Individuals using assistive technology may not be able to fully access the information contained in this file. For assistance, please call 800-835-4709 or 240-402-8010, extension 1. CBER Consumer Affairs Branch or send an e-mail to: <a href="https://occd@fda.hhs.gov">occd@fda.hhs.gov</a> and include 508 Accommodation and the title of the document in the subject line of your e-mail.

#### Andexanet in Life-Threatening or Uncontrolled Bleeding in Patients Receiving a Direct Oral Factor Xa Inhibitor

#### November 21, 2024

Cellular, Tissue and Gene Therapies Advisory Committee AstraZeneca BioPharmaceuticals



## Introduction

Jeffy John, MBA

Director, Regulatory Affairs AstraZeneca BioPharmaceuticals

## FXa Inhibitors Are Standard of Care for Anticoagulation in Many Clinical Situations, and Use Continues to Rise<sup>1-3</sup>

- FXa inhibitor specific reversal agents are needed to restore physiologic coagulation during major, life-threatening bleeds<sup>4,5</sup>
- Andexanet (ANDEXXA<sup>®</sup>) received accelerated approval in 2018
  - High unmet medical need
  - Evidence demonstrating potent reversal of FXa inhibition
- No other approved FXa inhibitor specific reversal therapies exist

## Andexanet is important component in bundle of care used to manage FXa inhibitor related uncontrolled, life-threatening bleeds

## Andexanet Is a Modified Recombinant FXa Protein that Sequesters FXa Inhibitors



 FXa inhibitors target active site of FXa, blocking enzymatic activity and preventing thrombin generation



 Andexanet acts as a decoy and binds directly to FXa inhibitors

## Andexanet binds to and sequesters FXa inhibitors



- Rapidly reduces free-plasma concentration
- Neutralizes anticoagulant effect

### Clinical Development Program Includes 4 Studies Supporting the Benefit-Risk of Andexanet



**PMR = Post-Marketing Requirement** 

### ANNEXA-A: Andexanet Reduced Anti-FXa Activity and Restored Thrombin Generation within 2 Minutes in Healthy Participants



ETP = Endogenous thrombin potential; ANNEXA-A = Apixaban-Treated Participants

## ANNEXA-4 Clearly Demonstrated that Andexanet is an Effective Reversal Agent for FXa Inhibitors

#### ANNEXA-A & ANNEXA-R

- Two prospective, randomized, placebo-controlled Phase III studies of Andexanet
- Older, healthy volunteers

Evidence supporting accelerated approval

#### ANNEXA-4

- Multinational, prospective, single-arm, open-label
   Phase IIIb/IV study
- Patients presenting with acute major bleeding, including all bleeding locations, within 18 hours of taking an FXa inhibitor

**Evidence supporting** accelerated approval

Demonstrated rapid reversal of FXa inhibitor activity Demonstrated hemostatic benefit in indicated population

# ANNEXA-I Initiated as PMR and Confirmed Hemostatic Benefit of Andexanet with Consistent Safety Profile

#### **ANNEXA-A & ANNEXA-R**

- Two prospective, randomized, placebo-controlled Phase III studies of Andexanet
- Older, healthy volunteers

Evidence supporting accelerated approval

#### **ANNEXA-4**

- Multinational, prospective, single-arm, open-label
   Phase IIIb/IV study
- Patients presenting with acute major bleeding, including all bleeding locations, within 18 hours of taking an FXa inhibitor

Evidence supporting accelerated approval

#### **ANNEXA-I**

- Randomized, open-label
   Phase IV study comparing
   Andexanet with usual care
- Patients presenting with acute intracerebral hemorrhage (ICH) within 15 hours of taking an FXa inhibitor

Post-marketing requirement trial to confirm superiority to usual care on effective hemostasis

Confirmed hemostatic benefit with acceptable and consistent safety profile

Demonstrated rapid reversal of FXa inhibitor activity Demonstrated hemostatic benefit in indicated population

## PMR Issued by FDA in 2018 Included Several Key Factors Which Were Reflected in ANNEXA-I Design

Post-Marketing Requirement for ANNEXA-I

- Include  $\geq$  440 adult patients
- Verify hemostatic effect in acute ICH
- Include assessments of NIHSS and CT / MRI at 12-hours post-randomization
- 30-day safety follow-up
- Hemostatic efficacy determined by blinded adjudication committee

