41 smaller adjusted hippocampal volumes,42–44 more pathologic levels of CSF abeta and tau,45,46 more senile plaques and neurofibrillary tangles postmortem,47 and poorer cognition.42,43 More recently, studies indicate that APOE4 carrier women also have greater tau burden based on CSF and tau PET imaging48,49 and have steeper rates of cognitive decline50 than women without an APOE4 allele. Findings from these previous studies are consistent with the unfolding story that APOE4 may dampen the beneficial relationships between lifestyle activities and cognitive reserve, with a unique effect in women, as found in the present study. Several mechanistic underpinnings may explain this unique decrement in APOE4 carrier women. For example, women experience dramatic reductions in estrogen production after menopause, which can lead to metabolic deficiencies and ultimately cognitive decline. The double-hit of APOE4 carriage exacerbates these negative effects by further reducing the availability of bioenergetic fuel.51

There are several notable limitations. The current study is focused on a cohort living in Northern Manhattan communities and thus excludes the activity patterns of suburban and rural community dwellers. The activity scores were derived from self-reported questionnaire answers, which may include some level of recall bias in reporting by sex or in those individuals with cognitive impairment. Notably, however, the observed associations between lifestyle activities and cognitive reserve were also detected in the impaired group. Structural and societal factors, which are reflected in part, by educational opportunities and attainment, are major determinants of cognitive reserve and were not directly measured or assessed in this study. Future studies with controlled activity interventions or the utilization of objective activity levels (e.g., accelerometers) to measure physical activity would help determine the validity of our findings. The present study is limited to participants who volunteered for MRI studies. Those who volunteer for MRI studies generally tend to self-report good or excellent health, thereby possibly restricting our findings to a healthier cohort. Finally, the present study uses an observational cross-sectional design limiting conclusions on causality.

Overall, the findings from the present study can be used to develop more precise lifestyle recommendations based on sex and APOE4 status. Future studies are needed to test the causal relationship between lifestyle activities and cognitive reserve and how causality is modified by sex and APOE4. Observing these sex- and APOE4-specific associations in a community-dwelling and racially-diverse cohort is a strength. Despite the large literature on modifiable lifestyle risk factors for AD, more studies are needed to establish causality to better inform dementia prevention approaches. It is indeed possible that a combination of modifiable lifestyle factors will need to be engaged for greater effect and that this may differ between women and men and by those with APOE4.

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| Name                      | Location                                                                 | Contribution                                                                 |
|---------------------------|--------------------------------------------------------------------------|------------------------------------------------------------------------------|
| Judy Pa, PhD              | Alzheimer’s Disease Cooperative Study, Department of Neurosciences, School of Medicine, UCSD Health, San Diego, CA; Mark and Mary Stevens Neuroimaging and Informatics Institute, USC Alzheimer Disease Research Center, Department of Neurology, University of Southern California, Los Angeles, CA | Drafting/revision of the manuscript for content, including medical writing for content; study concept or design; and analysis or interpretation of data |
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Appendix (continued)

| Name                       | Location                                                                 | Contribution                                                                 |
|---------------------------|--------------------------------------------------------------------------|------------------------------------------------------------------------------|
| Miguel Arce Renteria, PhD | Taub Institute for Research on Alzheimer’s Disease and the Aging Brain, Department of Neurology, Columbia University, New York City, NY | Drafting/revision of the manuscript for content, including medical writing for content, and analysis or interpretation of data |
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| Sarah E. Tom, PhD         | Department of Neurology, Vagelos College of Physicians and Surgeons and Department of Epidemiology, Mailman School of Public Health, Columbia University, New York City, NY | Drafting/revision of the manuscript for content, including medical writing for content, and analysis or interpretation of data |
| Nicole Armstrong, PhD     | Laboratory of Behavioral Neuroscience, National Institute on Aging, Bethesda, MD; Department of Psychiatry and Human Behavior, Warren Alpert Medical School of Brown University, Providence, RI | Drafting/revision of the manuscript for content, including medical writing for content, and analysis or interpretation of data |
| Kumar Rajan, PhD          | Department of Public Health Sciences, University of California, Davis, CA | Drafting/revision of the manuscript for content, including medical writing for content, and analysis or interpretation of data |
| Justina Avila-Rieger, PhD | Taub Institute for Research on Alzheimer’s Disease and the Aging Brain, Department of Neurology, Columbia University, New York City, NY | Drafting/revision of the manuscript for content, including medical writing for content, and analysis or interpretation of data |
| Yian Gu, PhD              | Taub Institute for Research on Alzheimer’s Disease and the Aging Brain, Department of Neurology, Columbia University, New York City, NY | Drafting/revision of the manuscript for content, including medical writing for content; major role in the acquisition of data; and analysis or interpretation of data |
| Nicole Schupf, PhD        | Taub Institute for Research on Alzheimer’s Disease and the Aging Brain, Department of Neurology, Columbia University, New York City, NY | Drafting/revision of the manuscript for content, including medical writing for content; major role in the acquisition of data; and analysis or interpretation of data |
| Jennifer J. Manly, PhD    | Taub Institute for Research on Alzheimer’s Disease and the Aging Brain, Department of Neurology, Columbia University, New York City, NY | Drafting/revision of the manuscript for content, including medical writing for content; major role in the acquisition of data; and analysis or interpretation of data |
| Adam Brickman, PhD        | Taub Institute for Research on Alzheimer’s Disease and the Aging Brain, Department of Neurology, Columbia University, New York City, NY | Drafting/revision of the manuscript for content, including medical writing for content; major role in the acquisition of data; and analysis or interpretation of data |
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