

# Sierra Oncology

Developing Transformative Therapies for Rare Cancers

January 2022



## SAFE HARBOR STATEMENT

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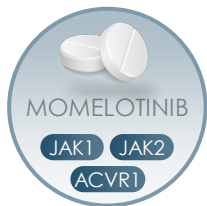
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## Momelotinib is the potential treatment of choice for myelofibrosis patients with anemia

- Profile has demonstrated anemia improvement, symptom and spleen control, without platelet decreases

## Less than 1 month from pivotal Phase 3 MOMENTUM data

- High probability of success trial; FDA approval and launch expected <18 months

## ~\$3B addressable market\* in anemic MF patients in the US

- ~15K prevalent patients with anemia in the US

## Could become the cornerstone of future combinations in myelofibrosis

- Momelotinib + SRA515 combination will provide proof-of-concept
- Targeting patient subsets where the unmet medical need remains

## Current market cap of ~\$506M

- ~22M FD shares outstanding<sup>(1)</sup>
- ~\$97.1M cash as of 9/30/21<sup>(2)</sup>



# Myelofibrosis: Disease Overview



**Myelofibrosis (MF):** a bone marrow cancer

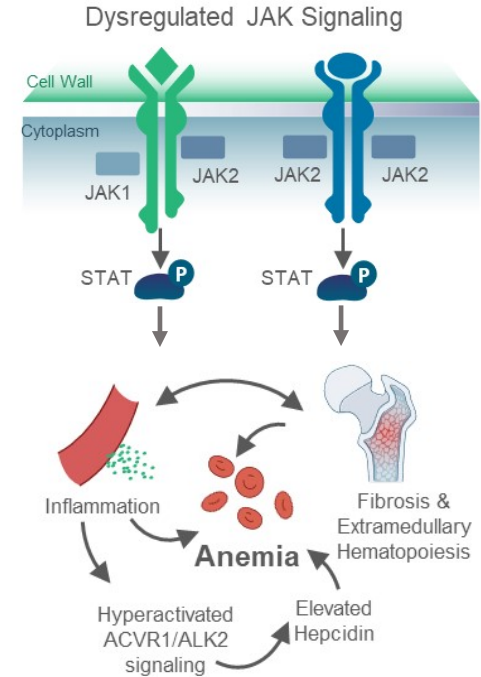
- Caused by constitutive activation of JAK-STAT signaling
- Inflammation and fibrosis impair red blood cell production



**Common manifestations** of disease include constitutional symptoms, enlarged spleen and **progressive anemia**



**Current treatments:** JAK inhibitors are the mainstay option for intermediate and high-risk patients



# Anemia and Hepcidin Predict Poor Survival in Myelofibrosis

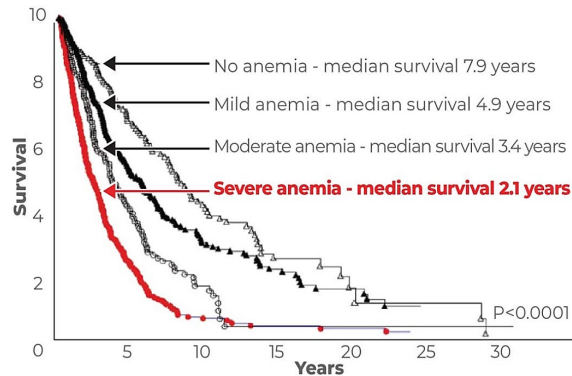
**Anemia of inflammation**  
driven by elevated  
hepcidin

**Elevated hepcidin** inhibits  
iron transport and iron  
homeostasis

Anemia and elevated  
hepcidin are **negative  
prognostic indicators**

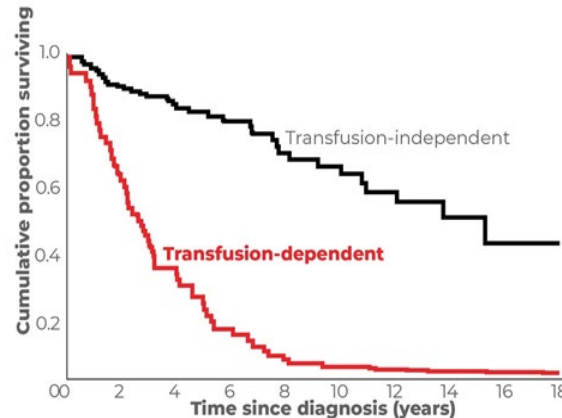
**New therapies should  
provide anemia benefits  
in addition to symptom,  
spleen benefits**