### ANNEXA-I Designed to Fulfill Regulatory Requirements Globally



- Enrollment aligned with approvals of Andexanet worldwide
- Sensitivity analyses of patients receiving Apixaban or Rivaroxaban supplement primary ITT results
  - Sensitivity analyses demonstrate consistent evidence of efficacy and safety

#### Timeline for Agency Feedback on Including Long-Term Neurological Functional Assessments in ANNEXA-I

- June 2019: First patient enrolled in ANNEXA-I
- April 2020: Sponsor prepared a protocol amendment with the addition of change in NIHSS and GCS through 24 hours as secondary endpoints
- August 2020 & October 2020: FDA feedback on amendment received
  - NIHSS and GCS through 24 hours do not translate to long-term outcomes
  - GOS and mRS at 90 Days represent valid functional outcome assessments

- **November 2020**: Sponsor responded to FDA feedback
  - Follow up timeline in ANNEXA-I remained through 30 days
  - GCS and NIHSS at 24H will be added as exploratory endpoints
  - Longer term follow up outside of scope for ANNEXA-I; goal is to confirm hemostatic efficacy

## ANNEXA-I Demonstrates Positive Benefit-Risk Profile and Supports Conversion to Full Approval

#### **Unmet Need**

 Patients receiving FXa inhibitors who experience life-threatening bleeding event need reversal agents to restore physiologic coagulation and improve outcomes

#### Efficacy

- Andexanet provides statistically significant and clinically meaningful improvements hemostatic efficacy compared to usual care
- Andexanet rapidly reverses anticoagulation of FXa inhibitors

#### Safety

- Higher rate of thrombotic events compared to usual care in ANNEXA-I
- No new safety signals or adverse drug reactions were identified
- Acceptable and consistent safety profile in setting of life-threatening bleed

#### Proposed Indication Consistent with Accelerated Approval Indication

**CC-13** 

ANDEXXA<sup>®</sup> is indicated for patients treated with rivaroxaban or apixaban, when reversal of anticoagulation is needed due to life-threatening or uncontrolled bleeding

#### Agenda

#### **Unmet Need**

#### **Andexanet Efficacy**

#### **Andexanet Safety**

#### **Clinical Perspective**

#### **Moderator for Q&A**

#### Paul A. Nyquist, MD, MPH

Professor of Neurology Co-Director, Johns Hopkins Bayview Neurocritical Care Unit Johns Hopkins School of Medicine

#### Per Ladenvall, MD, PhD

Global Clinical Head AstraZeneca BioPharmaceuticals

#### **Rohit Narayan, MBCHB**

Patient Safety Physician AstraZeneca BioPharmaceuticals

#### Ashkan Shoamanesh, MD, FRCPC

Marta and Owen Boris Chair in Stroke Research / Care Associate Professor, Medicine Department of Neurology McMaster University

#### Matthew Roe, MD, MHS

Cardiologist, Adjunct Professor of Medicine Duke University Medical Center Vice President, Head of Early CVRM Clinical Development AstraZeneca BioPharmaceuticals

#### **Additional Experts**

#### **Krister Bamberg, PhD**

Senior Principal Scientist AstraZeneca BioPharmaceuticals

#### Mikael Knutsson, PhD

Statistical Science Director AstraZeneca BioPharmaceuticals

#### Magnus Nord, MD, PhD

Vice President Global Patient Safety AstraZeneca BioPharmaceuticals

#### Anita Osborne, MSc, MBA

Senior Regulatory Affairs Director AstraZeneca BioPharmaceuticals

#### **Chris Penland, PhD**

Senior Director, Clinical Pharmacology and Pharmacometrics AstraZeneca BioPharmaceuticals

## Alex C. Spyropoulos, MD, FACP, FCCP, FRCPC

Professor, Institute of Health System Science, Feinstein Institutes for Medical Research Director, Anticoagulation and Clinical Thrombosis Services, Northwell Health

#### Anna Sundgren, PhD, MBA

Global Product Leader AstraZeneca BioPharmaceuticals



Burden of Life-Threatening Bleeds Related to FXa Inhibitors and Need for Effective Reversal Agents Paul A. Nyquist, MD, MPH

Professor of Neurology Co-Director, Johns Hopkins Bayview Neurocritical Care Unit Johns Hopkins School of Medicine

