

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

Endocrinologic and Metabolic Drugs Advisory Committee (EMDAC) Meeting

Tommy Douglas Conference Center
10000 New Hampshire Avenue, Silver Spring, Maryland
October 18, 2017

DRAFT AGENDA

The committee will discuss the safety and efficacy of new drug application (NDA) 209637 for semaglutide injection, submitted by Novo Nordisk, as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus.

8:00 a.m.	Call to Order and Introduction of Committee	Peter Wilson, MD Chairperson, EMDAC
8:05 a.m.	Conflict of Interest Statement	LaToya Bonner, PharmD, NCPS Designated Federal Officer, EMDAC
8:10 a.m.	FDA Introductory Remarks	William Chong, MD Clinical Team Lead Division of Metabolism and Endocrinology Products (DMEP), Office of Drug Evaluation II (ODE-II) Office of New Drugs (OND), CDER, FDA
8:20 a.m.	NIH PRESENTATION Review of Risk Factors Associated with Diabetic Retinopathy	Emily Y. Chew, MD Deputy Director Division of Epidemiology and Clinical Applications National Eye Institute, National Institutes of Health
8:35 a.m.	Clarifying Questions to Guest Speaker	
8:40 a.m.	APPLICANT PRESENTATIONS Introduction	Novo Nordisk Inc. Stephanie DeChiaro Director, Regulatory Affairs Novo Nordisk
	Design, Efficacy and Primary Outcomes	Anders Hvelplund, MD, PhD Senior Director, Medical and Science Novo Nordisk
	Safety	Stephen Gough, MD, FRCP (UK) Senior Principal Clinical Scientist Novo Nordisk

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DRAFT AGENDA (cont.)

APPLICANT PRESENTATIONS (CONT.)

Diabetic Retinopathy	Lloyd Paul Aiello, MD, PhD Professor and Vice Chair Department of Ophthalmology Harvard Medical School Director, Beetham Eye Institute and Vice President Joslin Diabetes Center
Retinal Safety	Stephen Gough, MD, FRCP (UK)
Clinical Perspective	Richard Pratley, MD Samuel E. Crockett, MD Chair in Diabetes Research Director, Florida Hospital Diabetes Institute Senior Scientist, Translational Research Institute for Metabolism and Diabetes
Benefit:Risk	Stephen Gough, MD, FRCP (UK)
10:10 a.m. Clarifying Questions to Applicant	
10:25 a.m. BREAK	
10:40 a.m. FDA PRESENTATIONS	
FDA Overview of Efficacy and Safety of Semaglutide	Andreea Lungu, MD Clinical Reviewer DMEP, ODE-II, OND, CDER, FDA
Statistical Assessment of Cardiovascular Safety and Retinopathy Safety of Semaglutide in the SUSTAIN 6 Trial	Ya-Hui Hsueh, PhD Mathematical Statistician Division of Biometrics VII Office of Biostatistics (OB) Office of Translational Sciences (OTS), CDER, FDA
Further Discussion of Findings for Diabetic Retinopathy	Andreea Lungu, MD
Summary of FDA Findings for Semaglutide	Andreea Lungu, MD
11:30 p.m. Clarifying Questions to FDA	
12:00 p.m. LUNCH	
1:00 p.m. OPEN PUBLIC HEARING	

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DRAFT AGENDA (cont.)

- 2:00 p.m. Questions to the Committee/Committee Discussion
- 3:45 p.m. **BREAK**
- 4:00 p.m. Questions to the Committee/Committee Discussion
- 5:00 p.m. **ADJOURNMENT**

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