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https://escholarship.org/uc/item/8gt5304p

Alzheimer's & dementia (New York, N. Y.), 3(1)

2352-8737

Zhou, Yan
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et al.

2017

10.1016/j.trci.2016.09.004

Peer reviewed
African Americans are less likely to enroll in preclinical Alzheimer’s disease clinical trials

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Abstract

Introduction: Alzheimer’s disease (AD) incidence is disproportionately high in African Americans, yet, recruitment of this community to AD clinical trials is challenging.

Methods: We compared 47 African Americans and 78 whites in their willingness to enroll in a hypothetical preclinical AD trial and examined barriers and facilitators in their decision making.

Results: African American race (OR = 0.45; 95% CI, 0.22–0.93) and score on the research attitude questionnaire (OR = 1.12; 95% CI, 1.04–1.22) were independently associated with willingness to participate. African Americans rated study risks, the requirement of a study partner, study procedures, the ratio of drug to placebo, and study location as more important factors in the decision whether to enroll than did whites.

Discussion: These results suggest that researchers will encounter challenges in recruiting African Americans to preclinical AD trials. Future research will be necessary to understand the optimal means to improve recruitment of underrepresented populations.

Keywords: Recruitment; Race; Minority; Prevention; Clinical trial

1. Introduction

Studies show substantial variation in the incidence of Alzheimer’s disease (AD) dementia across racial and ethnic groups, and African Americans may be at highest risk [1]. The incidence of AD in African Americans may be twice as high as that of whites [1–6]. In addition, the rates of dementia diagnosis [7–9], use of approved AD treatments [10], and survival [11,12] are all reduced in this community. Research is the key to understand and address these problems. Unfortunately, few African Americans participate in research [13,14]. Preclinical AD trials, such as the anti-amyloid treatment in asymptomatic AD study (A4 study) [15], are testing interventions in participants before the onset of cognitive impairment or dementia. If these studies do not include adequate representation of African American participants, then it risks perpetuating the disparities in understanding and addressing the burden of AD in this racial group.

Preclinical AD trials, similar to other clinical trials, face challenges in recruitment, especially for participants from diverse racial and ethnic populations. Strategies to enhance minority participation in AD research are well described [16,17], but a more thorough understanding of the barriers to minority recruitment based on empirical data remains an area of need. In the present study, we compared the willingness to participate in a hypothetical preclinical AD
trial between whites and African Americans. We hypothesized that the groups would differ in their approach to deciding whether to participate in AD prevention trials. We examined the influence of specific trial factors in decision making and probed for potential incentives for each racial group by analyzing both quantitative and qualitative data.

2. Methods

2.1. Participants

We performed a post hoc secondary analysis of an interview study that examined the impact of AD biomarker disclosure on AD prevention trial recruitment. The study used a mixed-methods experimental design. One-hundred thirty-two cognitively normal participants were randomly assigned to one of two hypothetical AD prevention trials that either did or did not require disclosure of the results of an amyloid positron emission tomography scan. We probed for responses of participant willingness to enroll and factors that might affect the decision. In the primary analysis [18], we found no difference in participant willingness to enroll, based on whether disclosure of amyloid status was required. In addition, we observed no interaction effects between the disclosure requirement and any other variable, including race. In the current analysis, we compared quantitative and qualitative data between 78 white and 47 African American participants.

All participants were aged 65 years or more and were interviewed in English. They had no previous diagnosis of dementia, mild cognitive impairment, or other neurological or psychiatric disease and no auditory or visual impairments that prevented the conduct of the study interview. They were recruited through a variety of mechanisms, including community education events on AD, and multiple referral sources (Table 1).

2.2. Procedure

In a face-to-face interview with a research assistant, participants were given an informed consent form (ICF) describing a hypothetical AD prevention clinical trial that was 36-months long, double-blind, 1:1 randomized, and required visits at a medical center every 6 months. Two versions of the ICF were used based on experimental assignment (details can be found in the primary article [18]). After checking for participants’ comprehension, the research assistant used structured and open-ended questions to assess participant willingness to enroll, as well as which trial factors and potential incentives affected their decision. Participants received a $25 gift card to a national retail store for their participation.

The UCLA Institutional Review Board (IRB) approved the study, and all participants underwent IRB-approved informed consent.

2.3. Measures

2.3.1. Demographics

In addition to race, we collected participant age, gender, ethnicity, education level, employment status, job freedom, family history of AD, caregiver status, perceived health condition, and residential distance to the medical center.

