Guidance for Industry and FDA Staff
Collection of Race and Ethnicity Data in Clinical Trials

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Overview

- Objectives
- Background
- Clinically Relevant Enrollment
- Collection of Race and Ethnicity Data
- Presentation of Race and Ethnicity Data
Objectives

• Clarify FDA’s expectations for enrolling clinically relevant populations in clinical trials

• Recommend during study design stage development of a plan to address inclusion of clinically relevant subpopulations (age, sex/gender, race, ethnicity)

• Outline FDA’s guidelines for the collection of race and ethnicity data in clinical trials
Background
1998 FDA Regulation: The “Demographic Rule”

Investigational New Drug Annual Reports & New Drug Application (NDA) Submissions:

Requires information on:

- Demographic Subgroup Trial participation
- Safety
- Effectiveness

Data by:

- Sex (Gender)
- Age
- Race

Investigational New Drug Applications (INDs) and New Drug Applications (NDAs) 21 CFR 314.50 and 21 CFR 312.33
“Demographic Rule” Cont’d

• There is **no requirement** for a specific:
  ▪ Number
  ▪ Percentage

• IND regulations
  ▪ Require annual report data to be **tabulated** by age, sex, race

• NDA regulations
  ▪ Require data for safety and efficacy to be **analyzed** by age, sex, race
FDA Safety and Innovation Act (FDASIA) of 2012
Section 907

History
• American Heart Association, WomenHeart, and Society for Women’s Health Research lobbied Congress for legislation requiring FDA to publicly report data on the inclusion and analysis of women in FDA applications
  ▪ Sen. Debbie Stabenow (D-MI) and Rep. Lois Capps (D-CA)
• Provision added to include reporting of a race and ethnicity
  ▪ Sen. Benjamin Cardin (D-MD)
• Final FDASIA legislation reauthorizing FDA user fees (essential for Agency operations)
  ▪ Requirement for an initial public report on inclusion data from medical product applications
  ▪ Subsequent action plan required to address deficiencies
Section 907 Requirements: A Report

Within one year of enactment:

• August 2013: Provided report to Congress and posted on FDA website

• Extent of clinical trial participation and the inclusion of safety and effectiveness data by demographic subgroups (sex, age, race, ethnicity) is included in applications submitted to FDA
August 2014: FDASIA Section 907 Action Plan

Three overarching priorities:

• **Priority One:** Improve the completeness and quality of demographic subgroup data collection, reporting and analysis (**Quality**)

• **Priority Two:** Identify barriers to subgroup enrollment in clinical trials and employ strategies to encourage greater participation (**Participation**)

• **Priority Three:** Make demographic subgroup data more available and transparent (**Transparency**)

FDA Report

FDA ACTION PLAN TO ENHANCE THE COLLECTION AND AVAILABILITY OF DEMOGRAPHIC SUBGROUP DATA

August 2014
Collection of Race and Ethnicity Data in Clinical Trials

Guidance for Industry and Food and Drug Administration Staff

Document issued on October 26, 2016

For questions about this document, contact the FDA Office of Minority Health at 240-402-5084 or emh@fda.hhs.gov.

U.S. Department of Health and Human Services (HHS)
Food and Drug Administration (FDA)
Office of the Commissioner (OC)
Office of Minority Health (OMH)
Office of Women’s Health (OWH)
Center for Drug Evaluation and Research (CDER)
Center for Biologics Evaluation and Research (CBER)
Center for Devices and Radiologie Health (CDRH)

October 2016
Clinical Medical
Guidance Development

- FDASIA 907 Action Plan committed to update of 2005 Guidance
- Working group convened July 2015: CDER, CBER, CDRH, OWH, OMH
- Consultation, review, and clearance included NIH, CDC, and HHS
Clinically Relevant Enrollment

• FDA expectations are that sponsors enroll participants who reflect the demographics for clinically relevant populations with regard to age, gender, race, and ethnicity

• A plan to address inclusion of clinically relevant subpopulations should be submitted for discussion to the Agency at the earliest phase of development and, for drugs and biologics, no later than the end of the phase 2 meeting

• Inadequate participation and/or data analyses from clinically relevant subpopulations can lead to insufficient information pertaining to medical product safety and effectiveness for product labeling
Points to Consider: Subgroup Differences

For potential race and ethnicity differences relevant to the evaluation of the medical product for the disease/condition, consider:

• Prevalence
• Diagnosis and treatment patterns
• Previous subgroup inclusion in past studies for target indication
• Any clinically meaningful subgroup differences in safety or efficacy
Collection of Race and Ethnicity Data
US Office of Management & Budget
Directive 15: Self-Reporting

- FDA recommends that trial participants **self-report** race and ethnicity information and those individuals be permitted to designate a multiracial identity.
- When the collection of self-reported designations is not feasible (e.g., because of the subject’s inability to respond), we recommend that the information be requested from a first-degree relative or other knowledgeable source.
- Race and ethnicity should not be assigned by the study team conducting the trial.
Two Question Format

In order to be consistent with OMB and other recommended best practices, FDA recommends using the two-question format for requesting race and ethnicity information, with the ethnicity question preceding the question about race. Example:

• **Question 1 (answer first):** Do you consider yourself Hispanic/Latino or not Hispanic/Latino?

