

**FOOD AND DRUG ADMINISTRATION (FDA)  
Center for Drug Evaluation and Research (CDER)**

*Psychopharmacologic Drugs Advisory Committee (PDAC) Meeting*  
June 17, 2022

**AGENDA**

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*The committee will discuss supplemental new drug applications (sNDAs) 210793-s008 and 207318-s011, efficacy supplement resubmission for NUPLAZID (pimavanserin) tablets, submitted by Acadia Pharmaceuticals Inc., for the proposed treatment of hallucinations and delusions associated with Alzheimer's disease psychosis.*

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8:45 a.m.	Call to Order	<b>Rajesh Narendran, MD</b> Chairperson, PDAC
8:50 a.m.	Introduction of Committee/ Conflict of Interest Statement	<b>Joyce Frimpong, PharmD</b> Designated Federal Officer, PDAC
9:00 a.m.	FDA Opening Remarks	<b>Tiffany R. Farchione, MD</b> Director Division of Psychiatry (DP) Office of Neuroscience (ON) Office of New Drugs (OND), CDER, FDA
9:10 a.m.	<b>APPLICANT PRESENTATIONS</b>	<b>Acadia Pharmaceuticals Inc.</b>
	Introduction	<b>Daryl DeKarske, MPH</b> Sr. Vice President, Global Head of Regulatory Affairs and Translational Sciences Acadia Pharmaceuticals Inc.
	Unmet Need	<b>Pierre Tariot, MD</b> Director, Banner Alzheimer's Institute Research Professor of Psychiatry University of Arizona College of Medicine
	Efficacy	<b>Clive Ballard, MD</b> Pro-Vice Chancellor and Executive Dean Professor of Age-related Diseases College of Medicine and Health University of Exeter United Kingdom
		<b>Suzanne Hendrix, PhD</b> President and CEO, Pentara Corporation
	Overview of Safety	<b>Mary Ellen Turner, MD, MPH</b> Sr. Vice President, Pharmacovigilance and Corporate Safety Officer Acadia Pharmaceuticals Inc.

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**AGENDA (cont.)**

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**APPLICANT PRESENTATIONS (CONT.)**

Benefit-Risk

**Serge Stankovic, MD**  
President  
Acadia Pharmaceuticals Inc.

10:40 a.m. Clarifying Questions to Applicant

11:10 a.m. **BREAK**

11:20 a.m. **FDA PRESENTATIONS**

Pimavanserin (NUPLAZID) for the treatment of hallucinations and delusions associated with Alzheimer's disease psychosis

**Paul Bossie, MD**  
Clinical Reviewer  
DP, ON, OND, CDER, FDA

**Xiang Ling, PhD**  
Statistical Reviewer  
Division of Biometrics I (DBI)  
Office of Biostatistics (OB)  
Office of Translational Sciences (OTS)  
CDER, FDA

12:50 p.m. Clarifying Questions to FDA

1:20 p.m. **LUNCH**

2:00 p.m. **OPEN PUBLIC HEARING**

3:00 p.m. Questions to the Committee/Committee Discussion

4:30 p.m. **ADJOURNMENT**