2.3.2. Likelihood to enroll in a prevention trial

The primary outcome was assessed using a single question: “How likely would you be to enroll in the described Alzheimer’s disease prevention trial,” with a 6-point response scale from “extremely unlikely” to “extremely likely.”

2.3.3. Importance of trial factors

Participants received structured questions on seven trial factors: frequency of visits, location of visits, length of study, requirement of a study partner, study risks, likelihood of receiving placebo, and required procedures. Participants used a 6-point rating scale from “extremely unimportant” to “extremely important” to rate each factor.

2.3.4. Incentives for participation

Participants were asked an open-ended question “are there any things that would have made you more likely to participate?” Additional structured questions asked about six incentives including receiving overall study results, personal blood test results, personal genetic test results, personal cognitive test results, financial compensation, and estimated personal risk for getting AD. Participants rated each incentive on a 6-point scale ranging from making them “much less likely to enroll” to “much more likely to enroll.”

2.3.5. Covariates

We also assessed participants’ knowledge about AD, general attitudes toward research, perceived risk for AD, and subjective cognitive performance. Knowledge about AD was measured by the AD Knowledge Scale (ADKS) [19], a 30-item true or false questionnaire, with higher scores representing greater knowledge. The Research Attitude Questionnaire [20] is a 7-item, 5-point scale (score range, 7–35), for which

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Table 1

| Source                                | Whites (N = 78) | African Americans (N = 47) |
|---------------------------------------|-----------------|-----------------------------|
| UCLA ADRC Registry, n (%)            | 25 (32.0)       | 5 (10.6)                    |
| Community talk, n (%)                | 14 (17.9)       | 16 (34.0)                   |
| Community liaison, n (%)             | 3 (3.8)         | 17 (36.2)                   |
| Community referral, n (%)            | 6 (7.7)         | 4 (8.5)                     |
| Self-referral, n (%)                 | 7 (9.0)         | 0 (0)                       |
| UCLA ADRC control subjects, n (%)    | 7 (9.0)         | 0 (0)                       |
| Banner Alzheimer’s Prevention Initiative Registry, n (%) | 5 (6.4) | 1 (2.1)                     |
| Caregiver support program, n (%)     | 3 (3.8)         | 1 (2.1)                     |
| Clinical referral, n (%)             | 2 (2.6)         | 1 (2.1)                     |
| Unknown, n (%)                       | 6 (7.7)         | 2 (4.2)                     |

Abbreviation: UCLA ADRC, University of California, Los Angeles Alzheimer’s Disease Research Center.
higher scores represent a more favorable attitude toward research. Perceived risk for AD was measured by a 5-item, 5-point scale (score range, 5–25) [21], with higher scores reflecting a higher perceived risk for AD. Subjective cognitive performance was measured by the Cognitive Change Index [22], a 20-item, 5-point scale (score range, 20–100) that assesses participants’ perceived cognitive decline, relative to their own level of function 5 years prior. Higher scores represent greater subjective decline.

2.4. Data analyses

We compared whites and African Americans on demographics and other characteristics, using unpaired t-tests for continuous variables, chi-square tests or Fisher tests for categorical variables, and Cochran–Armitage trend tests for ordinal variables (e.g., self-rated health, distance to the medical center, and job freedom).

Racial differences on willingness to enroll were examined with ordered logistic regression models. We first used unadjusted univariate models to assess for the effects of race and each covariate. Subsequently, using a multivariable model, we examined the effect of race adjusting for effects of covariates, including those covariates with P values <.2 in univariate models. We also examined potential interaction effects between race and the covariates. Because previous analyses found no effect of the requirement of biomarker status disclosure, this variable was not included in any model.

We examined racial differences on each trial factor and each incentive using Cochran–Armitage trend tests. Within each racial group, we used Friedman tests to examine for an overall difference among the factors and the incentives and Wilcoxon signed rank tests for post hoc pairwise comparisons. For open-ended responses, one investigator separated participant interviews into separate comments and developed preliminary themes in which comments were included. Comments were placed on cards and three investigators engaged in a consensus-forming exercise, examining the developed themes and assigning individual comments to those themes, blinded to participant information [23]. Participants’ responses were coded dichotomously, indicating present or absent, for each theme. We compared the racial groups on the frequencies of these defined codes.

All analyses were performed in R, version 3.1.3 [24]. Results of statistical tests are reported with a significance level of 0.05.

