Sustained-release ruxolitinib: Findings from a phase 1 study in healthy subjects and a phase 2 study in patients with myelofibrosis

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Abstract
Ruxolitinib is the first Janus kinase (JAK) inhibitor approved for the treatment of myelofibrosis, where its efficacy is often associated with cytopenia. It is possible that the severity of cytopenia is in part driven by \( C_{\text{max}} \). A once-daily sustained-release (SR) formulation of ruxolitinib was therefore developed to decrease the \( C_{\text{max}}/C_{\text{min}} \) ratio relative to twice-daily immediate-release (IR) ruxolitinib. An SR formulation was identified based on pharmacokinetic evaluation in a phase 1 study in healthy adults (N = 9). This was followed by an open-label phase 2 study in patients with myelofibrosis (N = 41). Ruxolitinib SR treatment was well tolerated with blood cell counts relatively unchanged through week 16. In terms of efficacy, 7 patients (17.1%) had clinical improvement and 33 (80.5%) had stable disease. While this study has raised the possibility of an increased therapeutic index for ruxolitinib via an SR formulation, further studies are required to validate the hypothesis.

KEYWORDS
clinical trials, experimental, formulation, oncology, pharmacokinetics

1 | INTRODUCTION

A twice-daily (BID) immediate-release (IR) formulation of ruxolitinib is approved for the treatment of patients with intermediate-risk or high-risk myelofibrosis (MF), including primary MF (PMF), postpolycythemia vera MF (PPV-MF), and postessential thrombocythemia MF (PET-MF).1 Phase 3 randomized clinical trials demonstrated that ruxolitinib IR reduced splenomegaly and improved MF-related symptoms and quality of life compared with controls.2,3 Furthermore, data from these trials strongly suggested that ruxolitinib prolonged survival relative to placebo or best available therapy.4,5 However, in some patients, ruxolitinib can cause thrombocytopenia and anemia, typically during the first 8 to 12 weeks of therapy, which may result in dose reductions or treatment discontinuation.1

A once-daily (QD) sustained-release (SR) formulation of ruxolitinib was developed in an effort to generate an efficacious treatment option while limiting the risk and severity of cytopenias by decreasing the maximal plasma exposure to ruxolitinib. A QD formulation could also provide a more convenient treatment regimen for patients with MF. Across a variety of treatment settings and disease states, patient adherence to prescribed treatment is optimized by limiting the number of required daily doses.6,7

This report presents data from 2sequential studies that evaluated SR formulations of ruxolitinib. A phase 1 study in healthy subjects (INCB 18424-139) compared the bioavailability of ruxolitinib SR and IR to
determine a suitable formulation and dose for evaluation in patients with MF. A phase 2 study (INCB 18424-260; NCT01340651) subsequently evaluated the pharmacokinetic properties and clinical activity of ruxolitinib SR in patients with MF.

## METHODS

### Ethics

Both studies were conducted in accordance with the Declaration of Helsinki and Good Clinical Practice: Consolidated Guideline approved by the International Conference on Harmonisation. The respective clinical study protocols, amendments, informed consent documents, and other appropriate study-related documents were reviewed and approved by independent ethics committees/institutional review boards. Written informed consent was obtained from all participants.

### Phase 1 study: healthy subjects

Details about enrolled subjects, study design, study endpoints, and statistical analyses for the phase 1 study of healthy subjects can be found in the Supplemental Appendix.

### Phase 2 study: patients with myelofibrosis

#### Patients

All enrolled participants were adults (aged ≥18 years) diagnosed with PMF, PPV-MF, or PET-MF for which treatment was indicated as per physician assessment. Study participants had a life expectancy ≥6 months and a spleen length ≥5 cm below the costal margin (determined by palpation).

#### Study design and dosing

INCB 18424-260 was a single-arm, open-label phase 2 trial of ruxolitinib SR tablets QD. Study visits occurred at screening; baseline; day 1; weeks 2, 4, 8, 12, 16, 20, and 24; and every 12 weeks thereafter. All patients began treatment with 25 mg ruxolitinib SR QD. After 8 or 12 weeks, the dose level of ruxolitinib could be titrated to 50 mg SR QD for inadequate efficacy. To address potential toxicity arising from the increased dose, the protocol was amended to include an optional titration to ruxolitinib 25 mg SR alternating with 50 mg SR every other day (QOD), depending on platelet count; this amendment occurred after the study began (patients entering the study before the amendment did not have this option). Efficacy was considered inadequate if patients had <40% reduction from baseline in palpable spleen length at the week 8 or 12 study visit.