## Anemia Predicts Poor Survival in MF



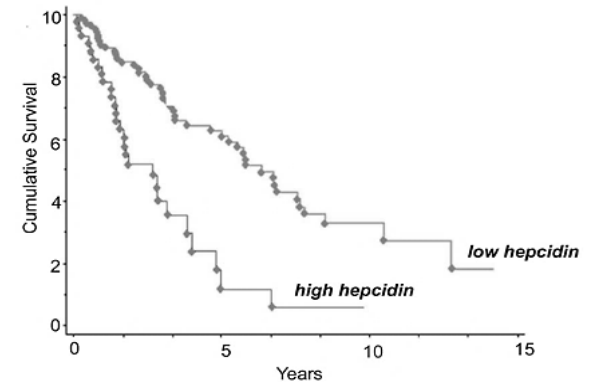
Nicolosi et al. *Leukemia*. 2018

## Transfusion Dependency Predicts Poor Survival in MF



Elena et al. *Haematologica*. 2011

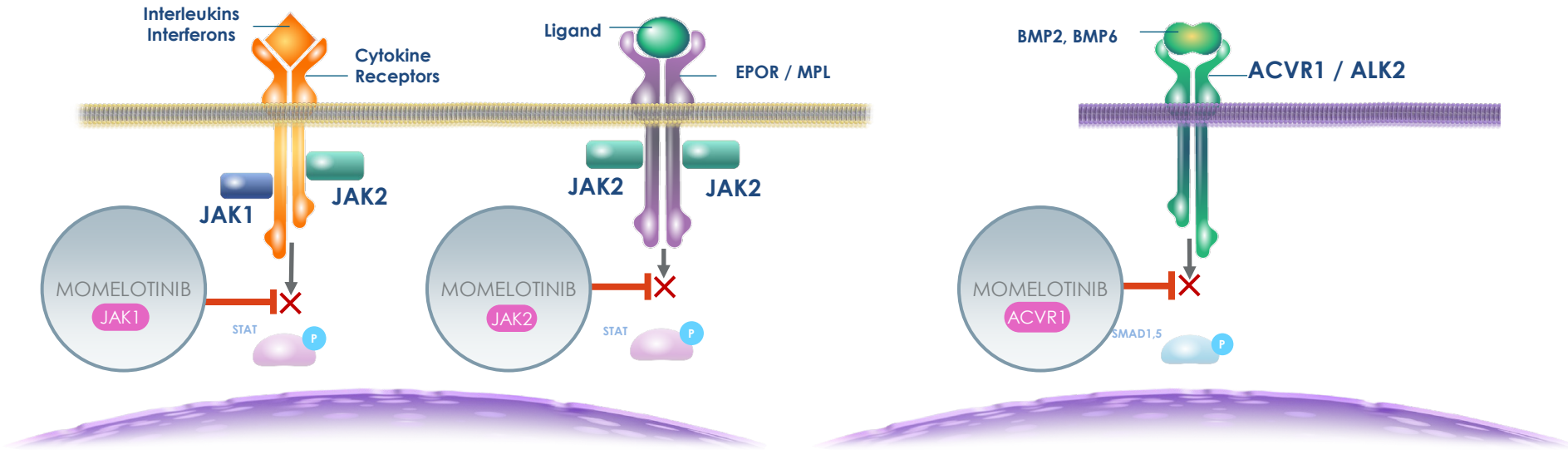
## Elevated Hepcidin Predicts Poor Survival in MF



Pardanani et al. *Am. J. Hematol.* 2013



# Momelotinib Inhibits JAK1, JAK2 and ACVR1/ALK2



Hyperactive JAK-STAT signaling is driving the disease in myelofibrosis

Preclinical and clinical studies suggest that the clinical anemia benefits of momelotinib result from suppression of ACVR1/ALK2-mediated hepcidin production

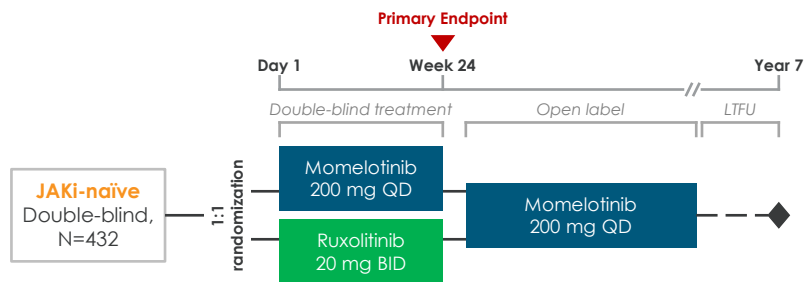
**Momelotinib Inhibits all Three Disease Drivers, Potentially Improving Splenomegaly and Symptoms of Myelofibrosis While Maintaining or Improving Hemoglobin**

