Sex-and race-specific associations of protein intake with change in muscle mass and physical function in older adults: the Health, Aging, and Body Composition (Health ABC) Study

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ABSTRACT

Background: Protein intake recommendations advise ≥0.8 g/kg body weight (BW)/d, whereas experts propose a higher intake for older adults (1.0–1.2 g/kg BW/d). It is unknown whether optimal protein intake differs by sex or race.

Objectives: We examined the shape of sex- and race-specific associations of dietary protein intake with 3- and 6-y changes in appendicular lean mass (aLM) and gait speed and also 6-y incidence of mobility limitation in community-dwelling older men and women.

Methods: We used data on men (n = 1163) and women (n = 1237) aged 70–81 y of the Health, Aging, and Body Composition Study. Protein intake was assessed using an FFQ (1998–1999). aLM and gait speed were measured at baseline and at 3 and 6 y. Difficulty walking one-quarter mile or climbing stairs was measured every 6 mo over 6 y. Prospective associations were evaluated with linear and Cox regression models, comparing fit of models with and without spline functions. All analyses were stratified by sex and additionally by race.

Results: Mean ± SD protein intake was 0.94 ± 0.36 g/kg adjusted body weight (aBW)/d in men and 0.95 ± 0.36 g/kg aBW/d in women. There were no strong indications of nonlinear associations. In women, higher protein intake was associated with less aLM loss over 3 y (adjusted B per 0.1 g/kg aBW/d: 39.4; 95% CI: 11.6, 67.2), specifically in black women, but not over 6 y or with gait speed decline. In men, protein intake was not associated with changes in aLM and gait speed. Higher protein intake was associated with a lower risk of mobility limitation in men (adjusted HR for 1.0 g/kg aBW/d: 0.55; 95% CI: 0.34, 0.91) and women (adjusted HR: 0.56; 95% CI: 0.33, 0.94), specifically white women.

Conclusions: Associations between protein intake and physical outcomes may vary by sex and race. Therefore, it is important to consider sex and race in future studies regarding protein needs in older adults. Am J Clin Nutr 2020;112:84–95.

Keywords: optimal intake, appendicular lean body mass, physical performance, gait speed, mobility limitation, community-dwelling, old age, spline functions

Introduction

Loss of muscle mass and loss of strength during aging contribute to an increased risk of frailty, disability, and mortality in old age (1). Dietary protein intake is a modifiable factor that may reduce these age-related processes (2). Short-term metabolic studies have shown that sufficient protein intake stimulates muscle protein synthesis (MPS) and suppresses breakdown of muscle protein (1, 3). Moreover, in observational studies, higher protein intake in old age has been associated with less decline in body weight (4), lean body mass (LM) (5–7), and muscle strength (8, 9); better physical function (9); and a lower risk of disabilities (10, 11). However, evidence from randomized controlled trials (RCTs) is less conclusive: a meta-analysis of 36 RCTs on protein supplementation showed no effects on muscular/skeletal outcomes in nonfrail older adults (12), whereas some recent trials found positive effects on LM (13), appendicular LM (aLM) (14), and gait speed (14).

The current protein recommendation established by the Institute of Medicine is 0.8 g/kg body weight (BW)/d for adults, irrespective of age, sex, and race (15), but there is an ongoing debate (16, 17) whether or not healthy older people should be recommended 1.0–1.2 g/kg BW/d (18, 19).
recommendations are the same for older men and women, despite sex differences in body composition and hormonal milieu, which may influence protein needs (20, 21). Men have relatively more muscle mass and less fat mass compared with women (22). Although with aging muscle mass generally declines and fat mass increases (23), men retain their higher muscle mass per kilogram body weight, which suggests that the protein needs of men may exceed those of women. However, women may have higher MPS rates compared with men (24, 25), suggesting that women’s protein needs may be higher. Differences in body composition also exist between white (Caucasian) and black (African American) people. Whites have more abdominal visceral fat and less bone mineral content (26–30) and lose less lean mass compared with blacks (31). Consequently, protein needs may differ between races.

The few previous studies that have studied sex-specific associations between protein intake and functional outcomes showed inconsistent findings. Some found associations in both men and women (32), whereas others found associations in women only (33, 34). Furthermore, to the best of our knowledge, none of these previous studies examined the shape of the associations—that is, whether a certain amount of protein intake is optimal.

In the Health, Aging, and Body Composition (Health ABC) Study, protein intake, muscle mass, and physical function have previously been studied. A higher protein intake was associated with smaller 3-y losses in LM and aLM (7), as well as a lower risk of mobility limitation over 6 y (11). To extend this previous work, we investigated comparable associations in a priori sex-stratified data and studied the shape of the associations to potentially detect optimal amounts of protein intake. Next, we stratified our analyses by race within sex in order to further increase homogeneity. Thus, this study aims to examine the shape of sex-specific associations of baseline dietary protein intake with 3- and 6-y changes in muscle mass and gait speed and also with 6-y incidence of mobility limitation in US older men and women. The secondary aim is to further examine whether race-specific optimal amounts of protein intake within sex can be found.

