2025 Current Fiscal Year Report: Patient Engagement Advisory Committee

Report Run Date: 07/02/2025 04:21:41 AM

1. Department or Agency		2. Fis	2. Fiscal Year	
Department of Health and Human				
Services		2025		
3. Committee or Subcommittee		3b. G No.	3b. GSA Committee No.	
Patient Engagement Advisory Committee		2532	2532	
4. Is this New D	uring 5. Current 6.	Expected	7. Expected	
Fiscal Year?	Charter R	enewal Date	Term Date	
No	10/06/2023 10)/06/2025		
8a. Was Terminated During8b. Specific Termination Authority8c. Actual Term Date				
Yes	2025 S Directiv	ecretary /e	04/01/2025	
9. Agency Recommendati FiscalYear	on for Next	egislation Terminate?	10b. Legislation Pending?	
Continue	Not Ap	plicable	Not Applicable	
11. Establishme	ent Authority Autho	rized by Law		
12. Specific	13.	14.		
Establishment	Effective	Commitee	14c.	
Authority	Date	Туре	Presidential?	
21 U.S.C. 394	10/06/2015	5 Continuing	No	
15. Description Board	of Committee Non	-		
16a. Total Number of Reports	No Reports for this FiscalYear			
17a. 0 17b. Closed 0 17c. Partially Closed 0 Other Activities 0 17d. Total 0 Open				
Meetings and D				

Current Next

	FY	FY
18a(1). Personnel Pmts to	\$0	00\$0.00
Non-Federal Members	ψ0.	ουψ0.00
18a(2). Personnel Pmts to	\$0	00\$0.00
Federal Members	ψ0.	οοφο.οο
18a(3). Personnel Pmts to	\$0	00\$0.00
Federal Staff	ψ0.	0000
18a(4). Personnel Pmts to	\$0	00\$0.00
Non-Member Consultants	ψ0.	0000
18b(1). Travel and Per Diem to	\$0	00\$0.00
Non-Federal Members	ψ0.	0000
18b(2). Travel and Per Diem to	\$0.	00\$0.00
Federal Members	ψ0.	0000
18b(3). Travel and Per Diem to	\$0.	00\$0.00
Federal Staff	φe.	
18b(4). Travel and Per Diem to	\$0.	00\$0.00
Non-member Consultants	φe.	
18c. Administrative Costs (FRNs,		
contractor support,	\$0.	00\$0.00
In-person/hybrid/virtual	·	·
meetings)		
18d. Other (all other funds not	^	
captured by any other cost	\$0.	00\$0.00
category)	•	
18e. Total Costs	\$0.	00\$0.00
19. Federal Staff Support Years	0.	00 0.00
(FTE)		

20a. How does the Committee accomplish its purpose?

The Committee provides advice to the Commissioner of Food and Drugs on complex scientific issues related to medical devices, the regulation of devices, and their use by patients. Agency guidance and policies, clinical trial or registry design, patient preference study design, benefit-risk determinations, device labeling, unmet clinical needs, post market safety, available alternatives, patient-reported outcomes, and device-related quality of life or health status issues are among the topics that may be considered by the Committee. The Committee provides relevant skills and perspectives to improve communication of benefits, risks and clinical outcomes, and increase integration of patient perspectives into the regulatory process for medical devices. It performs its duties by identifying new approaches, promoting innovation, recognizing unforeseen risks or barriers, and identifying unintended consequences that could result from FDA policy.

20b. How does the Committee balance its membership?

The Patient Engagement Advisory Committee charter specifies that the Committee shall consist of a core of 9 voting members including the Chair. Members and the Chair are selected by the Commissioner or designee from among authorities who are knowledgeable in areas such as clinical research, patient experience, healthcare needs of patient groups in the United States, or are experienced in the work of patient and health professional organizations, scientific methodologies for patient-reported outcomes and eliciting patient preferences, and strategies for communicating benefits, risks and clinical outcomes to patients and research subjects. As well as other relevant areas. The core of voting members may include 1 technically qualified member, selected by the Commissioner or designee, who is identified with consumer interests and are recommended by either a consortium of consumer-oriented organizations or other interested persons.

20c. How frequent and relevant are the

Committee Meetings?

The Patient Engagement Advisory Committee charter specifies that meetings shall be held approximately one to two times a year. During the meetings the Committee will provide advice to the Commissioner of Food and Drugs on complex issues relating to medical devices, the regulation of devices, and their use by patients. The Committee had no meeting in FY 2024. The committee plans to meet once in FY 2025.

20d. Why can't the advice or information this committee provides be obtained elsewhere?

The Committee provides relevant skills and perspectives in order to improve communication of benefits, risks and clinical outcomes, and increase integration of patient perspectives into the regulatory process for medical devices. The alternate means of accessing this advice would involve the recruitment of large numbers of specialists on a full-time basis at maximum rates of compensation.

20e. Why is it necessary to close and/or partially closed committee meetings? Not Applicable

21. Remarks

Per "2025 Secretary Directive" this FACA Committee has been terminated. Recruitment effort for new chairperson: Now that the preferred candidate term with a NIH Committee will officially end soon (by October 31, 2024), the Designed Federal Officer of the Patient Engagement Advisory Committee (PEAC) will work on preparing a nomination package to have that individual appointed as a Member/Chairperson of PEAC effective January 1, 2025. Selecting this start date, will allow appropriate persons at NIH enough time to take care of any outstanding pay issues and process his/her termination appointment.