3. Results

3.1. Participants

Participants were recruited through multiple sources (Table 1). Community lectures served as an effective

| Characteristic                        | White     | African American | P value |
|---------------------------------------|-----------|------------------|---------|
| n                                     | 78        | 47               | .10     |
| Mean age, years ± SD (range)          | 73.9 ± 6.6 (65–89) | 72.0 ± 5.2 (66–83) | .10     |
| Female gender, n (%)                  | 51 (65.4) | 37 (78.7)        | .17     |
| Hispanic ethnicity, n (%)             | 5 (6.4)   | 1 (2.1)          | .41     |
| Mean Education, years ± SD (range)    | 16.5 ± 2.7 (11–24) | 16.1 ± 2.4 (12–24) | .51     |
| Retired, n (%)                        | 61 (79.2) | 42 (89.4)        | .22     |
| Job freedom                           |           | .75              |         |
| Never, n (%)                          | 0         | 1 (2.1)          |         |
| Rarely, n (%)                         | 3 (3.8)   | 1 (2.1)          |         |
| Frequently, n (%)                     | 13 (16.7) | 8 (17.0)         |         |
| Always, n (%)                         | 62 (79.5) | 37 (78.7)        |         |
| ADKS score, mean ± SD (range)         | 24.0 ± 2.9 (18–29) | 22.5 ± 3.5 (15–29) | .01     |
| RAQ score, mean ± SD (range)          | 29.3 ± 4.3 (7–35) | 29.9 ± 3.3 (23–35) | .35     |
| AD Caregivers, n (%)                  | 6 (7.7)   | 4 (8.5)          | >.99    |
| Family History of AD, n (%)           | 21 (27.3) | 11 (23.9)        | .84     |
| Do you know someone with AD?          | 64 (82.1) | 38 (80.9)        | >.99    |
| Risk for AD score, mean ± SD (range)  | 16.6 ± 3.8 (5–24) | 15.8 ± 4.6 (5–24) | .28     |
| Rating of overall health               |           | .02              |         |
| Excellent, n (%)                      | 23 (29.5) | 4 (8.5)          |         |
| Very good, n (%)                      | 33 (42.3) | 25 (53.2)        |         |
| Good, n (%)                           | 19 (24.4) | 15 (31.9)        |         |
| Fair, n (%)                           | 3 (3.8)   | 3 (6.4)          |         |
| Poor, n (%)                           | 0         | 0                |         |
| Distance to the medical center        |           | .14              |         |
| 0–5 miles, n (%)                      | 40 (51.3) | 13 (27.7)        |         |
| 5–15 miles, n (%)                     | 22 (28.2) | 23 (48.9)        |         |
| 15–30 miles, n (%)                    | 10 (12.8) | 8 (17.0)         |         |
| >30 miles, n (%)                      | 6 (7.7)   | 3 (6.4)          |         |

Abbreviations: ADKS, Alzheimer’s Disease Knowledge Scale; RAQ, Research Attitude Questionnaire; AD, Alzheimer’s disease.
recruitment tool for both races (18% of whites and 34% of African Americans). A third of whites were recruited from the UCLA AD Research Center potential participant registry [25]. In contrast, a high proportion (36%) of African Americans were recruited through the work of a community liaison who attended establishments such as senior centers and beauty salons, discussed the study, and distributed flyers.

The two racial groups were similar in age, education, gender, employment status, perceived risk for AD, attitudes toward research, and residential distance to the medical center (Table 2). Similar proportions of each group knew someone with AD, had a family history of AD, or were caregivers for a patient with AD. Whites had higher scores on the ADKS ($P = .01$) and better self-rated health ($P = .02$) than African Americans.

### 3.2. Willingness to participate

Seventy-two percent (56 of 78) of whites and 51% (24 of 47) of African Americans reported that they were likely to enroll in the AD prevention trial (Table 3). Univariate-ordered logistic regression showed that African Americans were less likely to enroll than whites (odds ratio = 0.45; 95% confidence interval, 0.23–0.86; Table 4). In additional univariate analyses, RAQ score (higher scores associated with greater willingness), retirement status (not retired more willing than retired), and perceived risk for AD (higher perceived risk associated with greater willingness) were significantly associated with likelihood to enroll ($P < .05$; Table 4). ADKS score ($P = .07$), Cognitive Change Index score ($P = .05$), and distance from the medical center ($P = .16$) were also included in the subsequent multivariable model.