Methods

Study population

The Health ABC Study is a prospective cohort study in the United States that focuses on risk factors for functional decline and disability, including changes in body composition, in initially healthier community-dwelling older persons. At the start in 1997–1998, 3075 black and white men and women aged 70–79 y were recruited from a random sample of white Medicare-eligible residents and all black Medicare-eligible residents in the metropolitan areas of Memphis, Tennessee, and Pittsburgh, Pennsylvania. Participants were eligible if they were free of reported difficulty walking one-quarter mile and climbing up 10 steps. After the baseline measurements, follow-up data were collected annually during a clinic visit, followed by a telephone interview after 6 mo. Written informed consent was provided by all participants, and approval for the study was given by the institutional review boards of the University of Tennessee and the University of Pittsburgh.

Participants

At the first 12-mo follow-up examination, dietary intake was assessed, which served as the baseline of the current study. Participants were excluded if they dropped out before the study baseline (n = 77), had no dietary data (n = 285), had serious errors on the FFQ (n = 57), or had implausible energy intake [<800 kcal or >4000 kcal for men and <500 kcal or >3500 kcal for women (35)] (n = 59), leaving 2597 participants. For each outcome measure and each follow-up period, additional exclusions were made, as shown in Figure 1. For the 3-y analyses of aLM, participants for whom there were missing data on aLM at baseline (n = 43) or after 3 y (n = 465) or on covariates (n = 41) were excluded, leaving 2048 participants for the analytical sample. For the 6-y aLM analyses, participants for whom there were missing data on aLM at baseline (n = 43) or after 6 y (n = 1028) or on covariates (n = 31) were excluded (analytical sample: n = 1495). The analytical sample for the 3-y analyses of gait speed consisted of 2047 participants, after exclusion of those with missing data on gait speed at baseline (n = 29) or after 3 y (n = 482) or on covariates (n = 39). For the 6-y gait speed analyses, participants with missing data on gait speed at baseline (n = 29) or after 6 y (n = 1064) or on covariates (n = 23) were excluded (analytical sample: n = 1481). For the 6-y analyses of mobility limitation, participants with difficulty walking one-quarter mile and/or climbing 10 steps before or at baseline (2 consecutive reports; n = 500) were excluded, as were those with missing data on covariates (n = 45), leaving 2052 participants for the analytical sample.

Outcome measurements

Body composition was measured annually by DXA scans (Hologic 4500A, version 8.20a). aLM was used as an indicator of muscle mass and calculated as the sum of LM in arms and
Protein intake

Dietary intake during the past year was assessed at baseline (1998–1999) by a 108-item, interviewer-administered modified version of the Block FFQ (36). The list of foods was specifically developed for the Health ABC Study on the basis of 24-h recall data from the NHANES-III for older (>65 y) non-Hispanic white and black adults. Trained interviewers used wood blocks, food models, standard kitchen measures, and flash cards to assist participants in estimating their portion sizes. Intake of nutrients was determined by Block Dietary Data Systems.

Dietary protein intake was expressed in grams per kilogram adjusted body weight per day (g/kg aBW/d). Adjusted BW is the nearest BW that would put participants with underweight or overweight into a healthy BMI (in kg/m²) range: 18.5–25.0 for those aged ≤70 y and 22.0–27.0 for those aged >70 y (37). We chose to use aBW because underweight people require extra protein for building muscle tissue, whereas in overweight people, much of the excess BW consists of fat tissue requiring less protein.

Other variables

Demographic, lifestyle, and health-related factors were collected by an interviewer-administered questionnaire. Self-reported sex and age were used. Race was based on 3 sources: 1) self-report, 2) record by examiners of the physical measurements, and 3) administrative files of the Health Care Financing Administration (HCFA). If there were any discrepancies between these 3 sources, the final race variable (white/black) was based on the largest agreement between these 3 sources. For example, in case of agreement between examiners and HCFA but not with self-report, the category from HCFA was accepted. The