• **Question 2 (answer second):** Which of the following five racial designations best describes you? More than one choice is acceptable.
Question 1: Ethnicity

For ethnicity, we recommend the following minimum choices be offered:

• **Hispanic or Latino**: A person of Cuban, Mexican, Puerto Rican, South or Central American, or other Spanish culture or origin, regardless of race. The term, “Spanish origin,” can be used in addition to “Hispanic or Latino.”

• **Not Hispanic or Latino**
Question 2: Race

For race, we recommend the following minimum choices be offered:

- **American Indian or Alaska Native:** A person having origins in any of the original peoples of North and South America (including Central America), and who maintains tribal affiliation or community attachment.

- **Asian:** A person having origins in any of the original peoples of the Far East, Southeast Asia, or the Indian subcontinent, including, for example, Cambodia, China, India, Japan, Korea, Malaysia, Pakistan, the Philippine Islands, Thailand, and Vietnam.

- **Black or African American:** A person having origins in any of the black racial groups of Africa. Terms such as “Haitian” or “Negro” can be used in addition to “Black or African American.”

- **Native Hawaiian or Other Pacific Islander:** A person having origins in any of the original peoples of Hawaii, Guam, Samoa, or other Pacific Islands.

- **White:** A person having origins in any of the original peoples of Europe, the Middle East, or North Africa.
Use of More Detailed Categories

- In certain situations, as recommended in OMB Policy Directive 15, more detailed race and ethnicity information may be desired.
- For example, disease/condition may warrant more granular race data.
- For clinical trials conducted outside the United States, FDA recognizes that the recommended categories for race and ethnicity were developed in the United States and that these categories may not adequately describe racial and ethnic groups in foreign countries.
- Furthermore, White can reflect origins in Europe, the Middle East, or North Africa.
- Asian can reflect origins from areas ranging from India to Japan.
- Where concerns exist in the representation of race or ethnicity categories, sponsors are encouraged to discuss the race or ethnicity issues with the appropriate review division.
## Ethnicity Data Standard

Are you Hispanic, Latino/a, or of Spanish origin? (One or more categories may be selected)

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>a.</td>
<td>____No, not of Hispanic, Latino/a, or Spanish origin</td>
</tr>
<tr>
<td>b.</td>
<td>____Yes, Mexican, Mexican American, Chicano/a</td>
</tr>
<tr>
<td>c.</td>
<td>____Yes, Puerto Rican</td>
</tr>
<tr>
<td>d.</td>
<td>____Yes, Cuban</td>
</tr>
<tr>
<td>e.</td>
<td>____Yes, Another Hispanic, Latino/a or Spanish origin</td>
</tr>
</tbody>
</table>

These categories roll up to the Hispanic or Latino category of the OMB standard
Race Data Standard

What is your race? (One or more categories may be selected)

a. _____White
b. _____Black or African American
c. _____American Indian or Alaska Native
d. _____Asian Indian
e. _____Chinese
f. _____Filipino
g. _____Japanese
h. _____Korean
i. _____Vietnamese
j. _____Other Asian
k. _____Native Hawaiian
l. _____Guamanian or Chamorro
m. ___ Samoan
n. ____Other Pacific Islander

These categories are part of the current OMB standard.

These categories roll up to the Asian category of the OMB standard.

These categories roll-up to the Native Hawaiian or Other Pacific Islander category of the OMB standard.
Electronic Submission Standards

- Beginning in May 2017, CDER and CBER will require marketing applications to be submitted electronically (use ICH M4E eCTD)
  - For INDs, NDAs, and BLAs, we recommend the submission of tabulated demographic data based on the Demographic Rule for all clinical trials using the characterizations of race and ethnicity described in this guidance.
  - When submitting an electronic application, presentation of demographic data is described in ICH M4E eCTD Guidance (section 2.7.4.1.3 and table 2.7.4.2) which suggests a tabular display of demographic characteristics (e.g., age, sex, race) by treatment group (e.g., active drug, placebo).
Drugs and Biologics

- FDA recommends that sponsors provide the level of detail described in *Section IV of this guidance*

Devices

- FDA recommends following the guidance presented here and, when finalized, the recommendations in the draft guidance document *Evaluation and Reporting of Age, Race, and Ethnicity Data in Medical Device Clinical Studies*
Summary

• Overview of legislative mandate and AP commitment
• Recommendation of a plan to address clinically relevant populations in clinical trials
• Regulatory framework for how to report the data (OMB Directive 15)
Comments to the Docket

Stakeholders can submit comments on the guidance by October 25, 2021:

- Federal eRulemaking Portal: FDA-2016-D-3561
- http://www.regulations.gov
  - Comments are public and appear unchanged, including attachments
- Mail/Hand delivery/Courier (for written/paper submissions): Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852
Thank you!

U.S. Food and Drug Administration
Office of Minority Health

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Comments & Questions