# Completed Phase 3 Studies SIMPLIFY-1 and 2

## SIMPLIFY-1

### 1<sup>st</sup>-Line Population

JAK inhibitor naïve



Goal:	Non-Inferiority	
Endpoints at Week 24:	MMB	RUX
SRR $\geq 35\%$ (primary)*	27%	29%
Symptom score reduction $\geq 50\%$	28%	42%
TI for $\geq 12$ weeks	67%	49%

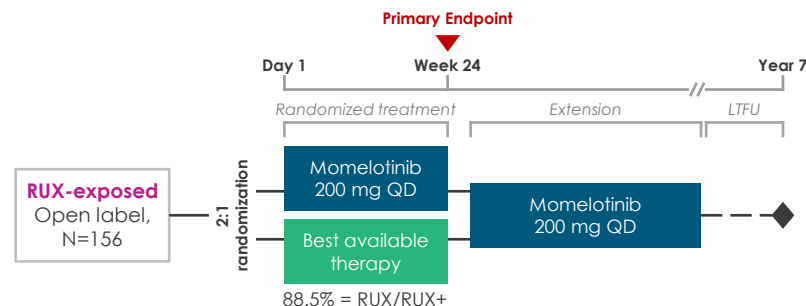
\*Met endpoint  
*Journal of Clinical Oncology*, 2017 35(34):3844



## SIMPLIFY-2

### 2<sup>nd</sup>-Line Population

Prior ruxolitinib complicated by hematologic toxicity



Goal:	Superiority	
Endpoints at Week 24:	MMB	RUX/BAT
SRR $\geq 35\%$ (primary)	7%	6%
Symptom score reduction $\geq 50\%$	26%	6%
TI for $\geq 12$ weeks	43%	21%

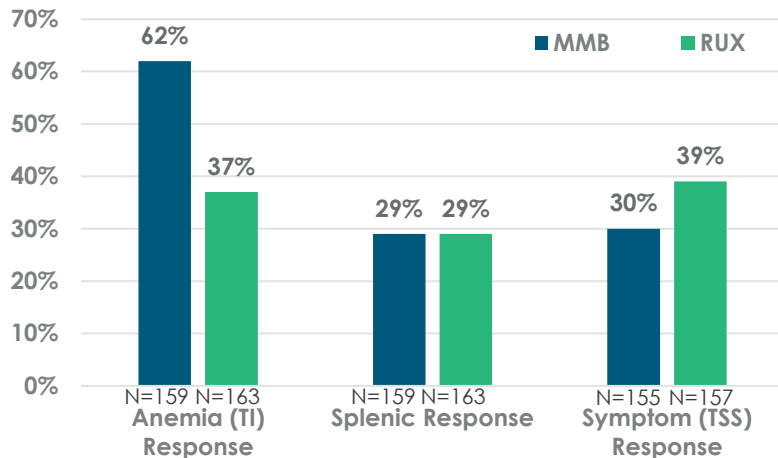
*The Lancet Haematology*, 2018 5(2): 7

# Comparative Efficacy MMB vs RUX/BAT in Anemic Patients

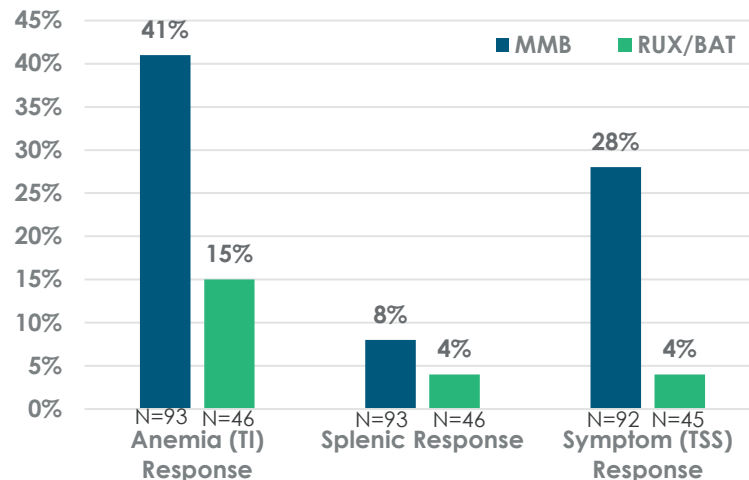
MMB's anemia benefits are accompanied by similar splenic and symptomatic response rates in SIMPLIFY-1 and significantly better symptom control relative to BAT in SIMPLIFY-2