FIGURE 1 Flowchart of participants included in the statistical analyses. aLM, appendicular lean mass; Health ABC, Health, Aging, and Body Composition.
highest attained level of education was categorized into low (less than high school), medium (high school graduation), and high (postsecondary education). Physical activity was based on the reported time spent walking in the past 7 days (minutes per week). Categories of smoking status were never, former, and current smoker. Current use of alcohol (yes/no), derived from the FFQ, was defined as the consumption of any alcoholic beverages in the past year. Total energy intake and the Healthy Eating Index (HEI) (38) is a diet quality index that reflects compliance with the 1995 Dietary Guidelines for Americans (39) and the Food Guide Pyramid of 1992 (40) and variety in the diet. Body height was measured with a Harpenden stadiometer (Holtain). The number of chronic diseases was derived from questions on a physician’s diagnosis of arthritis, cardiac diseases, stroke, peripheral arterial disease, hypertension, pulmonary diseases, diabetes mellitus, and cancer. Use of oral steroids was determined from drug data coded by using the Iowa Drug Information System codes (41). Overnight hospitalizations in the year before the study baseline were dichotomized as none or ≥1 hospitalization. Serum creatinine was used to estimate glomerular filtration rate (eGFR), indicating kidney function, using the Modification of Diet in Renal Disease equation (42). Depressive symptoms were assessed with the 20-item Center for Epidemiological Studies Depression Scale (CES-D) (43), and cognitive function was measured with the Modified Mini-Mental State Examination (3MSE; only used in sensitivity analyses) (44). All covariates were measured at study baseline when diet was assessed (1998–1999), except for education, smoking status, height, number of chronic diseases, eGFR, depressive symptoms, and cognitive function, which were assessed 1 y earlier in 1997–1998. Last, absolute change in fat mass, measured by DXA, was calculated over 3 and 6 y.

Statistical analyses

Because our main interest was sex-specific associations, all analyses were a priori stratified by sex. For the secondary aim, we additionally stratified by race. Descriptive statistics (means and SDs or percentages) were used to summarize participants’ baseline characteristics. Absolute changes in aLM and gait speed over 3 and 6 y were checked for normality.

Prospective associations of protein intake (g/kg aBW/d, continuous) with change in aLM and change in gait speed (both continuous) were analyzed by multivariable linear regression analysis. The associations between protein intake and incidence of mobility limitation over 6 y were examined by estimating HRs using multivariable Cox proportional hazards models. The time to event was calculated as the time between study baseline (1998–1999) and 1 of the following: 1) participant’s last examination (2004–2005) for those who survived without evidence of incident mobility limitation, 2) first occurrence of incident mobility limitation, 3) participant’s last examination for those lost to follow-up, or 4) participant’s date of death for those who died with no occurrence of incident mobility limitations, whichever occurred first. The proportional hazard assumption was tested using Schoenfeld residuals.

To potentially identify optimal cutoff amounts of protein intake, spline functions were added to the linear and Cox regression models to estimate the shape of the associations between protein intake and the outcomes. A spline function is a piecewise function that can describe the association for multiple intervals of a continuous determinant, without the assumption of linear associations. The points at which 2 intervals smoothly join are called “knots”. Spline functions were added to the univariable regression models (crude model) and the models adjusted for only baseline outcome (model 1). Next to a model without a spline (no spline), we applied linear and restricted cubic splines, both with 3, 4, and 5 knots. Linear splines estimate linear functions between the knots, whereas restricted cubic splines estimate cubic functions. The latter are restricted to be linear in the end regions to provide more conservative estimates of the association where data are often sparse (45, 46). The fit of the models was tested with the likelihood-ratio test, which compares the model fit of models with and without a specific spline function. If improvement in fit was not statistically significant ($P < 0.05$) and visual inspection of the plots confirmed this, the model without spline was chosen and vice versa. We used the following consecutive steps: 1) no spline versus spline; 2) 3, 4, or 5 knots; and 3) optimal position of the inner knot(s). If a model with a spline function fitted best, the protein intake amount associated with the least decline was identified by visual inspection of the plot.

After choosing the best-fitting model per sex, outcome, and time period, potential confounders that appeared to be important based on the literature were added to the crude model, resulting in 4 additional models. Model 1 was adjusted for the baseline value of the outcome (except for incident mobility limitation because persons with mobility limitation at baseline were excluded). Model 2 was additionally adjusted for age, race, study site, educational level, walking activity, smoking, alcohol use, body height, number of chronic diseases, use of oral steroids, overnight hospitalizations, eGFR, and depressive symptoms. Model 3 was additionally adjusted for energy intake and the HEI score (fully adjusted model). To examine the influence of changes in fat mass, model 4 was additionally adjusted for 3- or 6-y change in fat mass. For our secondary aim, all analyses were additionally stratified by race—that is, performed in the 4 sex–race groups. Similar to the procedures per sex as described previously, regression models with and without spline functions were compared, after which the best-fitting model was chosen and adjustment for confounders was performed.

Sensitivity analyses were conducted by repeating the analyses of the fully adjusted model (model 3) in different samples. First, we excluded persons with a poor cognitive performance (3MSE < 80) 1 y before baseline because they might have problems recalling their dietary intake. Furthermore, we repeated the analyses for aLM and gait speed in an analytical sample of participants with outcome data at baseline and after both 3 and 6 y to fairly compare the 3-y and 6-y results.