## Use of FXa Inhibitors Increasing as They Become New Standard of Care for Anticoagulation



Guidelines recommend FXa inhibitors in patients with venous thromboembolism and patients with atrial fibrillation<sup>2–5</sup>

**CC-17** 

- American Heart Association
- American College of Cardiology
- National Institute for Health and Care Excellence
- Heart Rhythm Society
- European Heart Rhythm Association

**3–5%** of patients on FXa inhibitors require hospitalization due to life-threatening bleeding<sup>2-4</sup>

1. https://data.cms.gov/summary-statistics-on-use-and-payments/medicare-medicaid-spending-by-drug/medicare-part-d-spending-by-drug/data 2. January, 2019; 3. Joglar, 2023; 4. National Institute for Health and Care Excellence: NG196/NG158, 2024; 5. Steffel, 2021

### FXa Inhibitor-Associated Major Bleeds Leading to Hospitalization Can Occur at Various Body Sites

**CC-18** 



Coleman, 2021 (US-based multicentre, retrospective, electronic medical record survey conducted between 2016 and 2019. N=45 hospitals and N=3030 hospitalizations)

### In Emergency Setting, Primary Goal is to Stop Bleeding Event and Minimize Hematoma Expansion

#### **Stop bleeding to reduce brain injury and death**<sup>1</sup>

- Door: Stabilize patient, rapid imaging, coagulation tests
- < 30 min: Reverse anticoagulant, start intensive BP lowering</p>
- **< 60 min:** SBP < 140 mmHg, consult neurosurgery, achieve temp < 37.5°C
- 7 days: Maintain SBP < 140 mmHg, temp < 37.5°C, maintain normoglycemia

**CC-19** 

#### Bundle of Care

Goal

## Multiple simultaneous, specific and fast-acting interventions required to stabilize patient and control the bleeding

- Blood pressure management
- Surgical procedures
- Targeted anticoagulant reversal agents

## For Patients on Anticoagulant Therapies Thrombotic Events are a Known Risk Associated with Effective Reversal



#### **TEs are Manageable in Acute Setting**

 Emergency and critical care teams ensure appropriate measures in place to effectively manage in ICU

#### Minimizing TE Risk is Key Aspect of Early Secondary Prevention

- Early VTE prophylaxis per AHA guidelines<sup>1</sup> (24-48 hours from index event)
- Restart anticoagulant therapy as early as possible based on individualized risk benefit assessment

1. Greenberg, 2022

## Reducing Hematoma Expansion is Critical Goal for Patients Experiencing a Life-Threatening Intracerebral Hemorrhage



#### Early Neurologic Deterioration<sup>1</sup>

OR (95% CI): 3.00 (2.05, 4.41), p < 0.001



Hematoma expansion predicts **Poor Clinical Outcomes** 



**Worsening Functional Outcomes**<sup>2</sup>

Cumulative OR (95% CI): 0.84 (0.75, 0.92), p < 0.001



For every **1 mL** increase in hematoma volume, there is a **5%** higher risk of death or dependency at 90 days<sup>3</sup>

1.Kuohn, 2022; 2.Davis, 2006; 3. Delcourt, 2012

#### Multiple Studies have Characterized Mortality Risk Associated with ICH in Patients Receiving FXa Inhibitors

**CC-22** 



1. Coleman, 2021; 2. Milling, 2018; 3. Xian, 2021; 4. Held, 2015; 5. Hankey, 2014

## Andexanet is the Only Approved FXa-Inhibitor Specific Reversal Agent for Patients Taking Apixaban or Rivaroxaban<sup>1</sup>

Anticoagulant	Class	Intervention	Mechanism of action	Primary Endpoint	
Apixaban	- EXa inhibitor	Andexanet <sup>1</sup>	Reverses FXa	Anti-FXa Activity	
Rivaroxaban	Γλαπησιτοι	Andexanet	inhibition	Efficacy <sup>2</sup>	
Warfarin	Vitamin K antagonist	4F-PCC <sup>3,4</sup>	Replaces factors II, VII, IX, and X	INR, Hemostatic Efficacy⁵	
Dabigatran	Direct thrombin inhibitor	Idarucizumab <sup>6</sup>	Restores thrombin generation	ECT/dTT Normalization <sup>7</sup>	