At week 16, all patients transitioned to ruxolitinib IR BID, with the starting dose depending on platelet count: Patients with a platelet count ≥200 × 10⁹/L initiated 20 mg IR BID, those with a platelet count 100 to <200 × 10⁹/L initiated 15 mg IR BID, and those with a platelet count 75 to <100 × 10⁹/L initiated 10 mg IR BID. All patients with platelet counts between 50 and <75 × 10⁹/L were subject to a mandatory dose reduction to ruxolitinib IR 5 mg BID. Further treatment with ruxolitinib was withheld from patients with platelet counts <50 × 10⁹/L. Doses were restarted or increased after platelet counts or absolute neutrophil count (ANC) levels recovered to acceptable levels. Patients had the option to remain on ruxolitinib IR treatment until ruxolitinib IR tablets became commercially available or until the last patient completed 36 weeks of treatment, whichever occurred earlier.

### Study endpoints

The primary study endpoints were safety/tolerability and overall response (OR). Safety/tolerability assessments included monitoring adverse events (AEs), vital signs, and clinical laboratory data. Changes from baseline in platelet count over time and the proportion of patients with grade 3/4 thrombocytopenia were also analyzed. To assess OR, the proportion of patients with International Working Group-Myelofibrosis Research and Treatment (IWG-MRT) responses was analyzed at each study visit and designated as clinical response, stable disease, or progressive disease. Although the IWG-MRT criteria include responses designated as complete remission (CR) and partial remission (PR), postbaseline bone marrow samples were not collected to confirm CR versus PR.

Secondary endpoints included changes from baseline in spleen volume, Modified Myelofibrosis Symptom Assessment Form (MFSAF) Total Symptom Score (TSS), and pharmacokinetic assessments. Changes from baseline in spleen volume were measured by magnetic resonance imaging, and spleen length was measured by palpation. The proportion of patients with a ≥35% reduction in spleen volume from baseline and a ≥50% reduction in MFSAF TSS from baseline were assessed at week 16. Assessed pharmacokinetic parameters included minimum plasma concentration (Cmin), maximum plasma concentration (Cmax), time to Cmax (tmax), area under the concentration time curve (AUC) from time 0 to 24 hours (steady state), AUC from time 0 to last measurable concentration (AUCt, last) (single dose), terminal elimination half-life (t1/2), oral-dose clearance (CL/F), and oral-dose volume of distribution (V/F).

### Statistical analyses

Descriptive summaries were included for continuous and categorical variables. Unless otherwise stated, all CIs were 95% CIs, unadjusted for multiplicity. The safety population was used for all safety analyses; the intent-to-treat population was used for efficacy analyses. Pharmacokinetic data were analyzed by population pharmacokinetic analysis using pharmacokinetic- evaluable subjects.

## RESULTS

### Phase 1 study: healthy subjects

#### Subject disposition and demographics

Nine healthy adults (6 men and 3 women) were enrolled in the phase 1 study and received 25 mg ruxolitinib IR. Eight of the 9 subjects remained on-study to receive treatment with 2 different ruxolitinib SR formulations (SR-1 and SR-2). The median (range) age of study participants was 27 (18-53) years. Eight subjects (88.9%) were white; 1 (11.1%) was African American.
Pharmacokinetic simulation indicated that ruxolitinib SR-1 and SR-2 could provide higher steady-state mean $C_{\text{max}}$ values compared with ruxolitinib IR (6.5-fold and 11.4-fold, respectively) and lower steady-state mean $C_{\text{max}}$ values (69% and 61%). The SR-2 formulation was ultimately selected for further development because of a slightly higher relative bioavailability compared with the SR-1 formulation (Table 1).

