

# PEVONEDISTAT (TAK-924): A POTENTIAL NEW TREATMENT FOR HR-MDS AND AML



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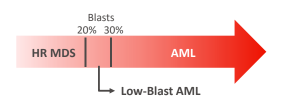
Better Health, Brighter Future

## **BUILDING ON THE TAKEDA ONCOLOGY FOUNDATION IN** Takeda **HEMATOLOGIC MALIGNANCIES** Cell therapies **Next Generation** Type I IFN Novel checkpoints 1/0 MDS/AML **GROWING** Phase 3 **LEADERSHIP** POSITION IN **HEMATOLOGIC MALIGNANCIES** Lymphoma Chronic Myeloid Leukemia **VADCETRIS ICLUSIG**® Improving Patient Outcomes in Multiple Myeloma **VELCADE** NINLARO (ixazomib) capsules

# HIGH RISK MYELODYSPLASTIC SYNDROME (HR-MDS) AND ACUTE MYELOID LEUKEMIA (AML) HAVE LIMITED TREATMENT OPTIONS

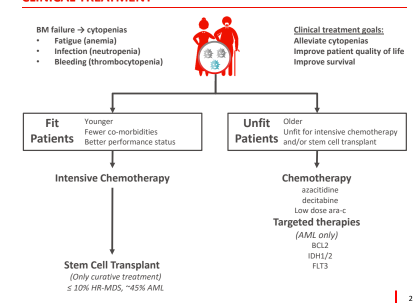


### **CONTINUUM OF HR-MDS AND AML**



- HR-MDS and AML are both rare bone marrowrelated cancers that share foundational biology, clinical features, and genetic mutations\*
- Incidence highest in elderly (>70 years old)
- Overall survival several months to a few years, depending on risk category

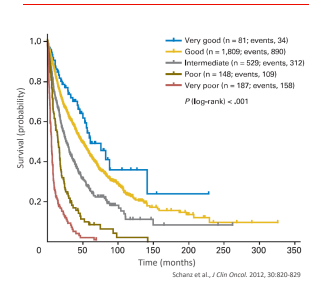
### **CLINICAL TREATMENT**



## **CURRENT STANDARD OF CARE IS INADEQUATE FOR HR-MDS PATIENTS**



### MDS SURVIVAL BY PROGNOSTIC RISK



Median survival ~6 months to 5 years

- No new treatments have been approved for MDS in over a decade
- Transplant ineligible patients treated with first line therapy:
   Median OS = 15mo; 2yr OS rate 35%
- Economic burden is substantial hospitalizations are common among patients and many are transfusion dependent

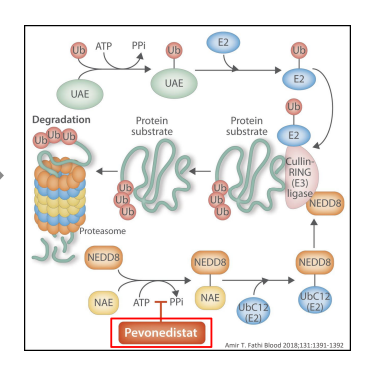
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<sup>\* 30%</sup> of HR-MDS patients progress to AML

## PEVONEDISTAT: A UNIQUE FIRST-IN-CLASS NAE INHIBITOR

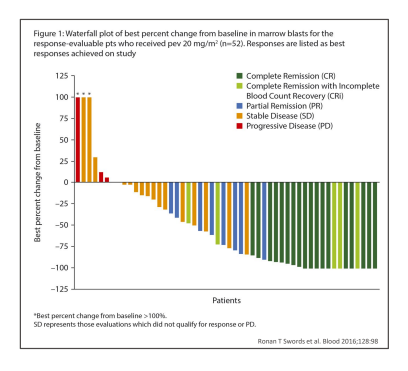


- Pevonedistat is a small molecule inhibitor of NAE (NEDD-8 activating enzyme), a protein involved in the ubiquitin-proteasome system
- NAE acts upstream of the proteasome and catalyzes the first step in the neddylation pathway



# ENCOURAGING RESPONSES IN AML PATIENTS TREATED WITH PEOVNEDISTAT + AZACITIDINE





60% ORR with a trend towards improved survival in secondary AML

Response rates not influenced by AML genetic risk or leukemia burden

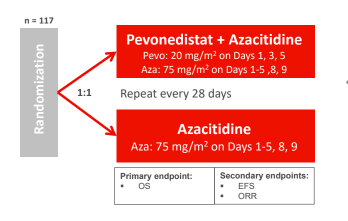


Initial data drove interest to move to registration

# A PHASE 2 STUDY IN HR-MDS TO CONFIRM THE RISK / BENEFIT PROFILE OBSERVED IN AML



Phase 2, Randomized, Open-label, Global, Multicenter Study Comparing Pevonedistat Plus Azacitidine vs. Azacitidine in Patients with Higher-Risk MDS, CMML, or Low-Blast AML

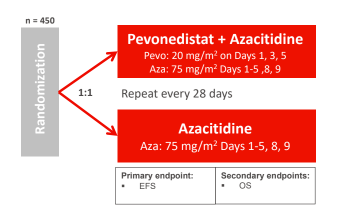


- Mature OS data will be available in November
- Data will be presented in upcoming congress
- Potential approval in FY21\*

# THE PHASE 3 PANTHER STUDY WAS INITIATED AT RISK TO ACCELERATE DEVELOPMENT



Phase 3, Randomized controlled trial of Pevonedistat Plus Azacitidine Versus Single-Agent Azacitidine as First-Line Treatment for Patients with Higher risk-MDS/CMML, or Low-blast AML





- Completed global enrollment 10 months earlier than originally projected\*
- Indicative of demand for new innovative therapies

<sup>\*</sup> Projected approval date assumes filing on Phase 2 data

## **EXPANDING PATIENT-CENTRIC DEVELOPMENT OF PEVONEDISTAT**



## **Continuum of disease**

## **HR-MDS**

Ph2 (P2001)

Ph3 (P3001)

Potential approval in FY21\*

PANTHER

## **NEW STUDIES IN UNFIT AML**

### **Ph3 PEVOLAM**

pevo + aza vs. aza
Currently enrolling patients

Utilizing partnership (PETHEMA) for efficient development

### Ph2 (P2002) Combo

pevo + venetoclax + aza vs. venetoclax + aza Study will open in 2020 Unique MOA and biologic hypothesis to support combination

\* Projected approval date assumes filing on Phase 2 data

## **SUMMARY**



1

Unmet need in Highrisk MDS and AML remain high with few treatment options 2

Pevonedistat is a selective first-in-class inhibitor with potential to be first new therapy in over a decade for HR-MDS

3

The Ph2 HR-MDS trial has reached the updated OS endpoint data readout and the PANTHER Ph3 trial has completed global enrollment