The final multivariable model showed that, after adjusting for covariates, African Americans remained significantly less likely to enroll than whites (OR = 0.45; 95% CI, 0.22–0.93). The only other predictor that remained significant was RAQ score (OR = 1.12; 95% CI, 1.04–1.22), with every one point higher score associated with 12% higher likelihood to enroll. There was no interaction effect between race and RAQ total score or between race and any of the individual RAQ items (data not shown).

### 3.3. Importance of trial factors

African American participants rated five of seven trial factors as being of greater importance to the decision whether to enroll than did whites (Fig. 1), including study risks, the requirement of a study partner, study procedures, ratio of drug to placebo, and study location (Cochran–Armitage test, $P < .05$). Frequency of study visits and total study length were rated similarly by the two groups.

Within each racial group, Friedman tests showed that participants’ ratings of importance significantly differed among the seven trial factors ($P < .0001$). In both groups, study risks and the requirement of a study partner were rated as significantly more important than the remaining five factors (Wilcoxon signed rank test, $P < .05$ for all pairwise comparisons).
3.4. Incentives for participation

Quantitative and qualitative data showed mixed results for potential incentives for enrollment. In response to open-ended questions, African Americans more frequently mentioned financial compensation (23% vs. 14%) and returning of research results (19% vs. 6%) as potential incentives than did whites. In structured questions that examined six potential incentives (Fig. 2), there was no difference between African Americans and whites in the impact that financial incentives would have on enrollment. More whites than African Americans responded that returning cognitive test results and returning overall study results would make them more likely to enroll (Cochran Armitage test, $P < .05$). No difference between the groups was found for the remaining incentives.

Within each racial group, Friedman tests confirmed that the six incentives were rated differently ($P < .0001$ for whites, $P < .05$ for African Americans). Whites reported that receiving personal cognitive test results, personal risk estimates for getting AD, personal genetic test results, and overall study results would make them more likely to enroll than receiving personal blood test results (Wilcoxon signed rank test, $P < .05$ for all pairwise comparisons); and that financial compensation was less effective than any other incentive ($P < .05$ for all pairwise comparisons). African Americans reported that receiving personal genetic test results would make them more likely to enroll than receiving personal blood test results or receiving financial compensation ($P < .05$); and that receiving personal cognitive results would make them more likely to enroll than receiving personal blood test results ($P < .05$).

4. Discussion

In this study, African Americans were less likely to express a willingness to participate in AD prevention trials
than were whites, a finding that remained after adjusting for potential confounders such as knowledge about AD, perceived risk for AD, attitudes toward research, perceived cognitive decline, retirement status, and residential distance from the medical center. This sample is representative of community members that researchers will attempt to recruit to preclinical AD trials, as they demonstrated interest and favorable attitudes toward AD research. Therefore, our results suggest that researchers may encounter challenges in recruiting African Americans and are in contrast to some recent studies that suggest that African Americans are just as likely as whites to participate in research when presented with the opportunity [26].

African American reluctance in participating in clinical trials and medical research is well-documented [27–30]. Past research practices with African American participants that were self-serving and unethical have had profound influences on this community, resulting in distrust in doctors, scientists, and the medical system [28,31]. The consent process, with the goal of informing patients of study risks and benefits, may be misinterpreted by some African Americans as relinquishing their autonomy and as a legal protection for doctors [28]. Our findings that African Americans more heavily weighted a variety of trial aspects, including study risks and study procedures, in their decision than did whites, may partly reflect these issues. Efforts to instill trust in clinical relationships by involving African American personnel who can explain and perform trial procedures may reduce this skepticism [32]. In this study, we used an African American community liaison to aid in recruitment, and this was the greatest source of African American participants.

Both race and research attitudes were independently associated with willingness to enroll, suggesting that research attitudes alone cannot explain the reluctance of African Americans to participate in AD prevention trials. In fact, African Americans’ RAQ scores did not differ from the scores of white participants. Others have reported divergent findings. Neugroschl et al. [33] administered the RAQ to 123 diverse attendees of community talks on cognitive aging in New York City and found that nearly half of participants had a less than positive response for the item “participating in medical research is generally safe.” The authors noted that these scores were lower than previously reported means from predominantly white participants [34].

A relatively high proportion of our African American participants were recruited through community talks. African Americans in our study also had less factual knowledge about AD than did whites. African Americans may have fewer sources of information about AD than whites [35,36] and may view memory loss as a natural and expected part of aging, instead of as a sign of disease [37].