## Week 24 Response Rates

SIMPLIFY-1: Patients with Hgb  $\leq 12$  g/dL at Baseline

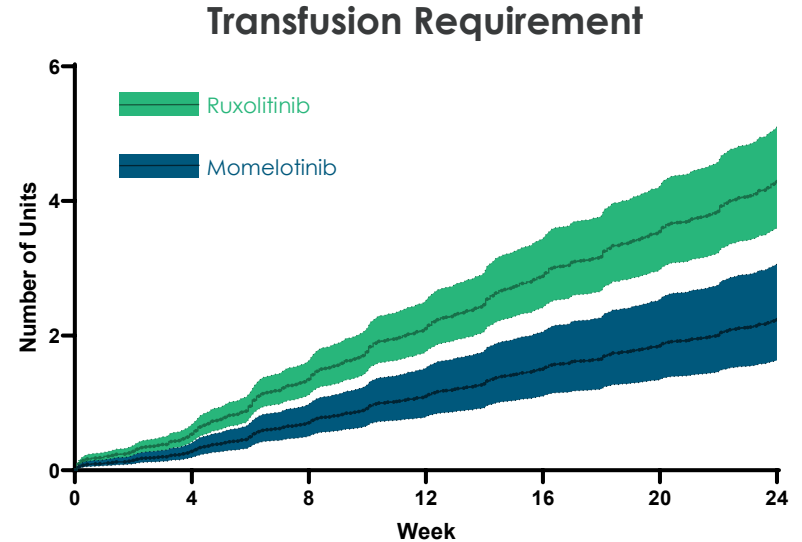
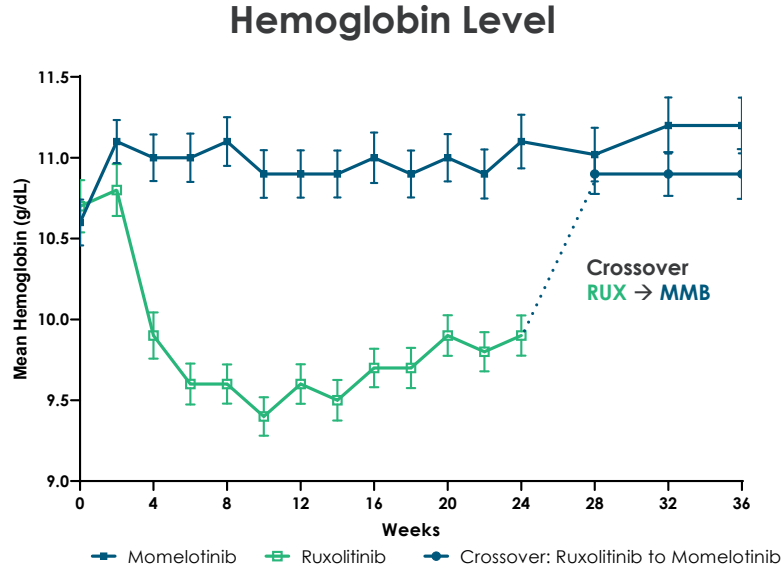


SIMPLIFY-2: Patients with Hgb  $\leq 12$  g/dL at Baseline





# SIMPLIFY-1 Highlighted MMB's Unique Impact on Hemoglobin and Transfusions



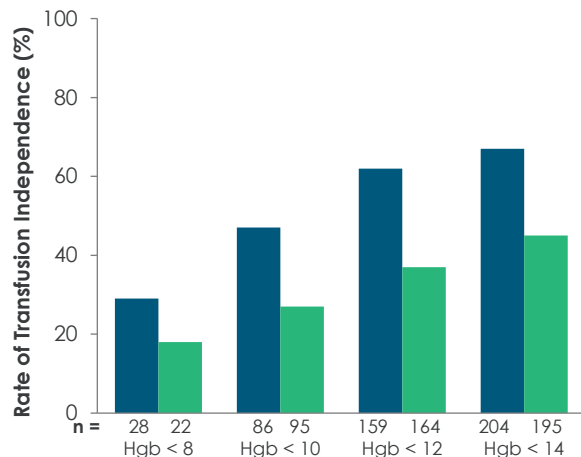
Transfusion requirement was ~half for MMB vs. RUX

