

# Overcoming the Safety Challenges of Aldose Reductase Inhibition: Development of AT-001 for Diabetic Cardiomyopathy



### **Disclosures**

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### Definition of Diabetic Cardiomyopathy (DbCM)<sup>1</sup>

- Abnormal cardiac structure and/or performance
  - Resulting from diabetes-associated metabolic alterations
  - In the absence of coronary artery disease (CAD) as well as hypertensive, valvular or congenital heart disorder
- Progresses to overt heart failure (HF)<sup>2,3</sup>

Myocardial dysfunction

Myocardial dysfunction

Diastolic dysfunction, Systolic dysfunction,

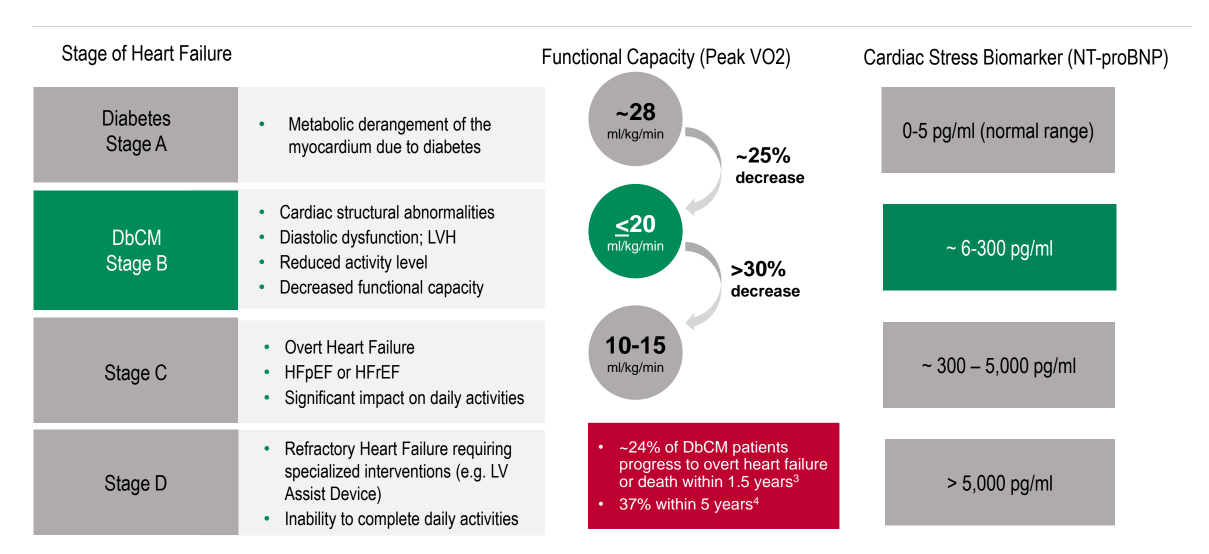
LV hypertrophy & concentric remodeling

Overt Heart Failure

Hospitalization, death

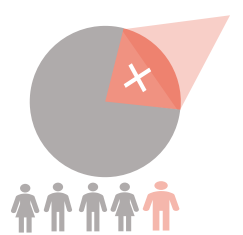


### Diabetic Cardiomyopathy as a Form of Stage B Heart Failure<sup>1-4</sup>





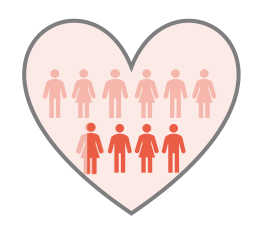
### Diabetic Cardiomyopathy: A High Unmet Medical Need



Approximately, **17-24%** of patients with diabetes have DbCM in the absence of other forms of heart disease. <sup>1,2</sup>

~77 M patients worldwide have DbCM<sup>3</sup>

- ~ 8.0M in North America
- ~ 10.0M in Europe



- ~24% of DbCM patients progress to overt heart failure or death within 1.5 years<sup>4</sup>
- 37% within 5 years<sup>5</sup>

- Patients with diabetes are counseled on HF risk reduction:
  - Lifestyle modification
  - Hypertension
  - Dyslipidemia