A $P$ value $<0.05$ (2-sided) was considered statistically significant. Descriptive analyses were performed in SPSS version 26 software (SPSS) and the spline regression models in R version 3.6.1 software (R Foundation for Statistical Computing) (47).

Results

Characteristics

At baseline, participants included in any of the analytical samples ($n = 2400$) had a mean ± SD age of 74.6 ± 2.9 y, 51.5% were women, and 36.2% were black. Baseline characteristics
for men and women as well as per sex–race group are shown in Table 1. Protein intake was 71.3 ± 26.6 g/d in men and 60.7 ± 22.3 g/d in women. Protein intake below the current recommendation (0.8 g/kg aBW/d) was observed in 39.7% of men and 37.6% of women. Protein intake was 0.93, 0.94, 0.95, and 0.95 g/kg aBW/d in white men, black men, white women, and black women, respectively, and a low protein intake was also comparable between the 4 groups. Compared with the included participants (n = 2400), the excluded participants (n = 598) were more likely to be older, black, less educated, and a current smoker. They also walked less; had more depressive symptoms, chronic diseases, and overnight hospitalizations; used alcohol less often; had a poorer cognitive performance; and had a higher energy and protein intake. Baseline aLM was higher and baseline gait speed lower in those excluded, but changes in both outcome measures were similar (data not shown).

Outcome measurements

Measures of body composition, gait speed, and mobility limitation per sex and sex–race group are presented in Table 2. At baseline, mean aLM was 23.7 ± 3.5 kg and 16.5 ± 3.1 kg in men and women, respectively. The mean loss of aLM over 3 y (611 ± 1164 g in men and 349 ± 943 g in women) was doubled to, respectively, 1169 ± 1480 g (−4.9%) and 705 ± 1117 g (−4.3%) over 6 y. Men and women had comparable mean gait speed values at baseline (1.21 ± 0.20 m/s and 1.12 ± 0.20 m/s, respectively) and similar declines over time (6 y: −0.15 ± 0.17 m/s and −0.13 ± 0.17 m/s, respectively). Over 6 y, 405/1052 (38.5%) men and 430/1000 (43.0%) women developed mobility limitation.

Protein intake and change in appendicular lean mass

For the association between protein intake and change in aLM, linear or restricted cubic spline functions did not improve the regression models in men or women based on likelihood-ratio tests and visual inspection, so linear models are presented. Figure 2 shows plots of the fully adjusted models (model 3) by sex and time period. We did not observe significant associations in men over 3 and 6 y. For women, a higher protein intake at sex and time period. We did not observe significant associations

### Table 1: Baseline characteristics of older participants of the Health ABC Study, according to sex and sex–race group, 1998–1999