**Momelotinib Demonstrated an Increase in Hemoglobin and a Decreased Transfusion Requirement vs. Ruxolitinib**

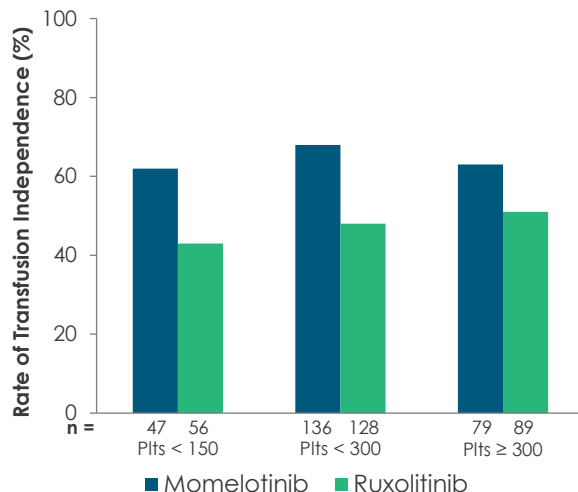


# Transfusion Independence is Achieved for Anemic Patients and all Patients, Irrespective of Baseline Platelets or Transfusion Status

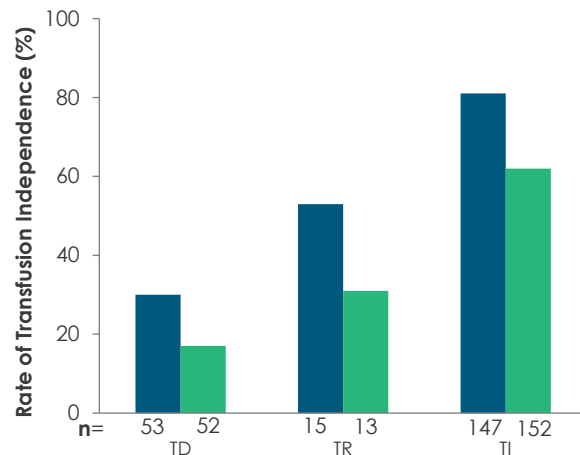
## SIMPLIFY-1 Anemic Patients W24 TI-Response by Baseline Hgb



## SIMPLIFY-1 W24 TI-Response by Baseline PLT



## SIMPLIFY-1 W24 TI-Response by Baseline Transfusion Status



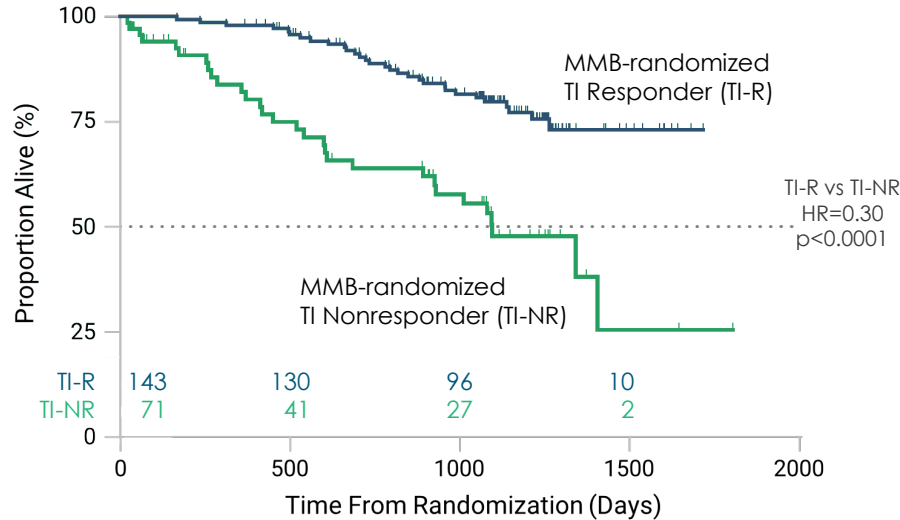
**The W24 TI-R Rate in S1 Was Higher in Patients Randomized to MMB vs RUX, Irrespective of the Degree of Baseline Anemia, or the Baseline PLT Count or Transfusion Status**

Week 24 Transfusion Independence Response (TI-R): no RBC transfusion within ≥ 12 weeks immediately prior to Week 24, with Hgb ≥ 8 g/dL  
Transfusion Dependent (TD): ≥ 4 units of RBCs or Hgb level, ≤ 8 g/dL in the 8 weeks prior to randomization  
Transfusion Independent (TI): absence of RBC transfusions and no Hgb < 8 g/dL in the 12 weeks prior to randomization  
Transfusion Requiring (TR): neither TD nor TI  
Hgb = hemoglobin, MMB = momelotinib, PLTs = platelets, RUX = ruxolitinib, TD = transfusion dependent, TI = transfusion independent, TR = transfusion requiring  
Kiladjian J.J. et.al. *European Hematology Association*, June 2021, poster EP1081; Virtual.