### 3.2 Phase 2 study: Patients with myelofibrosis

#### 3.2.1 Patient disposition and characteristics

For weeks 1 to 16 of the phase 2 study, patients received 25 mg ruxolitinib SR QD; at week 16, all patients transitioned to 25 mg ruxolitinib IR BID. Of the 41 patients enrolled, 39 (95.1%) completed through week 16, and 24 (58.5%) completed through week 24. Reasons for study withdrawal before week 16 included AEs ($n = 1$) and transferring to treatment with a commercial product ($n = 1$). Reasons for withdrawal from weeks 16 to 24 included transferring to treatment with a commercial product ($n = 11$), consent withdrawn ($n = 2$), and AEs ($n = 2$).

Baseline demographics, disease characteristics, and laboratory values are presented in Table 2. Forty-one adult patients (median [range] age, 68 [50-81] years) with PMF (78.0%), PPV-MF (17.1%), or PET-MF (4.9%) enrolled in this study. A majority of patients were men (65.9%) and were white (97.6%).

Overall, 38 patients (92.7%) were transfusion-independent and 22 (53.7%) were designated intermediate-2 risk according to International Prognostic Scoring System risk category; 19 (46.3%) were designated high risk. A Janus kinase (JAK) 2 mutation was present in 61.0% of patients, and the median spleen volume at baseline was 2592.7 cm$^3$, which is consistent with massive splenomegaly.

For most patients, platelet counts were within normal limits at baseline. The median hemoglobin level of patients was below the lower limit of normal ($130 \times 10^9$/L for men and $120 \times 10^9$/L for women); 37 (90.2%) had decreased hemoglobin of at least grade 1 at baseline. The median ANC was above the upper limit of normal ($7.9 \times 10^9$/L). The median ANC was above the upper limit of normal ($7.9 \times 10^9$/L).

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### Table 1 Phase 1 study of healthy subjects: pharmacokinetic parameters

| Parameter | IR (n = 9) | SR-1 (n = 8) | SR-2 (n = 8) | P Value | Bioavailability, % (90% CI)$^a$ |
|-----------|-----------|-------------|-------------|---------|-------------------------------|
| $C_{\text{max}}, \text{nM}$ | 1100 (332) | 333 (76.1) | 394 (126) | <.0001 | 30.4 (25.4-36.4) |
| $t_{\text{max}}, \text{h}$ | 0.9 (0.5) | 2.4 (1.0) | 2.9 (1.6) | .0003 | NA |
| $C_{12h}, \text{nM}$ | 45.6 (38.1) | 121 (46.8) | 104 (43.2) | NA | NA |
| $C_{\text{max}}/C_{12h}$ | 40 (24) | 3.0 (1.0) | 4.7 (3.1) | NA | NA |
| $t_{\text{el}}, \text{h}$ | 2.8 (0.7) | 5.3 (1.8) | 6.1 (2.1) | <.0001 | NA |
| area under the concentration-time curve from time 0 to last measurable concentration; $C_{\text{max}}$, maximum plasma concentration; $C_{12h}$, plasma concentration at 12 hours after dosing; $C_{\text{max}}/C_{12h}$, ratio of the maximum plasma concentration to the plasma concentration at 12 hours after dosing; $t_{\text{el}}$, elimination half-life; $t_{\text{max}}$, time to maximum plasma concentration.

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$^a$Geometric mean relative bioavailability.

$^b$P values from a crossover analysis of variance of log-transformed data.
3.2.2 Safety/tolerability

All enrolled patients were in the safety-evaluable population (n = 41). Through week 16, the median duration of exposure to ruxolitinib SR was 113.0 days, with an average daily dose of 26.1 mg.

Through week 16, 32 of 41 patients (78.0%) receiving ruxolitinib SR experienced a TEAE. The most common TEAEs through week 16 were diarrhea (19.5%), peripheral edema (17.1%), and thrombocytopenia/decreased platelet count (17.1%). Other TEAEs occurring in ≥5% of patients were bone pain (9.8%) and anemia, asthenia, nausea, pruritus, and sinusitis (7.3% each). A grade ≥3 TEAE was observed in 7 of 41 patients (17.1%), including 1 patient (2.4%) who experienced grade 3 thrombocytopenia. Treatment-emergent cytopenias occurring after week 16 included thrombocytopenia (4.9%) and anemia (2.4%).