INR = international normalized ratio; ECT = ecarin clotting time; dTT = diluted thrombin time

1. ANDEXXA (andexanet alfa) USPI; 2. Milling, 2023; 3. KCENTRA (4F-PCC) USPI; 4. BALFAXAR (4F-PCC) USPI; 5. Ansell, 2008; 6. PRAXBIND (Idarucizumab) USPI; 7. Pollack, 2017

## Multiple Guidelines Support Use of a Specific Reversal Agent for FXa Inhibitor-Related Major Bleeding Episodes

American Heart Association

#### American Stroke Association

#### Andexanet alfa is

reasonable to reverse anticoagulant effect of FXa inhibitors in patients with direct FXa inhibitorassociated ICH (Class 2a recommendation)<sup>1</sup> American College of Cardiology

#### Administer and exanet alfa

first line for reversal of anticoagulation in patients taking FXa inhibitors with major critical site or major bleeding on oral direct FXa inhibitors; PCC or activated PCC is suggested only if andexanet is not available<sup>2</sup> American College of Emergency Physicians

Andexanet alfa is a tier 1 recommendation for anticoagulation in reversal of apixaban- or rivaroxabantreated patients experiencing life-threatening or critical site bleeds; PCC is suggested as a tier 2 recommendation if a tier 1 agent is not available<sup>3</sup>

#### **Code ICH Protocol Bundled Care** *Neurocritical Care Society, and Neurocritical Care Foundation*

- Rapid neuroimaging
- Blood pressure control
- Anticoagulant reversal >>>
- Neurological assessments
- Surgical therapy
- Glucose control
- Temperature control

#### **Anticoagulant Reversal Goals**

Choose the right agent, and treat within 60 minutes:

- Vitamin K antagonists (warfarin, coumarins):
   IV vitamin K and 4F-PCC achieve INR ≤ 1.4
- Factor II inhibitor (dabigatran): Idarucizumab
- Factor Xa inhibitor (rivaroxaban, apixaban): Andexanet if available; 4F-PCC second line
- Antiplatelet agents (aspirin, clopidogrel): Desmopressin

## Patients Experiencing an FXa Inhibitor-Related Life-Threatening Bleed Need Specific Reversal Agents

- Timely intervention is crucial to prevent complications
- Need amplified by increasing number of hospital admissions due to FXa inhibitor-related bleeding events
- Andexanet rapidly neutralizes effects of FXa inhibitors
  - Offering a targeted solution for managing bleeding events

## Andexanet is vital component in bundle of care used to rapidly manage FXa inhibitor-related, life-threatening bleeding events



## **ANNEXA-I Efficacy**

Per Ladenvall, MD, PhD

Global Clinical Head AstraZeneca BioPharmaceuticals R&D

## ANNEXA-I: Randomized, Open-Label, Multicenter, Clinical Phase IV Study



- Imaging performed at baseline and 12 hours following randomization
- Neurologic assessments (NIHSS) performed at baseline, 2, 3, 6, 12, 24, and 72 hours
- Safety follow-up for 30 days after treatment

### ANNEXA-I: Primary Efficacy Endpoint Covering Different Aspects of Hemostasis

Primary Endpoint: Effective hemostasis 12 hours post-randomization

**CC-29** 

Hematoma Expansion: ≤ 35% increase in hematoma volume AND Neurological Deterioration: < 7-point change in NIHSS AND No Rescue Therapy: Between 3- and 12-hours post-randomization

### ANNEXA-I: DSMB Recommended Study be Stopped Based on Results from Planned Interim Analysis



### **ANNEXA-I:** Baseline Demographics

Primary Effic	cacy Population	Andexanet N = 224	Usual Care N = 228
Age (years)	), <b>Mean</b> (SD)	<b>78.9</b> (8.5)	<b>78.9</b> (8.5)
Male		58%	50%
Base	Black/ African American	2%	1%
Race	White	93%	94%
Ethnicity	Hispanic or Latino	6%	5%
Ethnicity	Not Hispanic or Latino	85%	90%
Dogion	Europe	88%	89%
Region	North America	12%	11%