| Characteristic | Men | White men | Black men | Women | White women | Black women |
|----------------|-----|-----------|-----------|-------|-------------|-------------|
| Participants, n (%) | 1163 (48.5) | 800 (33.3) | 363 (15.1) | 1237 (51.5) | 731 (30.5) | 506 (21.1) |
| Characteristics | | | | | | |
| Age, y | 74.8 ± 2.9 | 74.9 ± 2.9 | 74.6 ± 2.8 | 74.4 ± 2.8 | 74.6 ± 2.8 | 74.3 ± 2.9 |
| White race, % | 68.8 | 59.1 |
| Memphis study site, % | | | |
| Educational level | | | |
| Less than high school | 22.5 | 13.1 | 43.3 | 20.4 | 9.6 | 36.0 |
| High school graduation | 25.9 | 25.5 | 26.7 | 39.1 | 42.1 | 34.8 |
| Postsecondary education | 51.6 | 61.4 | 30.0 | 40.5 | 48.3 | 29.2 |
| Walking, min/wk | 165 ± 295 | 176 ± 308 | 140 ± 264 | 116 ± 228 | 135 ± 252 | 89 ± 184 |
| Smoking, % | | | | | | |
| Never | 30.6 | 29.8 | 32.5 | 58.2 | 58.8 | 57.3 |
| Former | 60.4 | 65.6 | 48.8 | 34.0 | 34.6 | 33.0 |
| Current | 9.0 | 4.6 | 18.7 | 7.8 | 6.6 | 9.7 |
| Current alcohol use, % | 45.5 | 52.0 | 31.1 | 30.5 | 38.7 | 18.6 |
| Height, cm | 173.5 ± 64.5 | 173.6 ± 62.3 | 173.3 ± 69.2 | 159.6 ± 60.8 | 159.5 ± 60.6 | 159.7 ± 61.2 |
| Weight, kg | 81.1 ± 12.8 | 80.9 ± 12.2 | 81.5 ± 13.9 | 69.8 ± 14.5 | 65.7 ± 11.8 | 75.7 ± 15.9 |
| BMI, kg/m² | 26.9 ± 3.8 | 26.8 ± 3.6 | 27.1 ± 4.2 | 27.4 ± 5.4 | 25.8 ± 4.4 | 29.7 ± 5.9 |
| No. of chronic diseases | | | | | | |
| 0 diseases | 14.8 | 15.0 | 14.3 | 11.2 | 13.8 | 7.5 |
| 1 disease | 27.4 | 27.8 | 26.7 | 28.7 | 31.2 | 25.1 |
| ≥ 2 diseases | 57.8 | 57.3 | 59.0 | 60.1 | 55.0 | 67.4 |
| Oral steroid use, % | 2.1 | 2.8 | 0.8 | 3.2 | 4.0 | 2.0 |
| Hospitalization in past year, % | 15.2 | 16.1 | 13.2 | 11.7 | 11.1 | 12.6 |
| eGFR | 73.8 ± 16.3 | 72.2 ± 14.5 | 77.4 ± 19.2 | 71.7 ± 14.9 | 68.6 ± 12.7 | 76.3 ± 16.6 |
| Depressive symptoms, CES-D score | 4.0 ± 4.8 | 3.9 ± 4.7 | 4.3 ± 4.8 | 5.0 ± 5.5 | 5.2 ± 5.8 | 4.7 ± 5.0 |
| Cognitive function, 3MSE score | 89.5 ± 8.7 | 92.5 ± 5.8 | 85.8 ± 9.5 | 90.6 ± 8.1 | 94.0 ± 5.1 | 88.2 ± 8.4 |
| Dietary intake | | | | | | |
| Total energy intake, kcal | 2003 ± 658 | 1974 ± 633 | 2067 ± 707 | 1688 ± 565 | 1629 ± 513 | 1773 ± 624 |
| HEI score | 69.0 ± 11.8 | 71.0 ± 11.3 | 64.5 ± 11.8 | 71.1 ± 11.8 | 72.6 ± 11.7 | 68.9 ± 11.6 |
| Protein intake, g/d | 71.3 ± 26.6 | 71.3 ± 25.6 | 71.2 ± 28.7 | 60.7 ± 22.3 | 59.5 ± 20.7 | 62.6 ± 24.4 |
| Protein intake, g/kg aBW/d | 0.94 ± 0.36 | 0.93 ± 0.35 | 0.94 ± 0.40 | 0.95 ± 0.36 | 0.95 ± 0.34 | 0.95 ± 0.38 |
| <0.8 g/kg aBW/d | 39.7 | 39.1 | 41.0 | 37.6 | 36.5 | 39.1 |
| ≥1.0 g/kg aBW/d | 63 | 63.6 | 61.7 | 60.7 | 61.6 | 59.5 |

1Values are means ± SDs or %, n = 2400: persons included the analytical sample of ≥1 outcome measure. aBW, adjusted body weight; CES-D, Center for Epidemiologic Studies Depression Scale; eGFR, estimated glomerular filtration rate; Health ABC, Health, Aging, and Body Composition; HEI, Healthy Eating Index; 3MSE, Modified Mini-Mental State Examination.

2Assessed 12 mo before the study baseline (i.e., in 1997–1998).
### Table 2

| Measure                        | Women | Men |
|-------------------------------|-------|-----|
| Baseline aLM, kg              |       |     |
| 3-y change in aLM, kg         |       |     |
| 6-y change in aLM, kg         |       |     |
| Baseline fat mass, kg         |       |     |
| 3-y change in fat mass, g     |       |     |
| 6-y change in fat mass, g     |       |     |
| Baseline gait speed, m/s      |       |     |
| 3-y change in gait speed, m/s |       |     |
| 6-y change in gait speed, m/s |       |     |

| Measure                        | White women | Black women |
|-------------------------------|-------------|-------------|
| Baseline aLM, kg              | 655, 431, 371 (30.5%) | 240, 130, 123 (31.5%) |
| 3-y change in aLM, kg         | ± 3.5, ± 3.2, ± 3.7 | ± 2.4, ± 3.1, ± 2.4 |
| 6-y change in aLM, kg         | ± 1.0, ± 0.9, ± 0.8 | ± 1.0, ± 0.9, ± 0.8 |
| Baseline fat mass, kg         | 672, 290, 758 (24.7) | 312, 190, 578 (24.7) |
| 3-y change in fat mass, g     | ± 1072, ± 552, ± 750 | ± 312, ± 150, ± 180 |
| 6-y change in fat mass, g     | ± 1446, ± 1094, ± 1363 | ± 312, ± 150, ± 180 |
| Baseline gait speed, m/s      | 1.21, 1.1, 1.09 (1.1) | 1.3, 1.2, 1.12 (1.2) |
| 3-y change in gait speed, m/s | ± 0.052, ± 0.060, ± 0.031 | ± 0.13, ± 0.15, ± 0.19 |
| 6-y change in gait speed, m/s | ± 0.15, ± 0.15, ± 0.15 | ± 0.15, ± 0.15, ± 0.15 |