Through week 16, 1 patient (2.4%) experienced TEAEs leading to study withdrawal (day 8 gastritis and gastrointestinal hemorrhage [neither related to study treatment] with a platelet count of 305 × 10⁹/L). The gastrointestinal hemorrhage resolved following hospitalization.

### TABLE 2 Phase 2 study of patients with myelofibrosis: baseline demographics and disease characteristics, intent-to-treat population

| Characteristic                                      | Ruxolitinib (N = 41) |
|-----------------------------------------------------|----------------------|
| Median (range) age, years                           | 68.0 (50.0-81.0)     |
| Sex, n (%)                                          |                      |
| Male                                                | 27 (65.9)            |
| Female                                              | 14 (34.1)            |
| Race, n (%)                                         |                      |
| White                                               | 40 (97.6)            |
| Asian                                               | 1 (2.4)              |
| IPSS risk category, n (%)                           |                      |
| High, ≥3 factors                                    | 19 (46.3)            |
| Intermediate, 2 factors                             | 22 (53.7)            |
| ECOG performance status, n (%)                      |                      |
| 0                                                    | 3 (7.3)              |
| 1                                                    | 26 (63.4)            |
| 2                                                    | 11 (26.8)            |
| 3                                                    | 1 (2.4)              |
| MF subtype, n (%)                                   |                      |
| Primary                                             | 32 (78.0)            |
| Post-polycythemia vera                              | 7 (17.1)             |
| Post-essential thrombocytetmia                      | 2 (4.9)              |
| Prior hydroxyurea use, n (%)                         |                      |
| Yes                                                 | 18 (43.9)            |
| No                                                  | 23 (56.1)            |
| Transfusion status, n (%)                           |                      |
| Independent                                         | 38 (92.7)            |
| Dependent                                           | 3 (7.3)              |
| Median (range) palpable spleen size below costal margin, cm | 18.0 (7.0-30.0)     |
| Median (range) spleen volume, cm³a                  | 2592.7 (697.1-5926.1) |
| Mean (SD) total symptom score                       | 21.4 (12.03)         |
| JAK2 mutation, n (%)                                |                      |
| Positive                                            | 25 (61.0)            |
| Negative                                            | 7 (17.1)             |
| Missing                                             | 9 (22.0)             |
| Median (range) percentage V617F of positive history JAK2 mutationb | 79.0 (44.0-96.0) |
| Median (range) baseline laboratory values, ×10⁹/La  |                      |
| Platelet count                                       | 217.5 (108.0-1030.0) |
| Hemoglobin                                          | 107.0 (68.0-135.0)   |
| Absolute neutrophil count                           | 12.5 (1.3-35.3)      |
| Lymphocyte count                                     | 1.3 (0.21-4.54)      |

Abbreviations: ECOG, Eastern Cooperative Oncology Group; IPSS, International Prognostic Scoring System; MF, myelofibrosis.

an = 40.
bn = 25.
(gastritis was ongoing). After week 16, 2 patients (4.9%) withdrew due to TEAEs (anemia, n = 1; cardiac arrest, n = 1).

The median levels of platelets, hemoglobin, neutrophils, and leukocytes over time are shown in Figure 2A. Most patients (56.1%) maintained their baseline platelet count grade through week 16. Baseline hemoglobin levels were generally maintained throughout the study. Most patients (95.1%) had neutrophil counts in the normal range at baseline, and 92.3% of these patients maintained normal counts through week 16. Similarly, 95.1% of patients had leukocyte counts in the normal range at baseline, and 84.6% maintained normal counts through week 16.

From study initiation through week 16, 6 patients (14.6%) had SAEs; diarrhea and pneumonia were the most common (n = 2 [4.9%] each), while clostridium difficile colitis, dehydration, gastroenteritis, gastrointestinal hemorrhage, generalized edema, hip fracture, hypotension, patella fracture, and pulmonary embolism occurred in 1 patient each (2.4%). None of the SAEs observed through week 16 were deemed related to study treatment. Two patients (4.9%) died after week 16 while receiving ruxolitinib IR; 1 patient died of cardiac arrest, and 1 patient progressed to acute myeloid leukemia. The investigators deemed these events unrelated to ruxolitinib treatment.

