

**FOOD AND DRUG ADMINISTRATION (FDA)
Center for Biologics Evaluation and Research (CBER)
77th Meeting of the Cellular, Tissue, and Gene Therapies
Advisory Committee (CTGTAC)
November 21, 2024
FINAL AGENDA**

The committee will meet in open session to discuss and make recommendations on supplemental biologics license application (sBLA) 125586/546 from Astra Zeneca to confirm clinical benefit of Andexxa (coagulation factor Xa (recombinant), inactivated -zhzo) for patients treated with rivaroxaban or apixaban when reversal of anticoagulation is needed due to life-threatening or uncontrolled bleeding.

Time EDT	Presentation/Presenter
10:00 a.m.	<p><u>Opening Remarks: Call to Order and Welcome (5 Min)</u></p> <p>Tabassum (Taby) Ahsan, Ph.D., Chairperson, CTGTAC Vice President, Cell Therapy Operations City of Hope, Duarte, CA</p> <p><u>Administrative Announcements, Roll Call, Introduction of Committee, Conflict of Interest Statement (20 Min)</u></p> <p>Cicely Reese, Pharm.D., LCDR, USPHS, Designated Federal Officer Division of Scientific Advisors and Consultants, Office of Management CBER, FDA</p>
10:25 a.m.	<p><u>FDA Introduction (5 Min)</u></p> <p>Introductory Remarks</p> <ul style="list-style-type: none"> • Nicole Verdun, M.D. Super Office Director Office of Therapeutic Products (OTP), CBER, FDA
10:30 a.m.	<p><u>Applicant Presentations (60 Min including Q & A)</u></p> <p>Introduction Jeffy John, M.B.A. Director, Regulatory Affairs AstraZeneca BioPharmaceuticals</p> <p>Burden of Life- Threatening Bleeds Related to FXa Inhibitors and Need For Effective Reversal Agents Paul A. Nyquist, M.D., M.P.H. Professor of Neurology Co-Director, Johns Hopkins Bayview Neurocritical Unit, Johns Hopkins School of Medicine</p> <p>Andexanet Efficacy Per Ladenvall, M.D., Ph.D. Global Clinical Head, AstraZeneca BioPharmaceuticals</p>

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	<p>Andexanet Safety Rohit Narayan, M.B.Ch.B. Patient Safety Physician AstraZeneca BioPharmaceuticals</p> <p>Clinical Perspective Ashkan Shoamanesh, M.D., F.R.C.P.C. Associate Professor of Medicine (Neurology) Director, Hemorrhagic Stroke Research Program Mart and Owen Boris Chair in Stroke Research and Care, McMaster University/Population Health Research Institute</p> <p>Moderator of Q & A Matthew Roe, M.D., M.H.S. Cardiologist, Adjunct Professor of Medicine Duke University Medical Center, Vice President Head of Early CVRM Clinical Development AstraZeneca BioPharmaceuticals</p> <p>Q & A (15 Min)</p>
11:30 a.m.	<p><u>FDA Presentations</u> (60 Min including Q & A)</p> <p>sBLA 125586/546 Christine Knoll, M.D. and Karl Kasamon, M.D. Medical Officers Division of Clinical Evaluation and Hematology Office of Clinical Evaluation OTP, CBER, FDA</p> <p>Q & A (15 Min)</p>
12:30 p.m.	LUNCH (40 Min)
1:10 p.m.	<u>Open Public Hearing</u> (60 Min)
2:10 p.m.	BREAK (10 Min)
2:20 p.m.	<u>Committee Discussion</u> (95 min)
3:55 p.m.	<p><u>Closing Remarks</u> (5 Min)</p> <ul style="list-style-type: none"> • Nicole Verdun
4:00 p.m.	ADJOURNMENT