**OMB Control No**: 0910-0360 **Expiration Date**: 10/31/2023

Paperwork Reduction Act Statement: According to the Paperwork Reduction Act of 1995, an agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a valid OMB control number. The valid OMB control number for this information collection is 0910-0360. The time required to complete this information collection is estimated to average 5 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information.

Send comments regarding this burden estimate or any other aspects of this collection of information, including suggestions for reducing burden to PRAStaff@fda.hhs.gov.

**CDRH Advisory Committee Satisfaction Survey**

**Rating Scale for Questions 1-8**

(5 strongly agree, 4 agree, 3 neutral, 2 disagree, 1 strongly disagree, N/A)

1. How satisfied are you with the overall quality of the meeting?
2. How satisfied are you with the Audiovisual platform(s) used for the meeting?
3. How satisfied are you with the format of FDA’s presentation?
4. How satisfied are you with the start time of the meeting?
5. How satisfied are you with the length of the meeting?
6. How satisfied are you with your ability to communicate with the panel?
7. How satisfied are you with the format of the panelists’ discussion?
8. How satisfied are you with the voting process?
9. What are your recommendations to FDA for future virtual meetings?

(Open end question)

1. Please help us understand who you are by checking the box that applies to you.
* Patient/ Consumer/ Caregiver
* Health Care Provider/ Health Professional Organization
* Industry/ Industry Consultants/ Industry Trade Associations
* Academia
* Food and Drug Administration (FDA)
* Other Federal Agency
* Non-U.S. Regulator