| Measure                        | White men | Black men |
|-------------------------------|-----------|-----------|
| Baseline aLM, kg              | 731, 363, 463 (30.5%) | 371, 187, 225 (30.5%) |
| 3-y change in aLM, kg         | ± 3.5, ± 3.2, ± 3.7 | ± 2.4, ± 3.1, ± 2.4 |
| 6-y change in aLM, kg         | ± 1.0, ± 0.9, ± 0.8 | ± 1.0, ± 0.9, ± 0.8 |
| Baseline fat mass, kg         | 672, 290, 758 (24.7) | 312, 190, 578 (24.7) |
| 3-y change in fat mass, g     | ± 1072, ± 552, ± 750 | ± 312, ± 150, ± 180 |
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| 3-y change in gait speed, m/s | ± 0.052, ± 0.060, ± 0.031 | ± 0.13, ± 0.15, ± 0.19 |
| 6-y change in gait speed, m/s | ± 0.15, ± 0.15, ± 0.15 | ± 0.15, ± 0.15, ± 0.15 |

Protein intake and change in gait speed

Linear models were also found to be appropriate for change in gait speed. Although the regression coefficients were positive as well as negative in all models, no significant associations were found in men or women (Figure 3, Supplemental Table 1). The associations remained not significant when further stratified by race (Supplemental Table 2).

Protein intake and incidence of mobility limitation

For incident mobility limitation, again no strong indication for nonlinear associations was found. Also, in none of the analyses was the proportional hazard assumption violated. A higher protein intake was associated with a lower 6-y incidence of mobility limitations in men (HR, model 3: 0.55; 95% CI: 0.34, 0.91) and women (HR, model 3: 0.56; 95% CI: 0.33, 0.94) (Figure 4, Supplemental Table 1). Additional stratification by race showed that the magnitude of the association was comparable between black and white men. The association was no longer significant in black women (HR, model 3: 0.72; 95% CI: 0.31, 1.64) but became stronger and remained significant in white women (HR, model 3: 0.46; 95% CI: 0.23, 0.92) (Figure 4, Supplemental Table 2).

Sensitivity analyses

Findings of the associations of protein intake with any outcome measure by sex were similar when participants with a low cognitive status (3MSE < 80) were excluded (n = 54–92 in men; n = 31–57 in women). The findings were also similar in the 4 sex–race groups. When the analyses were repeated in smaller samples of participants with complete aLM data after both 3 and 6 y (n = 676 men; n = 777 women), the 3-y association in women was attenuated and lost significance (B, model 3: 174 g; 95% CI: −138, 486), whereas the 6-y association did not markedly change (B, model 3: 286 g; 95% CI: −96, 668). In black women, the 3-y association was attenuated but remained significant (B, model 3: 637 g; 95% CI: 31.9, 1242). In men, the aLM findings did not change. The 3- and 6-y gait speed findings by sex remained similar in smaller samples of participants with complete gait speed data after 3 and 6 y (n = 675 men; n = 758 women); findings in the 4 sex–race groups remained similar as well.
**Discussion**

This prospective study in US older black and white men and women showed some associations between protein intake and physical outcomes, dependent on sex and race. A higher baseline protein intake was associated with less loss of muscle mass over 3 y in women, specifically black women, whereas no associations were observed over 6 y in women or in men. Protein intake was not associated with gait speed decline over 3 and 6 y in either men or women. However, higher protein intake was associated with a lower risk of mobility limitation over 6 y in both sexes and specifically white women. For all outcome measures, we did not find indications of nonlinear associations, so we could not detect an optimal amount of protein intake for any of the outcomes.

We *a priori* stratified the analyses and found sex-specific associations only for the outcome measure aLM. We observed an association for change in aLM solely in women. A higher protein...
Protein intake and physical function by sex and race

