Sierra Oncology

Developing Transformative Therapies for Rare Cancers
January 2022



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Sierra Oncology Overview

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⁽¹⁾
- $\sim 97.1 M cash as of $9/30/21^{(2)}$



Myelofibrosis: Disease Overview



Myelofibrosis (MF): a bone marrow cancer

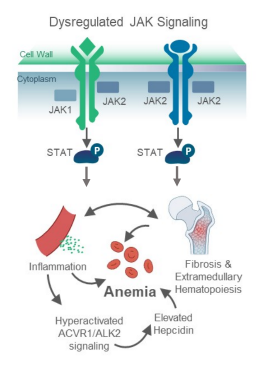
- Caused by constitutive activation of JAK-STAT signaling
- Inflammation and fibrosis impair red blood cell production



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

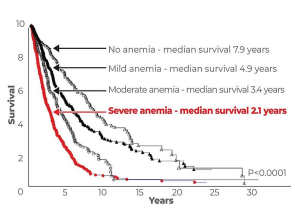
Anemia of inflammation driven by elevated hepcidin

Elevated hepcidin inhibits iron transport and iron homeostasis

Anemia and elevated hepcidin are *negative prognostic indicators*

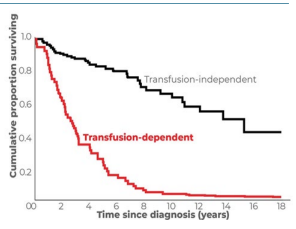
New therapies should provide anemia benefits in addition to symptom, spleen benefits

Anemia Predicts Poor Survival in MF



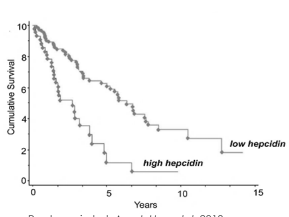
Nicolosi et al. Leukemia. 2018

Transfusion Dependency Predicts Poor Survival in MF



Elena et al. Haematologica. 2011

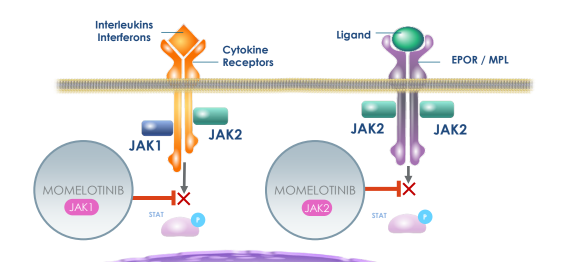
Elevated Hepcidin Predicts Poor Survival in MF

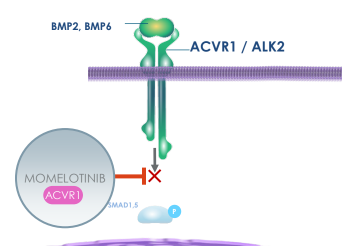


Pardanani et al. Am. J. Hematol. 2013



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

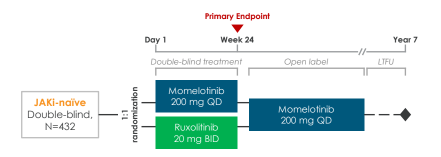
Asshoff, M. et. al. Blood. 2017;129(13):1823-1830. Oh, S. et al. Blood Advances. 2020;4(18):4282-4291.



Completed Phase 3 Studies SIMPLIFY-1 and 2

SIMPLIFY-1

1st-Line Population JAK inhibitor naïve



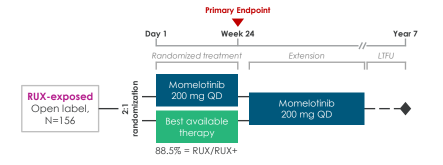
Goal:	Non-Inferiority		
Endpoints at Week 24:	MMB	RUX	
SRR ≥35% (primary)*	27%	29%	
Symptom score reduction ≥50%	28%	42%	
TI for ≥ 12weeks	67%	49%	

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

SIMPLIFY-2

2nd-Line Population

Prior ruxolitinib complicated by hematologic toxicity



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

The Lancet Haematology, 2018 5(2): 7

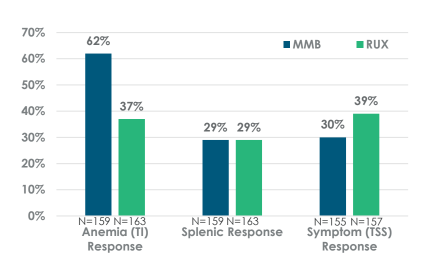


Comparative Efficacy MMB vs RUX/BAT in Anemic Patients

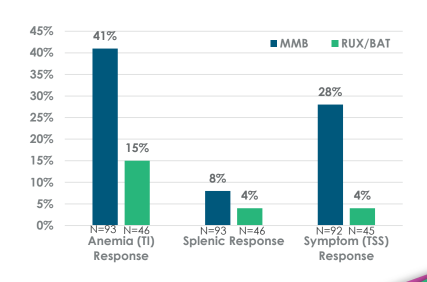
MMB's anemia benefits are accompanied by similar splenic and symptomatic response rates in SIMPLIFY-1 and significantly better symptom control relative to BAT in SIMPLIFY-2