![Fig. 2. Participant ratings of potential incentives for participation for each racial group. * indicates P < .05 for racial differences.](image-url)

|                | Amount | Probability | P-value |
|----------------|--------|-------------|---------|
| **Financial compensation** |        |             |         |
| White           | 60%    | 38%         |         |
| African American| 57%    | 43%         |         |
| **Overall study results** |        |             |         |
| White           | 27%    | *           | 72%     |
| African American| 49%    | 51%         |         |
| **Personal blood test results** |        |             |         |
| White           | 51%    |             | 49%     |
| African American| 57%    |             | 43%     |
| **Personal cognitive test results** |        |             |         |
| White           | 26%    | *           | 74%     |
| African American| 43%    |             | 55%     |
| **Personal genetic test results** |        |             |         |
| White           | 28%    |             | 72%     |
| African American| 38%    |             | 60%     |
| **Personal risk for AD** |        |             |         |
| White           | 27%    |             | 72%     |
| African American| 30%    |             | 62%     |

African American reluctance in participating in clinical trials and medical research is well-documented [27–30]. Past research practices with African American participants that were self-serving and unethical have had profound influences on this community, resulting in distrust in doctors, scientists, and the medical system [28,31]. The consent process, with the goal of informing patients of study risks and benefits, may be misinterpreted by some African Americans as relinquishing their autonomy and as a legal protection for doctors [28]. Our findings that African Americans more heavily weighted a variety of trial aspects, including study risks and study procedures, in their decision than did whites, may partly reflect these issues. Efforts to
Thus, community education, potentially partnering with trusted community members, may be a promising intervention to increase minority participation in AD prevention research. Unfortunately, our results do not explicitly instruct which educational topics will be most effective to improving participation rates. Nevertheless, community programs that describe the extensive precautions in place to ensure the voluntary nature of research and the safety of participants; that African Americans are at increased risk for AD as a community; and that diverse participation is needed to reduce health care disparities represent a logical starting point [38].

Among the limitations of this study is that it is a retrospective secondary analysis of a study that measured hypothetical behaviors, rather than actual enrollment decisions. The protocol was designed to examine the impact of disclosing amyloid status on recruitment, not to examine differences among racial groups. The questions that were asked about trial barriers and facilitators were developed for a general trial audience and did not address specific racial and cultural differences. Similarly, the RAQ focuses on general attitudes toward research, not race-specific elements. The sources of white and African American participants differed substantially (Table 1). More white than African American participants had previously enrolled in longitudinal research or potential participant registries, creating the possibility that differing levels of previous participation [39], rather than race accounted for the observed findings. In sub-analyses limited to those participants who were de novo recruited for this study; however, we observed similar trends suggesting that race is associated with willingness to participate (data not shown). Even among those enrolled in AD prevention registries, African Americans may be less likely to endorse participation in trials, especially those involving a drug [40]. Thus, novel recruitment methods to not only reach diverse participants but to overcome the barriers to their enrollment will likely be necessary to successfully increase minority participation. These methods are needed not only for African Americans, but also for other racial and ethnic groups that are traditionally underrepresented in AD research, such as Latinos and Asian Americans [13,17,41].

In conclusion, our results show a significant racial difference in willingness to participate in preclinical AD trials that is not explained by the other covariates. All of our findings should be viewed as preliminary and will require assessment in future research, which should seek to more fully understand the barriers to minority enrollment and the optimal means to improve recruitment of underrepresented populations. Nevertheless, differences in AD knowledge and well-described barriers such as lack of trust suggest that community education to inform African Americans about AD risk and the need for equitable research participation to overcome health disparities may be one of the keys to improving participation rates in preclinical AD trials in this group.

Acknowledgments

We thank the participants and community liaisons for making this study possible. This work was supported by Alzheimer’s Association NIRG 12-242511. Y.Z., D.E., S.K., E.T., and J.G. were also supported by NIA AG016570. J.G. is currently supported by NIA AG016573. J.K. was supported by NIA P30-AG01024. E.T., S.K., and J.G. were supported by the Sidell-Kagan Foundation. Y.Z. performed the statistical analyses, drafted the article, and approved the final draft. D.E. and J.K. designed the study, edited the article for content, and approved the final draft. S.K. and E.T. participated in the analyses, edited the article for content, and approved the final draft. J.G. secured the funding, designed and oversaw the study, edited the article for content, and approved the final draft.

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