### **ANNEXA-I:** Disease Characteristics

Primary Efficacy Population	Andovanat	Usual Caro
	N = $224$	N = 228
FXa inhibitor		
Apixaban	63%	59%
Rivaroxaban	29%	29%
Edoxaban	9%	11%
Hematoma volume (mL), median (Q1, Q3)	<b>10.6</b> (4.1, 24.6)	<b>9.0</b> (3.3, 22.8)
Baseline NIHSS, median (Q1, Q3)	<b>9.0</b> (6.0, 16.0)	<b>9.0</b> (4.0, 14.0)
Symptom onset to baseline scan (hours), median (min, max)	<b>2.3</b> (0.2, 11.4)	<b>2.4</b> (0.3, 11.9)
Symptom onset to treatment (hours), median (min, max)	<b>4.0</b> (1.3, 12.6)	<b>4.1</b> (1.2, 13.5)
Baseline scan to treatment (hours), median (min, max)	<b>1.5</b> (0.2, 4.5)	<b>1.7</b> (0.2, 4.0)

### **ANNEXA-I:** Baseline Medical History

Primary Efficacy Population – Safety Set	Andersonat		
Medical History	N = 223	N = 226	
Atrial fibrillation	90%	85%	
Diabetes	37%	26%	
Stroke	21%	21%	
Congestive heart failure	13%	20%	
Myocardial infarction	10%	14%	
Deep vein thrombosis	8%	10%	
Pulmonary embolism	8%	9%	
CHA <sub>2</sub> DS <sub>2</sub> -VASC score, median	4	4	

CHA<sub>2</sub>DS<sub>2</sub>-VASC score – stratification of risk of stroke in patients with AF estimated using the Congestive heart failure, Hypertension, Age, Diabetes mellitus, prior Stroke or TIA or thromboembolism, VAscular disease, Sex Category score

**ANNEXA-I Efficacy** 

#### ANNEXA-I: Primary Efficacy Endpoint Established Hemostatic Benefit of Andexanet vs Usual Care



# ANNEXA-I: Treatment Effect of Andexanet Consistent Across Key Patient Subgroups

Primary Efficad	y Population	Andowenet		Favors Andexanet
		Andexanet	Usual Care	
Overall		<b>67%</b> (150/224	) <b>53%</b> (121/228)	
	< 65	<b>77%</b> (10/13)	<b>67%</b> (10/15)	
Age	65 – 74	<b>62%</b> (28/45)	<b>41%</b> (19/46)	
	≥ 75	<b>67%</b> (112/166	<b>55%</b> (92/167)	
Cov	Male	<b>63%</b> (82/130)	<b>44%</b> (51/115)	
Sex Female	Female	<b>72%</b> (68/94)	<b>62%</b> (70/113)	
Location	North America	<b>50%</b> (13/26)	<b>52%</b> (13/25)	
Location	Europe	<b>69%</b> (137/198	<b>53%</b> (108/203)	
	Apixaban	<b>70%</b> (98/140)	<b>56%</b> (76/135)	
FXa Inhibitor	Rivaroxaban	<b>56%</b> (36/64)	<b>46%</b> (30/65)	
	Edoxaban	<b>80%</b> (16/20)	<b>56%</b> (14/25)	
	< 3	<b>67%</b> (130/195	<b>56%</b> (112/199)	
ICH Score	≥ 3	<b>69%</b> (20/29)	<b>31%</b> (9/29)	
Hematoma	< 30 mL	<b>69%</b> (125/180	<b>57%</b> (109/192)	
Volume	≥ 30 mL	<b>57%</b> (25/44)	<b>34%</b> (12/35)	
			-1	100 -50 0 50 100
				Effective Hemostasis at 12 hours

Treatment Difference (95% CI)

## ANNEXA-I: Andexanet Showed Numerical Improvements vs Usual Care in All Components of Primary Endpoint

**Primary Efficacy Population** 



CC<u>-37</u>

#### **ANNEXA-I:** And examet Rapidly Reduced Anti-FXa Activity



### ANNEXA-I: Confirmed Hemostatic Benefit of Andexanet Compared to Usual Care

- Statistically significant reduction in anti-FXa activity vs usual care
- Statistically significant and clinically meaningful improvement in effective hemostasis vs usual care
  - Consistent benefit across sensitivity analyses and exploratory patient subgroup analyses
- Improvement in all aspects of effective hemostasis
  - Hematoma expansion, neurological function, use of rescue therapy

**CC-39** 

ANNEXA-I confirms findings from ANNEXA-4 supporting benefit for patients with life-threatening, uncontrolled bleeding



