Impact of Individual Comorbidities on Survival of Patients with Myelofibrosis

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Simple Summary: The coexistence of cancer with other chronic conditions has substantial implications for treatment decisions and outcomes for both neoplasms and chronic disease. Reports have demonstrated the impact of comorbidities on survival in different hematologic disorders. Myelofibrosis (MF) guidelines do not consider the complex interrelations between MF and comorbidity. Several works have shown how MF patients have a wide variety and high burden of comorbidities and demonstrated that the comorbidity burden was significantly associated with an unfavorable impact on survival. These previous studies about comorbidity on MF are retrospective and consider the cumulative rather than individual comorbidity burden. The influence of individual comorbidities on outcome in MF patients has not been studied. We sought to identify the comorbidities in MF patients at diagnosis and to assess the influence of those different comorbidities on survival. Considering them individually may contribute to the personalization of MF management and optimizing outcomes.

Abstract: The comorbidity burden is an important risk factor for overall survival (OS) in several hematological malignancies. This observational prospective study was conducted to evaluate the impact of individual comorbidities on survival in a multicenter series of 668 patients with primary myelofibrosis (PMF) or MF secondary to polycythemia vera (PPV-MF) or essential thrombocytopenia (PET-MF). Hypertension (hazard ratio (HR) = 4.96, \( p < 0.001 \)), smoking (HR = 5.08, \( p < 0.001 \)), dyslipidemia (HR = 4.65, \( p < 0.001 \)) and hepatitis C virus (HCV) (HR = 4.26, \( p = 0.015 \)) were most adversely associated with OS. Diabetes (HR = 3.01, \( p < 0.001 \)), pulmonary disease (HR = 3.13, \( p < 0.001 \)) and renal dysfunction (HR = 1.82, \( p = 0.037 \)) were also associated with an increased risk of death.
Multivariate analysis showed that pulmonary disease (HR = 2.69, p = 0.001), smoking (HR = 3.34, p < 0.001), renal dysfunction (HR = 2.08, p = 0.043) and HCV (HR = 11.49, p = 0.001) had a negative impact on OS. When ruxolitinib exposure was included in the model, the effect of each comorbidity on survival was modified. Therefore, individual comorbidities should be taken into account in determining the survival prognosis for patients with MF.

**Keywords:** myelofibrosis; comorbidities; survival; prognosis

1. **Introduction**

Myelofibrosis (MF), either primary (PMF) or evolving after essential thrombocythemia MF (PET-MF) or polycythemia MF (PPV-MF), is a Philadelphia chromosome-negative myeloproliferative neoplasm (MPN) characterized by the mobilization of clonal hematopoietic cells from the fibrotic bone marrow to extramedullary organs, mainly the spleen and liver [1,2].

The estimated incidence of PMF ranges from 0.1 to 1.5 cases per 100,000 individuals per year, with a peak incidence in the sixth decade of life. The risk of progression to MF in polycythemia vera or essential thrombocythemia at 15 years is 6–14% and 4–11%, respectively [3–6].

About 90% of patients with MF present mutations in the JAK2, CALR or MPL genes, which activate the JAK/STAT signaling pathway, resulting in increased levels of inflammatory cytokines that may trigger constitutional symptoms [7–9]. Disease manifestations include progressive anemia, splenomegaly caused by extramedullary hematopoiesis and constitutional symptoms such as fatigue, itching, night sweats, low grade fever, weight loss and bone pain [10,11].

In clinical practice, treatment options include supportive care, cytoreductive agents or JAK inhibitors [12]. The only curative option for MF is allogeneic hematopoietic stem cell transplant (HSCT), but this procedure is restricted to a minority of patients with MF due to its significant morbidity and mortality [13,14]. Various prognostic scoring systems have been developed to predict the survival of MF patients [15–22]. However, comorbidities have not been included as risk factors in any of these prognostic models [23,24] despite their influence on survival in other hematological malignancies such as chronic myeloid leukemia [25], chronic lymphocytic leukemia [26], myelodysplastic syndromes [27], and acute myeloid leukemia [28] or HSCT [29].