Designated Federal Officer

Letise Williams Program Analyst, Center for				
Devices and Radiological Health, FDA				
Committee Members	Start	End	Occupation	Member Designation
Burkhart, Ian	05/01/2023	04/30/2027	Vice President, North American Spinal Cord Injury Consortium, Columbus, OH	Special Government Employee (SGE) Member
Diaz, Teresa	05/27/2022	04/30/2026	Co-Founder, Global Patient Advocacy Coalition, Orlando, FL	Special Government Employee (SGE) Member
Edwards, Necie	05/22/2017	04/30/2027	Health and Wellness Educator, Fibro Patient Education & Support Organization, Vernon Hills, IL	Special Government Employee (SGE) Member
Joniak-Grant, Elizabeth	06/29/2023	04/30/2027	Sociologist, Qualitative Research Consultant and Patient Experience Collaborator Injury Prevention Research Center, Univ. of North Carolina, Chapel Hill, NC	
Roy, Rita	07/23/2020	04/30/2028	CEO, National Spine Health Foundation, Reston, VA	Special Government Employee (SGE) Member
Rutherford, Philip	05/01/2022	04/30/2026	Chief Operating Officer, Faces & Voices of Recovery, Washington, DC	Special Government Employee (SGE) Member
Schrandt, Mary	05/22/2017	04/30/2026	Founder, CEO & Chief Patient Advocate, Exppect, LLC, Arlington, VA	Special Government Employee (SGE) Member

Lead Proofreader Special (Nights), Government White, David 05/01/2024 04/30/2028 Debevoise Employee Plimpton, New (SGE) York, NY Member

Number of Committee Members Listed: 8

Narrative Description

FDA's strategic priorities in responding to the public health challenges of the 21st century are to advance regulatory science and innovation; strengthen the safety and integrity of the global supply chain; strengthen compliance and enforcement activities to support public health; expand efforts to meet the needs of special populations; advance medical countermeasures and emergency preparedness; advance food safety and nutrition; promote public health by advancing the safety and effectiveness of medical products; establish an effective tobacco regulation, prevention, and control program; and manage for organizational excellence and accountability. The Patient Engagement Advisory Committee supports FDA's strategic priorities by providing advice to the Commissioner of Food and Drugs on complex, scientific issues relating to medical devices, the regulation of devices, and their use by patients. Topics to be considered by the Committee include Agency guidance and policies, clinical trial or registry design, patient preference study design, benefit-risk determinations, device labeling, unmet clinical needs, available alternatives, patient reported outcomes and device-related quality of life or health status issues. The Committee performs its duties by identifying new approaches, promoting innovation, recognizing unforeseen risks or barriers, and identifying unintended consequences that could result from FDA policy. This supports the development of safe and effective new medical technologies, and advances the status of the Agency as a science-based and science-led regulatory agency, providing global leadership in the protection of public health.

What are the most significant program outcomes associated with this committee?

Checked if Applies

Trust in government	✓
Major policy changes	~
Advance in scientific research	~
Effective grant making	
Improved service delivery	
Increased customer satisfaction	~
Implementation of laws or regulatory	~
requirements	
Other	

Outcome Comments

NA

What are the cost savings associated with this committee?

	Checked if Applies
None	
Unable to Determine	\checkmark
Under \$100,000	
\$100,000 - \$500,000	
\$500,001 - \$1,000,000	
\$1,000,001 - \$5,000,000	
\$5,000,001 - \$10,000,000	
Over \$10,000,000	
Cost Savings Other	

Cost Savings Comments

The utilization of the Patient Engagement Advisory Committee enables the Agency to obtain required services from experts who are knowledgeable in areas such as clinical research, primary care patient experience, and health care needs of patient groups in the United States not otherwise available to the Agency; and to obtain the services of these experts only on an as needed basis rather than on a full time basis. The service of the committee resulted in advice for the improvement of the public health, for which it is difficult to assign a financial value.

What is the approximate <u>Number</u> of recommendations produced by this committee for the life of the committee?

7

Number of Recommendations Comments

The Patient Engagement Advisory Committee made 7 recommendations since its establishment in FY 2016 through FY 24.

What is the approximate <u>Percentage</u> of these recommendations that have been or will be <u>Fully</u> implemented by the agency? 33%

% of Recommendations Fully Implemented Comments

The function of the committee is purely advisory in nature. Although the FDA most often accepts the recommendations from its committees, the advice is purely advisory in nature, and therefore, the Agency has the option of not implementing the advice. This number represents approximation of the percentage of recommendations that the agency has fully implemented or plans to fully implement.

What is the approximate <u>Percentage</u> of these recommendations that have been or will be <u>Partially</u> implemented by the agency?

67%

% of Recommendations Partially Implemented Comments

The function of an advisory committee is purely advisory in nature. Although the FDA most often accepts the recommendations from its committees, the advice is purely advisory in nature, and therefore, the Agency has the option of not implementing the advice.

Does the agency provide the committee with feedback regarding actions taken to implement recommendations or advice offered?

Yes 🗹 No 🗌 Not Applicable 🗌

Agency Feedback Comments

The Agency developed a guidance document to address the committee's recommendation. The guidance document and other general matters issues are available. Actions related to guidance documents or other general matters or issues are available publicly when implemented. Please see https://www.fda.gov/.

What other actions has the agency taken as a result of the committee's advice or recommendation?

Checked if Applies

Reorganized Priorities

Reallocated resources	
Issued new regulation	
Proposed legislation	
Approved grants or other payments	
Other	✓

Action Comments

The committee will address patient related issues and take appropriate programmatic steps.

Is the Committee engaged in the review of applications for grants?

No

Grant Review Comments

Not Applicable

How is access provided to the information for the Committee's documentation?

Checked if Applies

Contact DFO	\checkmark
Online Agency Web Site	\checkmark
Online Committee Web Site	\checkmark
Online GSA FACA Web Site	\checkmark
Publications	\checkmark
Other	

Access Comments

Not Applicable