## **ANNEXA-I** Safety

Rohit Narayan, MBCHB

Patient Safety Physician AstraZeneca BioPharmaceuticals

### Andexanet Has a Well-Established Safety Profile

	Healthy Volunteers	ANNEXA-4	ANNEXA-I Safety Set	
	Andexanet	Andexanet	Andexanet	Usual Care
Patients (N)	553	479	262	265

- Estimated\* cumulative global post-marketing exposure includes 70,158 patients
  - 36,597 patients in United States
- No new safety signals identified in clinical trials or post-marketing use

## ANNEXA-I: Overall Summary of Treatment-Emergent

	Safety Set		Apixaban and Rivaroxab	
<b>%</b> (n)	Andexanet N = 262	Usual Care N = 265	Andexanet N = 239	Usual Care N = 232
TEAE	<b>85.1%</b> (223)	<b>82.6%</b> (219)	<b>85.8%</b> (205)	<b>81.9%</b> (190)
TESAE	<b>45.8%</b> (120)	<b>36.2%</b> (96)	<b>46.4%</b> (111)	<b>37.1%</b> (86)
TEAE leading to withdrawal of study drug	0	0	0	0
TEAE leading to interruption of study drug	<b>0.4%</b> (1)	0	<b>0.4%</b> (1)	0
TEAE leading to death	<b>24.4%</b> (64)	<b>20.4%</b> (54)	<b>24.7%</b> (59)	<b>21.1%</b> (49)
All-cause mortality (30-days)	<b>28.2%</b> (74)	<b>26.4%</b> (70)	<b>28.0%</b> (67)	<b>26.3%</b> (61)

## ANNEXA-I: Treatment-Emergent SAEs Occurring in ≥ 2% of Patients

Safety Set	Andexanet	Usual Care
<b>%</b> (n)	N = 262	N = 265
TESAE	<b>46%</b> (120)	<b>36%</b> (96)
Nervous system disorders	<b>19%</b> (49)	<b>19%</b> (51)
Ischemic stroke	<b>5%</b> (13)	<b>0.8%</b> (2)
Hemorrhage intracranial	<b>3%</b> (8)	<b>4%</b> (11)
Cerebral hemorrhage	<b>3%</b> (7)	<b>4%</b> (11)
Hydrocephalus	<b>3%</b> (7)	<b>2%</b> (4)
Neurological decompensation	<b>0.8%</b> (2)	<b>3%</b> (7)
Infections and infestations	<b>16%</b> (43)	<b>11%</b> (28)
Pneumonia	<b>5%</b> (14)	<b>6%</b> (16)
Pneumonia aspiration	<b>5%</b> (14)	<b>3%</b> (7)
Sepsis	<b>2%</b> (6)	<b>0.8%</b> (2)
Cardiac disorders	<b>8%</b> (22)	<b>3%</b> (7)
Myocardial infarction	<b>3%</b> (8)	<b>0.4%</b> (1)
Respiratory, thoracic and mediastinal disorders	<b>6%</b> (17)	<b>5%</b> (12)
Pulmonary embolism	<b>0.8%</b> (2)	<b>3%</b> (7)

### **ANNEXA-I: TEAEs Leading to Death in > 2 Patients**

Safety Set	Andexanet	Usual Care
<b>%</b> (n)	N = 262	N = 265
TEAE leading to death	<b>24%</b> (64)	<b>20%</b> (54)
Nervous system disorders	<b>9%</b> (24)	<b>10%</b> (26)
Cerebral hemorrhage	<b>2%</b> (6)	<b>3%</b> (9)
Hemorrhage intracranial	<b>2%</b> (5)	<b>2%</b> (4)
Ischemic stroke	<b>1%</b> (3)	0
Infections and infestations	<b>7%</b> (18)	<b>6%</b> (15)
Pneumonia	<b>3%</b> (7)	<b>2%</b> (6)
Pneumonia aspiration	<b>3%</b> (7)	<b>2%</b> (5)
Sepsis	<b>1%</b> (3)	<b>0.4%</b> (1)
Respiratory, thoracic and mediastinal disorders	<b>4%</b> (11)	<b>2%</b> (5)
Respiratory failure	<b>2%</b> (4)	<b>2%</b> (4)
Cardiac disorders	<b>3%</b> (8)	<b>0.8%</b> (2)
Cardiac failure	<b>1%</b> (3)	0