Previous studies have shown that the coexistence of cancer and other chronic conditions has substantial implications for treatment decisions and outcomes for both neoplasms and chronic disease [30]. MF is a disabling condition mainly affecting elderly people [4–6]. Comorbidities are more prevalent in elderly patients and can potentially interact with the MF phenotype to increase long-term mortality [31–34].

Although few studies have assessed the impact of comorbidities on the outcome of patients with MF, retrospective studies have shown that a high burden of comorbidities as defined by the Adult Comorbidity Evaluation-27 (ACE-27) [31] is significantly associated with an unfavorable impact on survival in MF [32–34]. However, these studies considered the cumulative comorbidity burden rather than the impact of individual comorbidities on survival.

The main aim of the present study was to assess the influence of individual comorbidities on survival in a prospective multicenter series of MF patients.

2. **Materials and Methods**

2.1. **Study Design**

This observational prospective study was based on data obtained from a national multicenter MF register sponsored by the Spanish MPN Group (GEMFIN). The study design consisted of a baseline visit between February 2014 and October 2018 followed by...
two follow-ups at 6 and 12 months after inclusion. Each patient’s survival was monitored until death. Interviews, complementary tests and the treatment plan were carried out in line with routine clinical practice.

The inclusion criteria applied were that the patients should be aged ≥18 years, be diagnosed with MF (PMF, PPV-MF or PET-MF) as defined by the World Health Organization [1,2], and voluntarily provide signed informed consent.

The study data were recorded on a purpose-designed electronic case report form, which included standardized demographic, disease and treatment information and provided confidentiality, security and authenticity. The presence of individual comorbidities was considered at diagnosis (see the Table A1 for a list of the definitions of comorbidities). Risk stratification at diagnosis was performed using the International Prognostic Scoring System (IPSS) [15]. MF symptoms at the baseline visit were assessed by MPN-SAF [35]. This study was based on a prior review protocol (GEM-MIE-2014-01) approved by local ethics committees, and all research was performed in accordance with the provisions of the Declaration of Helsinki.

2.2. Statistical Analysis

The main endpoint was the impact of individual comorbidities on overall survival (OS) in patients with MF. Comorbidities were considered if they were present at the time of MF diagnosis. OS was calculated from the date of diagnosis until the date of death or the last follow-up (censored).

Univariate descriptive statistics of the study population were calculated. Quantitative variables were described by means of centralization and dispersion measures, and categorical data were presented as absolute numbers (N) with percentages (%).

Parametric (t-test) or non-parametric (Mann–Whitney) statistical tests were performed as appropriate to compare two independent means. According to the sample distribution, parametric (paired t-test) or non-parametric (Wilcoxon) statistical tests were performed to compare paired means. Differences in the distributions of categorical variables were evaluated by the chi-square test.

The evolution of a qualitative variable between two time points (before–after) was determined, and the associated p-value obtained, by the McNemar test.

Survival probability was determined by the Kaplan–Meier method. Multivariate adjusted hazard ratios for prognostic factors were estimated by Cox’s proportional hazard regression model. A p-value < 0.05 was deemed to indicate statistical significance. Variables with a p-value ≤ 0.05 in the univariate analysis (age (categorized into three groups defined by quartiles), hypertension, diabetes, dyslipidemia, pulmonary disease, renal dysfunction, smoking, IPSS, CALR gene mutation status and ruxolitinib treatment) were included in the multivariable analysis. Since ruxolitinib is not an intrinsic characteristic of the patient, but an external treatment that the patient may receive according to medical criteria, the model was applied with and without ruxolitinib.

All statistical analyses were performed using SPSS v24.0 (Dynamic, Madrid, Spain).