| Gait speed | 3 year                                                                 | 6 year                                                                 |
|-----------|------------------------------------------------------------------------|------------------------------------------------------------------------|
| Men       | ![Plots of associations of protein intake with 3- and 6-y change in gait speed in older participants of the Health, Aging, and Body Composition Study, according to sex. Model 3 is shown: adjusted for baseline gait speed, age, race, study site, educational level, walking, smoking, alcohol use, height, number of chronic diseases, oral steroid use, hospitalizations, eGFR, CES-D score, energy intake, and HEI score. BW, adjusted body weight; B, unstandardized regression coefficient; CES-D, Center for Epidemiologic Studies Depression Scale; eGFR, estimated glomerular filtration rate; HEI, Healthy Eating Index; y2, study baseline; y5, 3-y follow-up examination; y8, 6-y follow-up examination.](image) | ![Plots of associations of protein intake with 3- and 6-y change in gait speed in older participants of the Health, Aging, and Body Composition Study, according to sex. Model 3 is shown: adjusted for baseline gait speed, age, race, study site, educational level, walking, smoking, alcohol use, height, number of chronic diseases, oral steroid use, hospitalizations, eGFR, CES-D score, energy intake, and HEI score. BW, adjusted body weight; B, unstandardized regression coefficient; CES-D, Center for Epidemiologic Studies Depression Scale; eGFR, estimated glomerular filtration rate; HEI, Healthy Eating Index; y2, study baseline; y5, 3-y follow-up examination; y8, 6-y follow-up examination.](image) |
| Women     | ![Plots of associations of protein intake with 3- and 6-y change in gait speed in older participants of the Health, Aging, and Body Composition Study, according to sex. Model 3 is shown: adjusted for baseline gait speed, age, race, study site, educational level, walking, smoking, alcohol use, height, number of chronic diseases, oral steroid use, hospitalizations, eGFR, CES-D score, energy intake, and HEI score. BW, adjusted body weight; B, unstandardized regression coefficient; CES-D, Center for Epidemiologic Studies Depression Scale; eGFR, estimated glomerular filtration rate; HEI, Healthy Eating Index; y2, study baseline; y5, 3-y follow-up examination; y8, 6-y follow-up examination.](image) | ![Plots of associations of protein intake with 3- and 6-y change in gait speed in older participants of the Health, Aging, and Body Composition Study, according to sex. Model 3 is shown: adjusted for baseline gait speed, age, race, study site, educational level, walking, smoking, alcohol use, height, number of chronic diseases, oral steroid use, hospitalizations, eGFR, CES-D score, energy intake, and HEI score. BW, adjusted body weight; B, unstandardized regression coefficient; CES-D, Center for Epidemiologic Studies Depression Scale; eGFR, estimated glomerular filtration rate; HEI, Healthy Eating Index; y2, study baseline; y5, 3-y follow-up examination; y8, 6-y follow-up examination.](image) |

intake may thus reduce muscle mass loss in women, potentially indicating higher protein needs compared with those of men. This might be caused by older women’s higher MPS rates compared with those of older men found in some (24, 25) but not all (48–50) studies. A study in obese, older adults (aged 65–80 y) showed that the fractional synthesis rate of muscle protein was ∼30% higher in women than in men (24). Moreover, a study in healthy persons observed that muscle protein fractional synthesis rate as well as whole-body protein synthesis were higher in women than in men, at both young and old age (25). The higher MPS coincides with an accelerated loss of muscle in older women, which may imply that they also have an increased rate of protein breakdown. An upregulation of stimulatory and inhibitory muscle growth-regulatory genes was reported in postmenopausal women (51). Our observed association between protein intake and aLM in older women adds to these studies that found sex differences in MPS rates and suggests a higher dependency of dietary protein in older women compared with men. However, further study is needed to unravel these sex differences.

The race-specific analyses additionally showed that the aLM association remained only significant in black women. A potential explanation is that mean aLM loss in black women was much larger than in white women (see Table 2); the larger range of aLM change may have increased our ability to pick up an association. A higher MPS rate in black women might also explain our race-specific finding, although to our knowledge, no studies are available on race differences in MPS. Further research on the race difference of this association is needed. Another race-specific association was found for mobility limitation; only in white women was a higher protein intake significantly associated with a lower risk of mobility limitation, which is in contrast to our aLM association specifically in black women.

The effect sizes of our associations were moderate. A daily 0.2 g/kg aBW higher protein intake, for example from 0.8 to 1.0/kg aBW/d, was associated with 78.8 g (2 × 39.4) less loss of aLM in women during a 3-y period, which is 23% of the mean 3-y aLM loss (78.8/349 × 100). For mobility limitation, such increment was associated with an 11% lower risk \[\exp (-0.59/10 × 2) = 0.89\] of mobility limitation in both sexes. So our findings need to be interpreted in this light. Because we also did not adjust for multiple testing, our findings need to be interpreted with caution.

We hypothesized similar findings for gait speed and mobility limitation because of the relation between these 2 measures of physical function (52). However, although we found associations between protein intake and mobility limitation, we did not find any association for gait speed, our objective outcome measure. Because the variation in both protein intake and gait speed change was substantial, our null associations probably cannot be explained by a lack of variation. One of the few previous studies on protein intake and gait speed showed a cross-sectional association but—similar to our study—no prospective association in older women (9). Because the Health ABC
FIGURE 4  Plots of associations of protein intake with 6-y incidence of mobility limitation in older participants of the Health, Aging, and Body Composition Study, according to sex and sex–race group (if applicable). Model 3 is shown: adjusted for age, (race,) study site, educational level, walking, smoking, alcohol use, height, number of chronic diseases, oral steroid use, hospitalizations, eGFR, CES-D score, energy intake, and HEI score. aBW, adjusted body weight; CES-D, Center for Epidemiologic Studies Depression Scale; eGFR, estimated glomerular filtration rate; HEI, Healthy Eating Index.

cohort was well-functioning at recruitment, results may not be generalizable to populations with lower physical function. More studies on the protein intake and gait speed are warranted.