Week 24 Response Rates

SIMPLIFY-1: Patients with Hgb ≤12 g/dL at Baseline

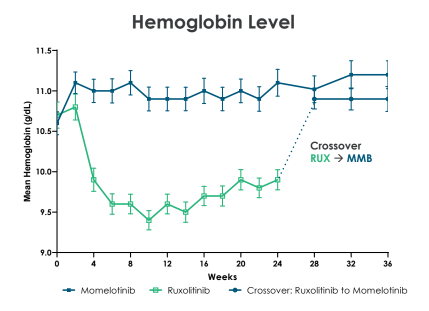


SIMPLIFY-2: Patients with Hgb < 12 g/dL at Baseline

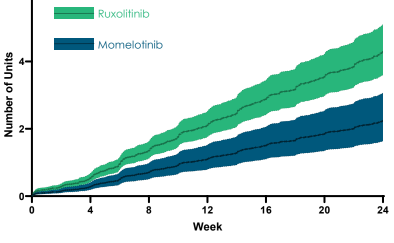




SIMPLIFY-1 Highlighted MMB's Unique Impact on Hemoglobin and Transfusions



Transfusion Requirement Ruxolitinib



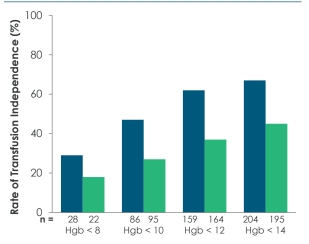
Transfusion requirement was ~half for MMB vs. RUX

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

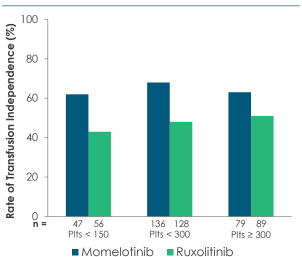


Transfusion Independence is Achieved for Anemic Patients and all Patients, Irrespective of Baseline Platelets or Transfusion Status

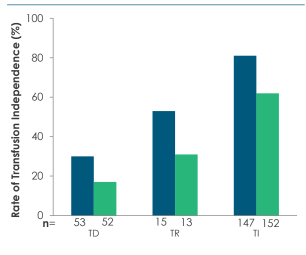
SIMPLIFY-1 Anemic Patients
W24 TI-Response by Baseline Hgb



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 \geq 12 weeks immediately prior to Week 24, with Hgb \geq 8 g/dL Transfusion Dependent (TD): \geq 4 units of RBCs or Hgb level, \leq 8 g/dL in the 8 weeks prior to randomization

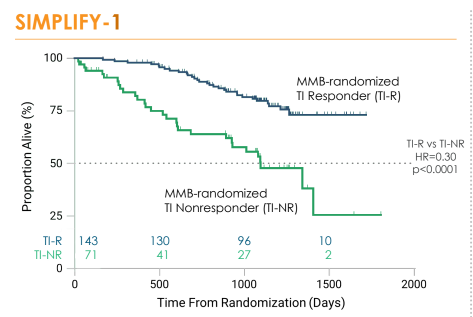
 $Transfusion \, Independent \, (TI): absence of RBC \, transfusions \, and \, no \, Hgb < 8 \, g/dL \, in \, the \, 12 \, weeks \, prior \, to \, randomization \, transfusions \, and \, random \, ran$

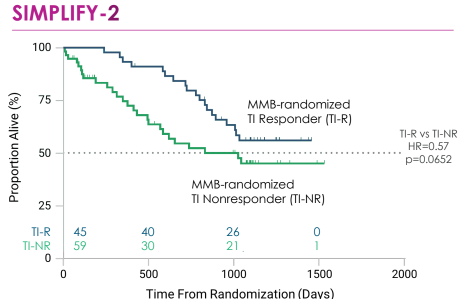
Transfusion Requiring (TR): neither TD nor TI

HgB = hemoglobin, MMB = momelotinib, Plts = platelets, RUX = ruxolitinib, TD = transfusion dependent, TI = transfusion independent, TR = transfusion requiring Kiladjian JJ. et.al. European Hematology Association, June 2021, poster EP1081; Virtual.



Transfusion Independence (TI) with Momelotinib is Associated with Improved Overall Survival





Achieving or Maintaining TI Predicted Better Survival in Patients Treated with Momelotinib – 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 \$202; 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	Randomized Treatment Period			
Frequent TEAEs ¹ by PT	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	Extended duration MMB		
Most Frequent TEAEs ¹ by PT	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%)		

1 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

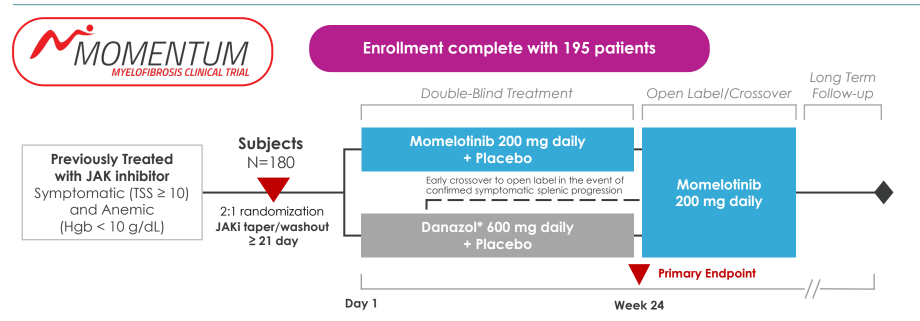


Anemia Could Become the Primary Driver of JAKi Choice

- 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



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 bode
 - 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



- 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 internallyand 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
Momelotinib monotherapy	Myelofibrosis	Topline	results expected by Febru	ary 2022	I
Momelotinib + SRA515*	Myelofibrosis	Planned for H1 2022			
SRA515 monotherapy and/or SRA515 + SRA737	Heme malignancies**	Finalizing Design			
SRA515 monotherapy and/or SRA515 + SRA737	Solid tumor in combo with SOC**	Finalizing Design			
SRA737 + IO/gemcitabine	Solid tumors	Finalizing Design			

^{*}Formerly AZD5153

^{**}Opportunities currently under consideration include co-operative trials