3. Results

3.1. Patient Characteristics

The study population was composed of 668 patients. Table 1 shows their demographic characteristics, disease history and comorbidities at diagnosis of MF. The median age of the patients at diagnosis was 68 years (range 25–89), 244 (36.5%) were female, and 61% had PMF. JAK2 and CALR mutations were present in 56.1% and 9.1% of these patients, respectively. On the IPSS 15, 64% (n = 431) were classed as intermediate-2 or high risk.

The most common comorbidities were hypertension (n = 282, 42.2%), diabetes (n = 124, 18.6%), dyslipidemia (n = 117, 16%) and cardiovascular disease (n = 105, 15.7%). Other comorbidities, which were less frequent but were observed in ≥5% of the patients, were pulmonary disease (n = 55, 8.2%), renal dysfunction (n = 58, 8.7%) and other neoplasms (n = 55, 8.2%). The prevalence of each comorbidity is detailed in Table 1.
Table 1. Demographic data, patient comorbidities and disease characteristics at diagnosis of MF.

| Variable                        | All Patients n = 668 |
|---------------------------------|----------------------|
| Median age, years               | 68 (25–89)           |
| Female, n (%)                   | 244 (36.5)           |
| Hypertension, n (%)             | 282 (42.2)           |
| Diabetes, n (%)                 | 124 (18.6)           |
| Dyslipidemia, n (%)             | 117 (16.0)           |
| Cardiovascular disease, n (%)   | 105 (15.7)           |
| Pulmonary disease, n (%)        | 55 (8.2)             |
| Renal dysfunction, n (%)        | 58 (8.7)             |
| Hepatic disease, n (%)          | 41 (6.2)             |
| HIV, n (%)                      | 0 (0)                |
| HBV, n (%)                      | 20 (3)               |
| HCV, n (%)                      | 10 (1.5)             |
| Other neoplasm, n (%)           | 55 (8.2)             |
| Smoking, n (%)                  | 161 (24.1)           |
| PMF, n (%)                      | 411 (61.5)           |
| JAK2, n (%)                     | 375 (56.1)           |
| CALR, n (%)                     | 61 (9.1)             |
| Splenomegaly +, n (%)           | 371 (55.5)           |
| IPSS                             |                      |
| Low risk, n (%)                 | 71 (10.6)            |
| Intermediate 1, n (%)           | 166 (24.9)           |
| Intermediate 2, n (%)           | 243 (36.4)           |
| High risk, n (%)                | 188 (28.1)           |

HIV, human immunodeficiency virus. HBV, hepatitis B virus. HCV, hepatitis C virus. IPSS, International Prognostic Scoring System. PMF, primary myelofibrosis. * Splenomegaly determined by imaging methods such as ultrasonography or computed tomography.

The median time elapsed from MF diagnosis to inclusion in the study was 5.9 years. At the time of inclusion, 15.7% (n = 105) of the patients were receiving ruxolitinib treatment, and the median MPN-SAF37 score was 19 (range 0–62).

3.2. Impact of Risk Factors on Survival

After a median follow-up of 2.49 years, 380 (56.8%) of the patients had died. The median survival was 4.01 years (3.45–4.57). Table 2 shows the results of the univariate analysis performed. Survival was greater among patients younger than 61.4 years than among those aged 61.4–76.3 years (HR = 2.04; 95% confidence interval (CI), 1.54–2.69, p < 0.001) and 76.3 years (HR = 4.43; 95% CI, 3.23–6.06, p < 0.001). In our cohort, the female patients survived longer than males (HR = 0.76; 95% CI, 0.61–0.95, p = 0.017). Intermediate-2 (HR = 3.65; 95% CI, 2.31–5.77, p < 0.001) and high-risk IPSS categories were associated with shorter survival (HR = 5.10; 95% CI, 3.20–8.11, p < 0.001). By contrast, patients presenting a CALR mutation (HR = 0.49; 95% CI, 0.29–0.83, p = 0.009) or treated with ruxolitinib (HR = 0.04; 95% CI, 0.01–0.12, p < 0.001) had better odds of survival.