The 2 previous articles on the association of protein intake with aLM and mobility limitation in the Health ABC Study showed findings comparable with those of the current study. Houston et al. (7) found a significant association between energy-adjusted protein intake (i.e., residuals, continuous, and in quintiles) and 3-y aLM loss. Interactions with sex and race were tested but not significant ($P > 0.15$). To our knowledge, this is the first study that a priori stratified the analyses by sex (and race), and we observed a significant association with aLM only in women, specifically black women. We examined aLM also over 6 y, showing an attenuated association but of similar effect size in women. When we restricted our sample to those with complete longitudinal aLM data, the 3-y association disappeared, possibly indicating that the association is driven by more unhealthy persons (who were lost to follow-up). Houston et al.’s (11) article on mobility limitation, with almost the same confounders as in this study, also did not find significant interactions with
sex or race ($P > 0.20$) but showed associations stratified by sex and race (whites compared with blacks). In men, women, whites, and blacks, associations between protein intake and incident mobility limitation were observed, and after adjustment for baseline lean mass, only the association in women and whites remained significant. Comparison with our study is difficult because Houston et al. used categories of protein intake ($<0.7$, $0.7–1.0$, and $>1.0$ g/kg BW/d), whereas we used protein intake continuously and used adjusted body weight (g/kg aBW/d) in order to correct the protein intake of underweight and overweight participants. We also stratified by race within sex, which showed that the associations remained only significant in white women. Last, we studied the association shape by using spline functions, but this did not reveal optimal amounts of protein intake.

It is noteworthy that no effect modification by sex or race seems to exist based on the nonsignificant interaction terms from the previous articles (7, 11), whereas the current study clearly did show differential associations between men and women (aLM) and between blacks and whites (aLM and mobility limitation) in stratified analyses. We performed post hoc tests of interaction terms, showing that—similar to previous articles—all had a $P$ value $>0.15$, except for the protein–race interaction in the 3-y aLM analyses in women: $P = 0.093$ (crude model) and $P = 0.063$ (model 3). Because sex is increasingly recognized as an important factor in health-related research, it can be advised to report data separately for men and women (53), especially if sex differences have consistently been shown for measures of interest, such as body composition and physical function (20–22). This can also be considered for other individual (demographic) factors, including race.

Only a few observational studies have investigated protein intake and physical outcomes separately for men and women. A study in older adults did not find prospective associations of energy-adjusted protein intake with change in LM or aLM in both sexes (54). This contrasts our association between protein intake and aLM change observed only in women. Moreover, 2 longitudinal studies showed that a higher protein intake was associated with hand grip strength and physical performance (33) as well as with maintenance of subjective physical function (34) in women but not in men. However, we found associations for a subjective physical function measure in both men and women. These discrepancies in the sparse literature on sex-specific associations of protein intake, muscle mass, and physical function might be explained by differences in study design, length of follow-up, study population (including race), used measures, and adjustment for confounders.

Strengths of this study include the long follow-up of 3 and 6 y and the large sample of community-dwelling older adults allowing stratification by sex and in 4 race–sex groups. In addition, we used adjusted or “healthy” body weight for the determinant protein intake, studied objective (aLM and gait speed) as well as subjective (mobility limitation) outcome measures, and were able to adjust for various potential confounders. Last, the use of spline functions to examine the shape of associations using all available data is a strength; nonetheless, strong indications for nonlinear associations were not observed for any outcome, so spline functions were not included in the final models. Some limitations also have to be considered. First, protein intake was assessed by an FFQ, a method that provides inaccurate estimates of absolute dietary intake (55). Potential memory problems of older subjects may also have led to misreporting (56). Indeed, in the Health ABC Study, a lower cognitive function was associated with more FFQ errors, which differed by race (57). However, underreporting did not differ between sex and race groups (58).

Second, the assessment of dietary intake only at baseline may limit the validity of our results because changes in diet over time were not captured. Finally, the Health ABC cohort was well-functioning and free of mobility limitation at recruitment; thus, the generalizability of our findings to the general older population is hampered.

In conclusion, our prospective study in a US older population showed that a higher protein intake at baseline was associated with less loss of muscle mass over 3 y in women, particularly black women, but not over 6 y. In men, no associations with muscle mass were observed. In both sexes, a higher protein intake was not associated with gait speed decline over 3 and 6 y but was associated with a lower 6-y risk of mobility limitation, particularly in white women. More prospective studies with a sex- and race-specific focus are needed to elucidate whether associations between protein intake, muscle mass, and physical function differ by sex and race. RCTs that examine the effects of increasing protein intake on age-related declines should preferably enroll sufficient numbers of men and women, whites and blacks, to be able to perform analyses by sex and race.

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