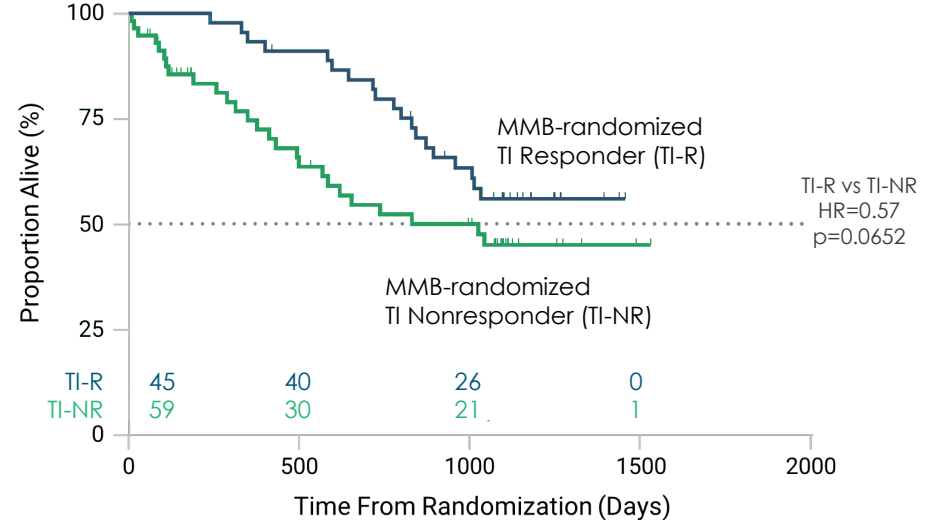


# Transfusion Independence (TI) with Mometotinib is Associated with Improved Overall Survival

## SIMPLIFY-1



## SIMPLIFY-2



**Achieving or Maintaining TI Predicted Better Survival in Patients Treated with Mometotinib – The Goal of Achieving TI Should Become an Important Driver of Treatment Decisions**

Week 24 TI response = no RBC transfusion for ≥ 12 weeks immediately prior to Week 24, Hgb level ≥ 8 g/dL.  
Mesa, R. et.al. *European Hematology Association*, June 2021, oral presentation S202; Virtual.



# Safety and Tolerability from the SIMPLIFY Phase 3 Trials

- Safety generally similar for momelotinib, ruxolitinib in the 24-week double-blind period
  - Anemia and thrombocytopenia were more common in the ruxolitinib arm
  - Nausea was more common with momelotinib, as was the early withdrawal rate in S-1
- Tolerability persists with extended treatment
  - No evidence of long-term cumulative toxicity observed
- Safety profile enables long duration of dosing
  - Several patients from early trials have now received >10 years of continuous momelotinib therapy
  - Many patients from SIMPLIFY-1 and -2 continue to receive momelotinib

## SIMPLIFY-1

### Frequent TEAEs<sup>1</sup> by PT

	Randomized Treatment Period	
	MMB (N=214)	RUX (N=216)
Pts with any TEAE, n (%)	198 (92.5%)	206 (95.4%)
Diarrhea	39 (18.2%)	43 (19.9%)
Anemia	31 (14.5%)	81 (37.5%)
Thrombocytopenia	40 (18.7%)	63 (29.2%)
Nausea	34 (15.9%)	8 (3.7%)
Fatigue	31 (14.5%)	26 (12.0%)

## S-1 Extended

### Most Frequent TEAEs<sup>1</sup> by PT

	Extended duration MMB Final Safety Analysis (N=411)
Pts with any TEAE, n (%)	397 (96.6%)
Diarrhea	99 (24.1%)
Anemia	93 (22.6%)
Thrombocytopenia	94 (22.9%)
Nausea	85 (20.7%)
Fatigue	84 (20.4%)

