## FOOD AND DRUG ADMINISTRATION (FDA)

Center for Biologics Evaluation and Research (ĆBER)
77<sup>th</sup> 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 lifethreatening or uncontrolled bleeding.

Time EDT		D/D/	
Time EDT	Presentation/Presenter		
10:00 a.m.	Opening Remarks: Call to Order and Welcome (5 Min)		
	Tabassum (Taby) Ahsan, Ph.D., Chairperson, CTGTAC Vice President, Cell Therapy Operations City of Hope, Duarte, CA		
	Administrative Annou Conflict of Interest Sta	ncements, Roll Call, Introduction of Committee, atement (20 Min)	
	Cicely Reese, Pharm.D., LCDR, USPHS, Designated Federal Officer Division of Scientific Advisors and Consultants, Office of Management CBER, FDA		
10:25 a.m.	FDA Introduction (5 Min)		
	Introductory Remarks		
	Nicole Verdun, M.D.		
	Super Office Director Office of Therapeutic Products (OTP), CBER, FDA		
	Office of Therapeutic Floudicts (OTF), OBEN, FDA		
10:30 a.m.	Applicant Presentations (60 Min including Q & A)		
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	Introduction	Jeffy John, M.B.A.	
		Director, Regulatory Affairs	
		AstraZeneca BioPharmaceuticals	
	Burden of Life-	Doul A Niversiat M.D. M.D.II	
	Threatening Bleeds	Paul A. Nyquist, M.D., M.P.H. Professor of Neurology	
	Related to FXa	Co-Director, Johns Hopkins Bayview Neurocritical	
	Inhibitors and Need	Unit, Johns Hopkins School of Medicine	
	For Effective Reversal	Offit, John's Hopkins School of Medicine	
	Agents		
	Andexanet Efficacy	Per Ladenvall, M.D., Ph.D.	
		Global Clinical Head,	
		AstraZeneca BioPharmaceuticals	

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	Andexanet Safety	Rohit Narayan, M.B.Ch.B.	
		Patient Safety Physician	
		AstraZeneca BioPharmaceuticals	
	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	
		Research institute	
	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	
	Q & A (15 Min)	Astrazerieca bior narmaceuticais	
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11:30 a.m.	FDA Presentations (60 Min including Q & A)		
	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	
		011, 022.1, 12,1	
	Q & A (15 Min)		
12:30 p.m.	LUNCH (40 Min)		
1:10 p.m.	Open Public Hearing (60 Min)		
2:10 p.m.	BREAK (10 Min)		
2:20 p.m.	Committee Discussio	<u>n</u> (95 min)	
3:55 p.m.	Closing Remarks (5 M	lin)	
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	Nicole Verdun		
4:00 p.m.	ADJOURNMENT		