

2023 CARE Registry Executive Summary

CARE Registry was launched on: October 15, 2020

Enrollment number:

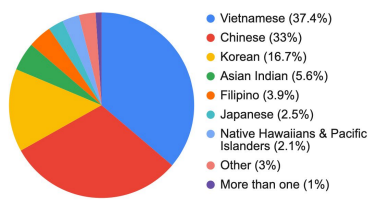
9,759 participants as of January 1, 2023

Purpose:

CARE has received 65 recruitment referral requests as of January 2023 and has connected >8,200 CARE participants with various research studies

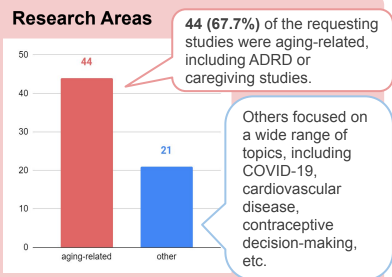
- To address the gap and reduce disparities in research participation in Alzheimer's disease and related dementias, aging, caregiving, and other health issues across the lifespan among Asian American, Native Hawaiian, and Pacific Islander (AANHPI) communities.
- Implement culturally appropriate and innovative recruitment strategies in collaboration with CARE's community partners to enroll 10,000 AANHPI participants.
- CARE aims to give AANHPI communities a voice and the opportunity to participate in research through enrollment in CARE.

Demographics of CARE Participants



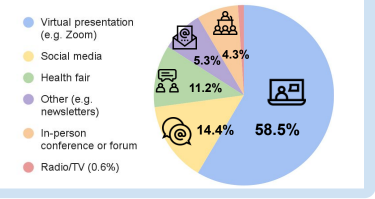
- Mean age: 53.4 (Range 18-100)
- Generale: 62.6% | Male: 35.7%
- 81% of participants do not have previous experience with research
- 13.2% of participants have self-reported ADRD-related symptoms
- 24.5% of participants are currently caregivers





Outreach Activities





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