#### ANNEXA-I: All-Cause Mortality at 30 Days Similar Between Treatment Groups



## **AESI: Thrombotic Events**

USPI contains boxed warning about risk of thrombotic events with Andexanet

## **Adjudication of Thrombotic Events in ANNEXA-I**

## Multiple sources identified potential thrombotic events for adjudication

- 1. Site reported event of special interest
- 2. Identified by medical review
- 3. Identified by adverse event code matching

Blinded Endpoint Adjudication Committee Followed prespecified charter determined whether events were thrombotic

	Safety Set		Apixaban / Rivaroxaban Subset	
<b>%</b> (n)	Andexanet N = 262	<b>Usual Care</b> N = 265	Andexanet N = 239	Usual Care N = 232
Blinded EAC Adjudicated Thrombotic Events	<b>10.3%</b> (27)	<b>5.7%</b> (15)	<b>10.9%</b> (26)	<b>5.6%</b> (13)

### Difference Between Independently Adjudicated Thrombotic Event Rates and FDA Assessment

Apixaban / Rivaroxaban Subset

Andexanet (N = 239)			
Blinded EAC Adjudicated TEs	10.9% (26)		
Total FDA TE Count	14.6% (35)		
Atrial thrombosis	+1		
Troponin increased	-1		
Ischaemic stroke	+2		
Cerebrovascular accident	+2		
Cerebral infarction	+1		
Pulmonary embolism	+3		
Embolism arterial	+1		

Usual Care (N = 232)			
Blinded EAC Adjudicated TEs	5.6% (13)		
Total FDA TE Count	6.9% (16)		
Sudden cardiac death	+1		
Central venous thrombosis	+1		
Pulmonary embolism	+1		

### **ANNEXA-I: Adjudicated Thrombotic Events**

Safety Set		
<b>%</b> (n)	Andexanet N = 262	Usual Care N = 265
Any thrombotic event	<b>10.3%</b> (27)	<b>5.7%</b> (15)
Ischemic stroke	<b>6%</b> (17)	<b>2%</b> (4)
Myocardial infarction	<b>4%</b> (11)	<b>2%</b> (4)
Arterial systemic embolism	<b>1%</b> (3)	<b>0.8%</b> (2)
Pulmonary embolism	<b>0.4%</b> (1)	<b>2%</b> (6)
Deep vein thrombosis	<b>0.4%</b> (1)	<b>0.8%</b> (2)
Thrombotic event leading to death	<b>2%</b> (6)	<b>0.8%</b> (2)

#### ANNEXA-I: Relationship Between Re-anticoagulation <sup>°</sup> and Thrombotic Events

Safety Set



## ANNEXA-I: Subgroup Analyses to Delineate Update to USPI Informing Use in Patients at High Baseline Thrombotic Risk

Safety Set

Medical History		Andexanet % (n/N)	Usual Care % (n/N)	Higher TE Event Rate	Difference (95% CI)
Cardiac Failure -	Yes	<b>17.4%</b> (8/46)	<b>3.3%</b> (2/61)	· · · · · · · · · · · · · · · · · · ·	<b>14.1%</b> (2.2, 26.0)
	No	<b>8.8%</b> (19/216)	<b>6.4%</b> (13/204)		<b>2.4%</b> (-2.6, 7.5)
Stroke or MI	Yes	<b>12.8%</b> (10/78)	<b>2.5%</b> (2/80)	· · · · · · · · · · · · · · · · · · ·	<b>10.5%</b> (2.1, 18.9)
	No	<b>9.2%</b> (17/184)	<b>7.0%</b> (13/185)		<b>2.2%</b> (-3.4, 7.8)
			-3	30 -20 -10 0 10 20 3	30

Difference in Thrombotic Event Rate 30 Days Post-Randomization (95% CI)

## ANNEXA-I Supports Acceptable Safety Profile of Andexanet In The Setting of Uncontrolled and Life-Threatening Bleeding

- Higher rate of thrombotic events with and exanet
  - Known risk when restoring physiologic coagulation in patients with an underlying thrombotic risk who are bleeding
  - Proposed USPI update informing use in specific patients with high baseline thrombotic risk
- 30-day mortality rates were similar between treatment groups
  - Causes of death in ANNEXA-I are in line with other studies
- Safety profile in ANNEXA-I is consistent with results from the clinical development program and consistent with MoA
- No new safety signals or adverse drug reactions identified