### 3.2.3 Week 16 efficacy

Efficacy endpoints at week 16 are summarized in Table 3. From baseline to week 16, 7 patients (17.1%) had clinical improvement, 33 (80.5%) had stable disease, and 1 (2.4%) had progressive disease.

At week 16, the median (range) percentage change from baseline in spleen volume was −21.7% (−64.6% to 43.6%), and 11 patients (26.8%) had a ≥35% reduction in spleen volume from baseline. Changes from baseline in palpable spleen length decreased steadily through week 12 (Figure 2B); the median (range) percentage change from baseline to week 16 was −25.8% (−100% to 50.0%).

The median (range) percentage change in MFSAF TSS from baseline to week 16 was −48.6% (−100% to 12.7%); 18 patients (43.9%) had a ≥50% reduction from baseline at week 16.

### 3.2.4 Pharmacokinetics

At weeks 4 and 12, the median $t_{\text{max}}$ after the first dose of ruxolitinib was 2.0 hours (Table 4). Plasma concentrations of ruxolitinib declined in a multiphasic fashion (geometric mean terminal-phase disposition $t_{1/2}$, 5.44–8.90 hours). Geometric mean CL/F ranged from 20.6 to 29.6 L/hour, and geometric mean $V_z/F$ ranged from 226 to 305 L.

Steady-state plasma concentrations of ruxolitinib at weeks 4 and 12 are shown in Figure 3. For ruxolitinib doses of 25 to 50 mg SR QD, the mean steady-state $C_{\text{max}}$ and AUC values increased less than proportionally to dose. The geometric mean peak-to-trough ratios for SR ranged from 9.3 (week 12; 25 mg SR QD) to 20.4 (week 4; 25 mg SR QD), which exceeded the estimated value of 6.8 based on the pharmacokinetic simulation from the phase 1 study of healthy subjects.

Sex-based differences were noted in the clearance and volume of distribution of ruxolitinib SR; women had lower CL/F (18.3 vs 20.6 L/hour) and higher $V_z/F$ (226 vs 305 L).
and apparent oral‐dose Vz/F (218 vs 304 L) than men. A sex‐based difference was also observed with ruxolitinib IR (1.25‐fold reduction of CL/F in women vs men).

**TABLE 4**  Phase 2 study of patients with myelofibrosis: sustained‐release ruxolitinib pharmacokinetic parameters at weeks 4 and 12

| Parameters, Mean ± SD (Geometric Mean) | Ruxolitinib 25 mg SR QD | Ruxolitinib 50 mg SR QD | Ruxolitinib 25 mg SR QODa |
|---------------------------------------|-------------------------|-------------------------|--------------------------|
|                                       | Week 4 (n = 36)         | Week 12 (n = 18)        | Week 12 (n = 11)          | Week 12 (n = 5) |
| Cmin, nM                              | 32.9 ± 42.0 (NC)        | 59.1 ± 49.3 (NC)        | 70.0 ± 79.4 (NC)          | 51.3 ± 94.9 (NC) |
| Cmax, nM                              | 411 ± 221 (371)         | 464 ± 223 (419)         | 730 ± 334 (664)           | 407 ± 142 (387) |
| PT ratio                              | 40.1 ± 55.4 (20.4)      | 11.7 ± 9.21 (9.30)      | 28.5 ± 37.5 (15.8)        | 45.9 ± 63.4 (16.84) |
| tmax, h                               | 7.21 ± 3.67 (6.37)      | 8.46 ± 3.62 (7.66)      | 6.12 ± 2.98 (5.44)        | 16.3 ± 17.8 (8.90) |
| AUC0‐t, nM·h                          | 2110 ± 1830 (2290)      | 2590 ± 1320 (2290)      | 4260 ± 1730 (3880)        | 2370 ± 1060 (2180) |
| AUC0‐0, nM·h                          | 3320 ± 2280 (2750)      | 4520 ± 2390 (3960)      | 6670 ± 3410 (5870)        | 4010 ± 2460 (3440) |
| CL/F, L/h                             | 34.8 ± 18.7 (29.6)      | 23.6 ± 12.9 (20.6)      | 32.2 ± 20.2 (27.8)        | 27.5 ± 15.6 (23.4) |
| Vz/F, L                               | 350 ± 212 (289)         | 287 ± 190 (237)         | 255 ± 156 (226)           | 470 ± 540 (305) |
| Median (range)                        | 2.0 (1.0‐6.0)           | 2.0 (0.5‐8.0)           | 3.0 (1.0‐8.0)             | 2.0 (0.5‐3.0) |