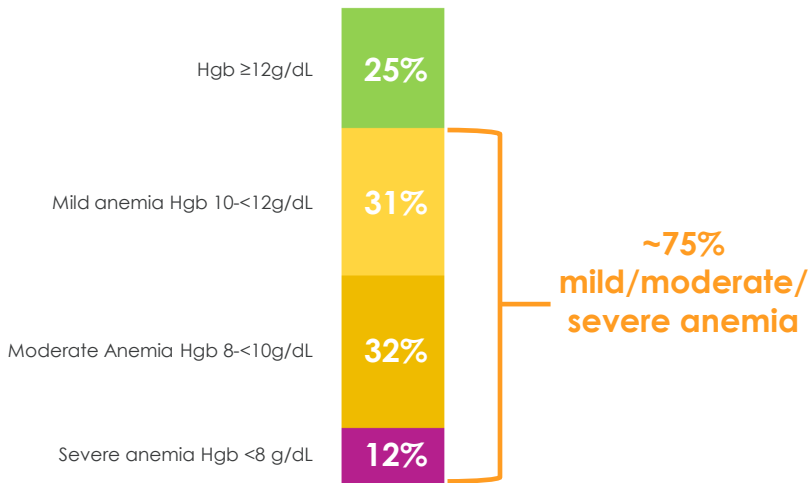
<sup>1</sup> TEAEs occurring in >20% pts in the "Overall exposed to MMB" population including the 214 subjects receiving blinded momelotinib and 197 additional subjects who received momelotinib after cross-over from ruxolitinib



# Anemia Could Become the Primary Driver of JAKi Choice

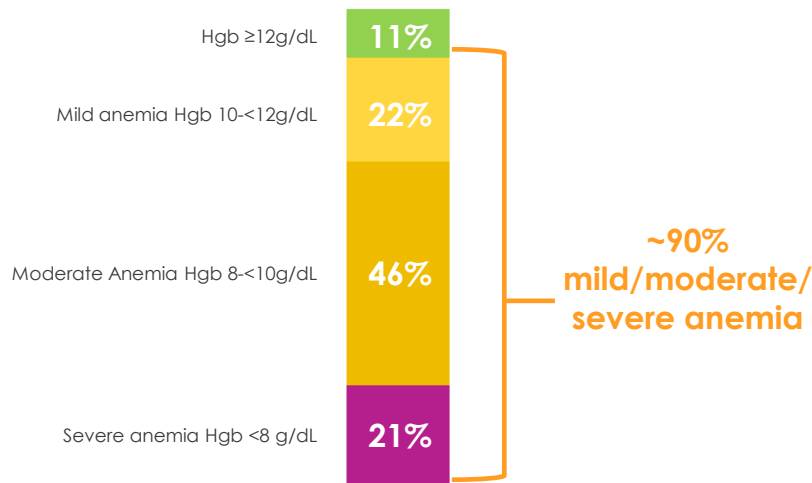
- Momelotinib could become the JAKi of choice for anemic myelofibrosis patients
  - In S-1 at baseline, ~75% of anemic with 30% requiring transfusions
  - In Rux-experienced patients, 90% are anemic with more than two thirds require transfusions
- A differentiated JAKi could allow hem/oncs to transition patients early in their treatment journey
- Upside potential with a broad label that allows for use in JAKi-naïve patients

