2024 Current Fiscal Year Report: Pharmacy Compounding Advisory Committee

Report Run Date: 04/19/2024 10:56:16 AM

1. Department or Agency 2. Fiscal Year

Department of Health and Human

Services

2024

3b. GSA

3. Committee or Subcommittee

Committee No.

Pharmacy Compounding Advisory

Committee

5220

4. Is this New During 5. Current 6. Expected 7. Expected Fiscal Year? Charter Renewal Date Term Date

No 04/25/2022 04/25/2024

8a. Was Terminated During 8b. Specific Termination Authority 8c. Actual Term Date

No

9. Agency 10b.

Recommendation for Next Req to Terminate?

| Continuous Continuous

Continue Not Applicable Not Applicable

11. Establishment Authority Statutory (Congress Created)

12. Specific 13. 14.

Establishment Effective Committee Presidential?

Authority Date Type

21 U.S.C. 353a 11/21/1998 Continuing No

15. Description of Committee Scientific Technical Program

Advisory Board

16a. Total

No Reports for this FiscalYear Reports

17a.

Open 0 17b. Closed 0 17c. Partially Closed 0 Other Activities 0 17d. Total 0

Meetings and Dates

No Meetings

	Current Next	
	FY	FY
18a(1). Personnel Pmts to	ድለ ሰላ	\$0.00
Non-Federal Members	φυ.υι	, φυ.υυ
18a(2). Personnel Pmts to	ድለ ሰላ	\$0.00
Federal Members	φυ.υι	, φυ.υυ
18a(3). Personnel Pmts to	ድስ በወ	\$0.00
Federal Staff	φυ.υι	, φυ.υυ
18a(4). Personnel Pmts to	ያስ በያ	\$0.00
Non-Member Consultants	φυ.υι	, φυ.υυ
18b(1). Travel and Per Diem to	\$0.00	\$0.00
Non-Federal Members	φυ.υι	, φυ.υυ
18b(2). Travel and Per Diem to	\$0.00	\$0.00
Federal Members	Ψ0.00	, ψυ.υυ
18b(3). Travel and Per Diem to	\$0.00	\$0.00
Federal Staff	Ψ0.00	, ψυ.υυ
18b(4). Travel and Per Diem to	\$0.00	\$0.00
Non-member Consultants	Ψ0.00	, ψυ.υυ
18c. Other(rents,user charges,	\$0.00	\$0.00
graphics, printing, mail, etc.)	Ψ0.00	, ψυ.υυ
18d. Total	\$0.00	\$0.00
19. Federal Staff Support Years	0.00	0.00
(FTE)	0.00	, 0.00

20a. How does the Committee accomplish its purpose?

The Committee provides advice on scientific, technical, and medical issues concerning drug compounding under sections 503A and 503B of the Federal Food, Drug, and Cosmetic Act, and, as required, any other product for which the Food and Drug Administration has regulatory responsibility, and make appropriate recommendations to the Commissioner of Food and Drugs.

20b. How does the Committee balance its membership?

Members are authorities in the fields of pharmacy compounding, pharmaceutical manufacturing, pharmacy, medicine, and related specialties. The core of voting members may include one or more technically qualified members, selected by the Commissioner or designee, who are identified with consumer interests and are recommended by either a consortium of consumer-oriented organizations or other interested persons. In addition to the voting members, the Committee may include one or more non-voting member(s) who are identified with industry interests.

20c. How frequent and relevant are the Committee Meetings?

The Committee did not meet during FY-23. It is expected that the Committee will meet 1-2 times during FY-24.

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

Members of the Committee are drawn from academia, research and/or clinical practice. Their advice and input lends credibility to FDA regulatory decisions. The alternate means of obtaining this advice would involve the recruitment of large numbers of scientists on a full-time basis at a maximum rate of compensation.

20e. Why is it necessary to close and/or partially closed committee meetings?

The Committee held no closed meetings in FY-23.