At week 4, there were 36 patients in the 25 mg SR QD group, 2 in the 5 mg IR BID group, and 2 who were not included (1 because of withdrawal from the study and 1 because of a dose interruption). At week 12, there were 18 patients in the 25 mg SR QD group, 11 in the 50 mg SR QD group, 7 in the 25/50 mg QOD group (5 patients took 25 mg SR and 2 took 50 mg SR that day), and 2 in the 5 mg IR BID group.

Abbreviations: AUC0‐t, area under the concentration‐time curve at the last measurable concentration; AUC0‐0, area under the concentration‐time curve from time 0 to the last measurable concentration; BID, twice daily; Cmax, maximum plasma concentration; Cmin, minimum plasma concentration; CL/F, oral dose clearance; IR, immediate release; NC, not calculated; PT, peak‐trough; QD, once daily; QOD, every other day; SR, sustained release; tmax, time to maximum plasma concentration; t1/2, elimination half‐life; Vz/F, volume of distribution.

aThe 25 mg SR QOD regimen was 25/50 mg SR QD with the 25 mg dose on the day of plasma concentration collection. The 25/50 mg QOD group that received 50 mg on the pharmacokinetic sampling day (n = 2) was not included in the table because of the small number of patients in this group.

4 | DISCUSSION

Pharmacokinetic simulations based on the single‐dose phase 1 study results in healthy subjects demonstrated higher steady‐state mean Cmin values and lower steady‐state mean Cmax values with ruxolitinib SR formulations compared with ruxolitinib IR. In fasted subjects, ruxolitinib SR provided extended absorption of ruxolitinib, with lower maximal exposure compared with ruxolitinib IR without compromising relative oral bioavailability. The ruxolitinib SR‐2 formulation was selected for further development because of a slightly higher relative bioavailability of 87% compared with SR‐1.

In the phase 2 study of patients with MF, the pharmacokinetic characteristics of ruxolitinib SR were similar to those observed in the phase 1 study of healthy subjects. Ruxolitinib SR clearance and volume of distribution were lower in women compared with men. Although smaller in magnitude than was observed with ruxolitinib SR, a similar difference was also observed between women and men receiving ruxolitinib IR. The reason for this difference remains unclear; however, it could not be explained by differences in body weight alone. No other covariates were significant predictors for pharmacokinetic parameters.

Patients receiving ruxolitinib SR exhibited reductions in spleen size compared with baseline, as well as improvement in symptoms as
assessed by the MFSAF version 2.0. Ruxolitinib SR was associated with stable hemoglobin and platelet levels, with decreased rates of thrombocytopenia and anemia compared to ruxolitinib IR; safety results were otherwise similar to those reported previously for ruxolitinib IR in patients with MF.4,5

Improvements in spleen size with ruxolitinib SR at week 16 were smaller than those reported for ruxolitinib IR in the pivotal COMFORT studies of patients with MF.2,3 The proportion of patients treated with ruxolitinib who had a ≥35% reduction in spleen volume from baseline was 26.8% at week 16 in the current phase 2 study compared with 41.9% at week 24 or weeks 24/48 in the COMFORT-I and COMFORT-II studies, respectively.2,3

Because ruxolitinib is a balanced JAK1/JAK2 inhibitor, it is unlikely that any potential differences in efficacy between the SR and IR formulations were attributable to JAK enzyme specificity. The variance is most likely due to the differences in peak, average, and minimum inhibition of JAK/signal transducer and activator of transcription signaling. However, an evaluation of isoform specificity was beyond the scope of our analysis.