## **Clinical Perspective**

Ashkan Shoamanesh, MD, FRCPC

Marta and Owen Boris Chair in Stroke Research and Care Associate Professor, Medicine Department of Neurology McMaster University

## Treatment of Patients with ICH Requires Rapid Decisions in Emergency Setting to Minimize Risk of Death



**3 cc** ICH Score: 0 0% mortality **42 cc** ICH Score: 3 72% mortality

#### Time is Brain: Therapies that Assist with our Rapid Response in These Settings are Critical



<sup>1.</sup> Adapted from Al-Shahi Salman, 2018

## Reversing Anticoagulation is Only One Aspect in Bundle of Care for Patients with ICH

EMS	CT-Scan	Bundle of Care	In-hospital Care and Secondary Prevention
<ul> <li>Symptom onset</li> <li>Vitals</li> <li>Glucose</li> <li>NIHSS</li> <li>Anticoagulant use (dose, time)</li> <li>eGFR</li> <li>Trauma vs spontaneous</li> </ul>	<ul> <li>Ischemic</li> <li>Hemorrhagic</li> <li>Tumor/cancer</li> <li>Trauma</li> </ul>	<ul> <li>Anticoagulation reversal, agent specific</li> <li>BP control</li> <li>Glucose control</li> <li>Surgical evacuation</li> <li>Normothermia</li> </ul>	<ul> <li>BP management</li> <li>Risk factor management</li> <li>Re-initiation of anticoagulants</li> <li>Infection control measures</li> <li>Early rehabilitation</li> </ul>
			<image/>

**CC-56** 

Ma, 2023; Li, 2024; Greenberg, 2022; Parry-Jones, 2023

#### ANNEXA-I: Primary Efficacy Endpoint Established Hemostatic Benefit of Andexanet vs Usual Care



# Minimizing Hematoma Expansion is Critical Outcome in FXa Inhibitor-Related Life-Threatening Bleeds



## Minimizing Door to Needle Time for ICH Treatment is Key Goal

	ANNEXA-I		
Time (hours), Median (Q1, Q3)	Andexanet N = 224	Usual care N = 228	Get With The Guidelines Stroke Registry (USA, N = 9,492) <sup>1</sup>
Symptom onset to baseline scan	<b>2.3</b> (1.5, 4.0)	<b>2.4</b> (1.4, 3.8)	-
Symptom onset to hospital arrival (door)	-	-	<b>2.6</b> (1.1, 7.0)
Baseline scan to randomization	<b>1.1</b> (0.7, 1.5)	<b>1.2</b> (0.7, 1.7)	-
Door-to-needle	<b>2.1</b> (1.5, 2.9)	<b>2.3</b> (1.7, 3.1)	<b>1.4</b> (1.0, 2.0)*

Door-to-needle time is a predictor of treatment effect

## Andexanet Has Well-Established Safety Profile

- Andexanet increased risk of TE in ANNEXA-I
  - Consistent with label and boxed warning in USPI
  - Similar rate as observed in ANNEXA-4
- TE risk must be assessed in context of management of life-threatening or uncontrolled bleeding event
  - Primary goal is to stop the bleed to limit brain injury
  - Well-equipped to monitor and manage TEs
  - Once patient stabilizes, focus shifts to managing long-term risks
  - Need to reinitiate anticoagulation therapy as soon as medically appropriate

## Reducing Hematoma Expansion is the Primary Goal of Medical Interventions in Acute ICH Setting

- Time is brain!
- ANNEXA-I supports that and example fills an important medical need as an effective, rapid reversal agent for FXa inhibitors
- Balancing reductions in hematoma expansion vs. increase in thrombotic risk indicate overall benefit of andexanet



### **Moderator for Q&A**

#### Matthew Roe, MD, MHS

Cardiologist, Adjunct Professor of Medicine Duke University Medical Center

Vice President, Head of Early CVRM Clinical Development AstraZeneca BioPharmaceuticals

#### Andexanet in Life-Threatening or Uncontrolled Bleeding in Patients Receiving a Direct Oral Factor Xa Inhibitor

#### November 21, 2024

Cellular, Tissue and Gene Therapies Advisory Committee AstraZeneca