**JAKi Naïve Patients (S-1) – at Baseline**



**~30% Require Transfusions**

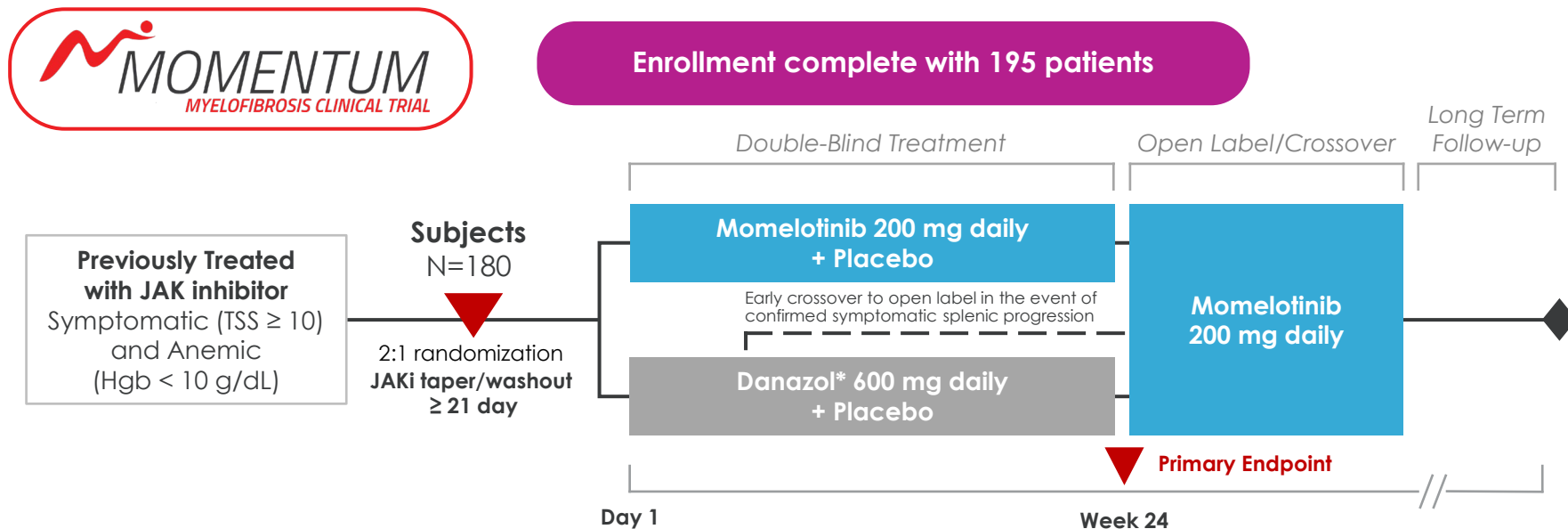
**RUX Experienced Patients (S-2) – at Baseline**



**~70% Require Transfusions**



# Pivotal Phase 3 'MOMENTUM' Study: Topline Results Expected by February 2022



## Primary Endpoint

- Total symptom score (TSS) response rate at Week 24

## Secondary Endpoints

- Transfusion independence (TI) rate at Week 24
- Splenic response rate (SRR) at Week 24

\*Danazol was selected as an appropriate comparator given its use to ameliorate anemia in MF patients, as recommended by NCCN, ESMO guidelines.



# Momelotinib + SRA515 Combinations Could Expand the Myelofibrosis Opportunity



## Myelofibrosis landscape is evolving with multiple combination studies ongoing

- BET inhibition has shown initial proof-of-concept with disease-modifying potential
- Unlike other BET inhibitors, SRA515 has a novel bivalent binding mode
  - A stronger connection on the cellular level leads to improved potency
- SRA515 has favorable PK, PD and safety profile when dosed as monotherapy and in combination



## Clinical validation of MMB + BET combination

- SRA515 has synergistic preclinical efficacy in combination with diverse agents
  - Best-in-class potential
- As a non-myelosuppressive inhibitor of JAK1, JAK2, and ACVR1, MMB is an ideal combination partner for novel agents
- SRA515+MMB has the potential to improve outcomes in patients with MF

**Momelotinib + SRA515 may provide the opportunity for longer and more durable responses for myelofibrosis patients**

# MMB + SRA515 May be the “Winning” Combination



- ✓ SRA515 potentially the **most potent BETi** with selective target inhibition
  - Novel bivalent binding mode; Allows for maintained dosing durability
- ✓ MMB may be the best combination agent as **only JAKi with anemia benefit in MF**
  - Novel JAK1, JAK2 and ACVR1/ALK2 MOA does not add to myelosuppression of BETi
- ✓ Sierra **wholly owns both compounds**, allowing for data-driven development approach
- ✓ Distinct advantage of extensive MMB clinical experience with almost **1,000 MF patients dosed**
- ✓ Can create **intelligent development plan** due to both internally- and externally-derived data sources





# Sierra Oncology Clinical Program

Sierra's pipeline affords numerous combination opportunities, with SOC and with other investigational agents

Program	Indication	Phase 1	Phase 2	Phase 3	Registration
<b>Momelotinib monotherapy</b>	Myelofibrosis	Topline results expected by February 2022			
<b>Momelotinib + SRA515*</b>	Myelofibrosis	Planned for H1 2022			
<b>SRA515 monotherapy and/or SRA515 + SRA737</b>	Heme malignancies**	Finalizing Design			
<b>SRA515 monotherapy and/or SRA515 + SRA737</b>	Solid tumor in combo with SOC**	Finalizing Design			
<b>SRA737 + IO/gemcitabine</b>	Solid tumors	Finalizing Design			

\*Formerly AZD5153

\*\*Opportunities currently under consideration include co-operative trials