21. Remarks

Although this Committee did not meet in FY-23, considerable time was devoted to appointing members, maintaining associated records for these activities, and streamlining paper processes within FDA. In addition, time was spent in the

routine care and maintenance of the Committee: the development of a financial report for this website; updating the roster and number of vacancies on our website; completing the annual ethics report; reviewing financial disclosures of current members and providing ethics training. Since the Committee did not meet, no reporting was required. Although the current charter states that the Committee shall hold meetings approximately 1-2 times a year, this is only an estimation based on data from previous years. As the FDA convenes advisory committees regarding based on the needs of the Agency, it should not be construed as an exact figure.

Designated Federal Officer

Takyiah Stevenson Designated Federal Officer

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Committee Members	Start	End	Occupation	Member Designation
Bassani, Gus	11/01/2019	10/31/2023	Chief Scientific Officer, Professional Compounding Centers of America, Inc.	Representative Member
Bogner, Robin	12/18/2020	09/30/2024	Professor, University of Connecticut School of Pharmacy	Special Government Employee (SGE) Member
Bui, Michael	11/01/2019	10/31/2023	Senior Vice-President, Global Regulatory Affairs, Pyxis Oncology	Representative Member
Desai, Seemal	10/01/2017	09/30/2025	Founder and Medical Director, Innovative Dermatology	Special Government Employee (SGE) Member
Fensky, Timothy	10/01/2018	09/30/2026	Chief Pharmacy Operations Officer, Sullivan's Pharmacy and Medical Supply, Inc., National Assoc. of Boards of Pharmacy	Representative Member

Gulur, Padma	12/18/2020	09/30/2024	Professor of Anesthesiology, Duke University Health System	Special Government Employee (SGE) Member
Gupta, Anita	12/18/2020	09/30/2024	Assistant Professor, Adjunct Johns Hopkins School of Medicine. Chief Executive Officer, Strata Group, Inc.	Special Government Employee (SGE) Member
Gura, Kathleen	12/18/2020	09/30/2024	Manager, Pharmacy Clinical Research Program, Boston Children's Hospital	Special Government Employee (SGE) Member
Rebello, Elizabeth	10/01/2018	09/30/2026	Professor, Department of Anesthesiology and Perioperative Medicine, MD Anderson Cancer Center	Special Government Employee (SGE) Member
Serumaga, Brian	04/23/2022	09/30/2026	Senior Manager, Personalized Medicines United States Pharmacopeial Convention	Representative Member
Vaida, Allen	06/03/2021	09/30/2025	Former Executive Vice President, Institute for Safe Medication Practices	Special Government Employee (SGE) Member

Number of Committee Members Listed: 11

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 Committee shall provide advice on scientific, technical, and medical issues concerning drug compounding under

sections 503A and 503B of the Federal Food, Drug, and Cosmetic Act, and, as required, any other product for which the Food and Drug Administration has regulatory responsibility, and make appropriate recommendations to the Commissioner of Food and Drugs. 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	
Improvements to health or safety	✓	
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 sovings appealed with thi	ia aammittaa?	

what are the cost savings associated with this committee?		
	Checked if Applies	
None		
Unable to Determine	✓	
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 Pharmacy Compounding Advisory Committee enabled the Agency to obtain required and frequently scarce professional services from medical and scientific experts 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?

91

Number of Recommendations Comments

The Committee made 91 recommendations from FY-12 through FY-23. The Committee was re-established in FY-12.

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

84%

% of Recommendations Fully 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, therefore, the Agency has the option of not implementing the advice. This number represents an 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? 10%

% 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 of Yes ✓ No Not Applicable	offered?	
• • •	ilable to the public. Actions related to guidance es are available publicly when implemented.	
What other actions has the agency taker recommendation?	n as a result of the committee's advice or	
	Checked if Applies	
Reorganized Priorities	~	
Reallocated resources	X	
Issued new regulation	∀	
Proposed legislation	~	
Approved grants or other payments		
Other	\(\sigma \)	
Action Comments FDA approves or chooses not to approve a Is the Committee engaged in the review		
No		
Grant Review Comments NA		
How is access provided to the information	on for the Committee's documentation? Checked if Applies	
Contact DFO	✓	
Online Agency Web Site	X	
Online Committee Web Site	Y	
Online GSA FACA Web Site	Y	
Publications	∀	
Other		

Access Comments

NA