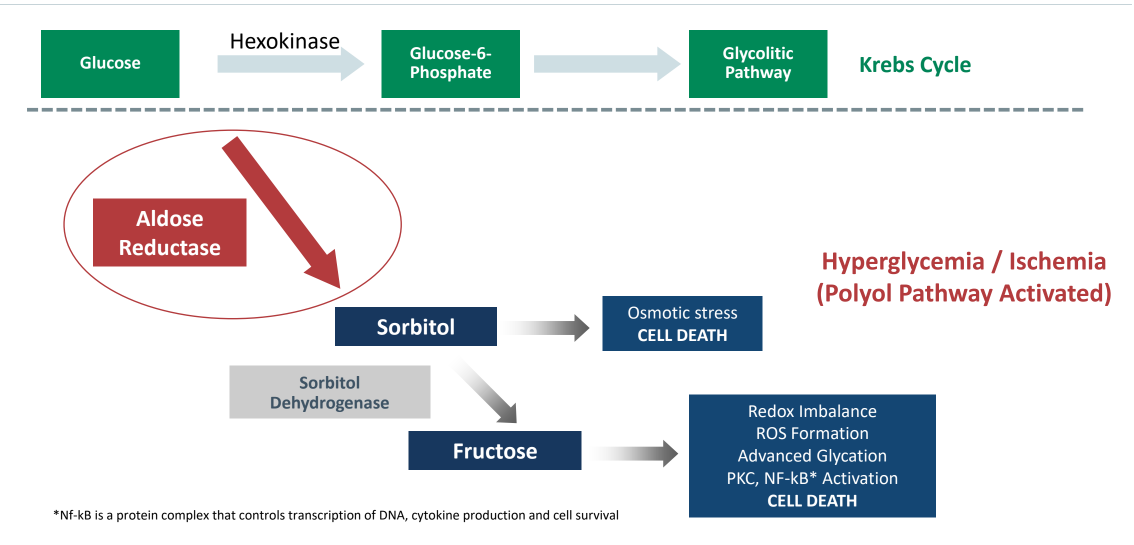
- Hyperglycemia
- Albuminuria

#### No Treatment for DbCM

- No therapies target the metabolic derangement responsible for DbCM and subsequent worsening to overt HF
- Heart Failure treatment is only initiated upon onset of clinical symptomatology (stage C heart failure)



### Pathogenesis of DbCM & Hyperactivation of Polyol Pathway<sup>1,2</sup>





## First Generation Aldose Reductase Inhibitor Zopolrestat (Pfizer)

# Inhibition of Aldose Reductase Clinical Efficacy Competitive Inhibition of Aldehyde Reductase

- First generation Aldose Reductase Inhibitor (zopolrestat) demonstrated clinical efficacy in Diabetic Cardiomyopathy<sup>1</sup>
- Hepatotoxicity was observed in the development program (presumably due to off target competitive binding with Aldehyde Reductase in liver)
- Clinical development was discontinued



(Off-Target) Hepatotoxicity

### AT-001: A Next Generation Highly Selective Aldose Reductase Inhibitor for Treatment of Diabetic Cardiomyopathy

# **AT-001**

- AT-001 was developed through rational drug design, using the geometric parameters of the active site of the Aldose Reductase enzyme determined via X-ray crystallography.
- Optimal target selectivity for Aldose Reductase and minimization of potential off-target activity with Aldehyde Reductase was achieved.
- Aldehyde Reductase plays an important role in detoxification mechanisms in the liver.
   Minimization of off-target activity is critical to ensure safety.

### AT-001 Increased Affinity for Aldose Reductase vs. Zopolrestat

Compound	Structure	IC <sub>50</sub>	MTD in animals	Tissue Penetration (in rats)			
				System ic/ Heart	Nerve	Retina	CNS
AT-001	$O$ $N$ $S$ $CF_3$	30pM	>2,000mg/kg	✓	✓	✓	X
zopolrestat	$O$ $N$ $S$ $CF_3$ $CO_2H$	10nM	100mg/kg	✓	✓	X	Х

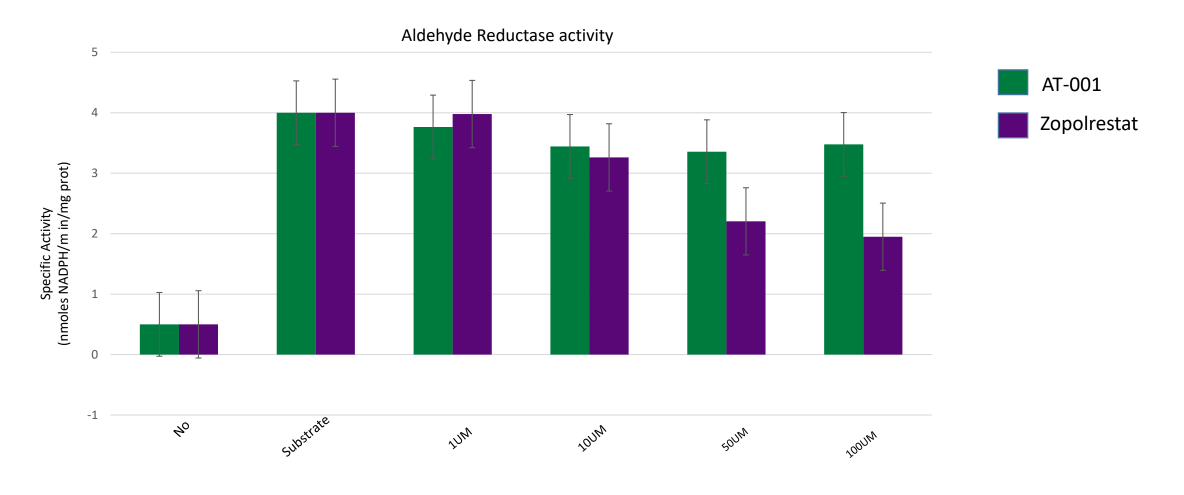


### **No AT-001 Off-Target Binding**

- Eurofins Panlabs Safety Screen Panel (consisting of 87 primary molecular targets including 13 enzyme and 74 binding assays) was used to evaluate potential off target binding activity of AT-001
- No off-target binding activity (defined as ≥50% inhibition or stimulation for biochemical assays) was observed



### **Zopolrestat (But Not AT-001) Inhibits Aldehyde Reductase**





### Conclusions

- AT-001 is logarithmically more potent than zopolrestat in inhibiting Aldose Reductase
- The unique structure and activity of AT-001 provide selectivity for Aldose Reductase and avoid off-target inhibition of Aldehyde Reductase
- The in vitro safety of this agent together with the positive safety data from the phase 1/2 program, support the ongoing pivotal study in DbCM