FIGURE 3 Phase 2 study of patients with myelofibrosis: steady-state plasma concentrations of ruxolitinib at weeks 4 (A) and 12 (B). BID, twice daily; IR, immediate release; QD, once daily; QOD, every other day; SR, sustained release. Data are mean ± SE. Two patients were not included (1 because of withdrawal from the study and 1 because of a dose interruption)

5 | STUDY LIMITATIONS

Given the small size of the phase 2 study population, comparisons of efficacy between dose groups could not be adequately assessed. Direct comparisons of the SR and IR formulations were not planned in this phase 2 proof-of-concept study, and post hoc comparisons are not feasible because the SR and IR formulations were administered sequentially without a washout period. In addition, several patients dropped out of the study after the week 16 visit before ruxolitinib IR was initiated.

A QOD dosing regimen that was available to some patients in the phase 2 study (25 mg SR QOD/50 mg SR QOD) was not available to the first several patients in the study. As such, dose titration was inconsistent and generally more challenging for the first few patients. Limited dosing options (ie, use of 1 25 mg tablet size) may have precluded effective dose titration and achievement of full clinical benefit for some patients. Finally, the phase 2 study was potentially limited by the exclusion of patients who were intolerant of ruxolitinib IR but may have derived clinical benefit from ruxolitinib SR.

6 | CONCLUSIONS

The current study results suggest that the higher peak inhibitions achieved with ruxolitinib IR may have incremental contributions to efficacy compared with ruxolitinib SR. However, there was a trend toward a lesser anemia effect with ruxolitinib SR, which could be beneficial for some patients with MF. Collectively, these data suggest that the development of effective SR formulations of ruxolitinib for treating patients with MF is feasible. However, further long-term studies with multiple ruxolitinib SR dosage strengths will be required to adequately compare efficacy and safety outcomes to those observed with ruxolitinib IR.
AUTHOR CONTRIBUTIONS
All authors participated in writing the manuscript and analyzing the study data. SV, SY, and SE-V participated in designing the study research. SV participated in performing the study research.

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CONFLICT OF INTEREST/DISCLOSURE
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REFERENCES
1. JAKAFI® [package insert]. Wilmington, DE, USA: Incyte Corporation; 2017.
2. Verstovsek S, Mesa RA, Gotlib J, et al. A double-blind, placebo-controlled trial of ruxolitinib for myelofibrosis. N Engl J Med. 2012;366(9):799-807.
3. Harrison C, Kiladjian JJ, Al-Ali HK, et al. JAK inhibition with ruxolitinib versus best available therapy for myelofibrosis. N Engl J Med. 2012;366(9):787-798.
4. Verstovsek S, Mesa RA, Gotlib J, et al. Efficacy, safety, and survival with ruxolitinib in patients with myelofibrosis: results of a median 3-year follow-up of COMFORT-I. Haematologica. 2015;100(4):479-488.
5. Cervantes F, Vannucchi AM, Kiladjian JJ, et al. Three-year efficacy, safety, and survival findings from COMFORT-II, a phase 3 study comparing ruxolitinib with best available therapy for myelofibrosis. Blood. 2013;122(25):4047-4053.
6. Saini SD, Schoenfeld P, Kaulback K, Dubinsky MC. Effect of medication dosing frequency on adherence in chronic diseases. Am J Manag Care. 2009;15(6):e22-e33.
7. Claxton AJ, Cramer J, Pierce C. A systematic review of the associations between dose regimens and medication compliance. Clin Ther. 2001;23(8):1296-1310.
8. Tefferi A, Barosi G, Mesa RA, et al. International working group (IWG) consensus criteria for treatment response in myelofibrosis with myeloid metaplasia, for the IWG for myelofibrosis research and treatment (IWG-MRT). Blood. 2006;108(5):1497-1503.
9. Shi JG, Chen X, McGee RF, et al. The pharmacokinetics, pharmacodynamics, and safety of orally dosed INCB018424 phosphate in healthy volunteers. J Clin Pharmacol. 2011;51(12):1644-1654.
10. Cervantes F, Dupriez B, Pereira A, et al. New prognostic scoring system for primary myelofibrosis based on a study of the international working Group for Myelofibrosis Research and Treatment. Blood. 2009;113(13):2895-2901.

SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section at the end of the article.

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