Hypertension (HR = 4.96; 95% CI, 3.26–7.55, p < 0.001), smoking (HR 5.08; 95% CI, 3.35–7.71, p < 0.001), dyslipidemia (HR 4.65; 95% CI, 3.11–6.95, p < 0.001) and HCV (HR 4.26; 95% CI, 1.32–13.75, p = 0.015) were all strongly associated with worse survival (HR > 4 for each factor). Diabetes (HR 3.01; 95% CI, 2.07–4.36, p < 0.001), pulmonary disease (HR 3.13; 95% CI, 1.86–5.26, p < 0.001) and renal dysfunction (HR 1.82; 95% CI, 1.04–3.19, p = 0.037) were significantly associated with an increased risk of death (Figure 1). Cardiovascular comorbidity and other neoplasms showed a trend toward worse survival, but the difference was not statistically significant (p = 0.186 and p = 0.052, respectively).
### Table 2. Univariate survival model.

| Variable                        | HR    | 95% CI          | p-Value |
|---------------------------------|-------|-----------------|---------|
| Age (61.4–76.3 y) *             | 2.04  | (1.54–2.69)     | <0.001  |
| Age (≥76.3 y) *                 | 4.43  | (3.23–6.06)     | <0.001  |
| Female                          | 0.76  | (0.61–0.95)     | 0.017   |
| Hypertension                    | 4.96  | (3.26–7.55)     | <0.001  |
| Diabetes                        | 3.01  | (2.07–4.36)     | <0.001  |
| Dyslipidemia                    | 4.65  | (3.11–6.95)     | <0.001  |
| Cardiovascular disease          | 1.41  | (0.85–2.35)     | 0.186   |
| Pulmonary disease               | 3.13  | (1.86–5.26)     | <0.001  |
| Renal dysfunction               | 1.82  | (1.04–3.19)     | 0.037   |
| HCV                             | 4.26  | (1.32–13.75)    | 0.015   |
| Other neoplasm                  | 1.76  | (0.99–3.12)     | 0.052   |
| Smoking                         | 5.08  | (3.35–7.71)     | <0.001  |
| PMF vs. PPV-MF or ET-MF         | 1.07  | (0.86–1.33)     | 0.534   |
| IPSS Intermediate-1 **          | 1.56  | (0.95–2.55)     | 0.740   |
| IPSS Intermediate-2 **          | 3.65  | (2.31–5.77)     | <0.001  |
| IPSS High Risk **               | 5.10  | (3.20–8.11)     | <0.001  |
| JAK2                            | 0.83  | (0.65–1.06)     | 0.140   |
| CALR                            | 0.49  | (0.29–0.83)     | 0.009   |
| Splenomegaly +                  | 1.12  | (0.80–1.58)     | 0.509   |
| MPN-SAF (9–31.5) ***            | 0.98  | (0.19–5.28)     | 0.985   |
| MPN-SAF (≥31.5) ***             | 0.60  | (0.05–5.82)     | 0.600   |
| Ruxolitinib                     | 0.04  | (0.01–0.12)     | <0.001  |

PPV-MF, myelofibrosis secondary to polycythemia vera. PET-MF, myelofibrosis essential thrombocythemia. Other abbreviations are explained in Table 1. * Reference category: <61.4 years. ** Reference category: low risk. *** Reference category: <9. + Splenomegaly determined by imaging methods such as ultrasonography or computed tomography.

#### 3.3. Multivariate Analysis

Table 3 shows the multivariate Cox’s proportional hazards models obtained. When ruxolitinib was excluded from the regression analysis, the individual comorbidities significantly associated with survival were pulmonary disease (HR = 2.69; 95% CI, 1.47–4.91, p = 0.001), smoking (HR = 3.34; 95% CI, 1.85–6.04, p < 0.001), renal dysfunction (HR = 2.08; 95% CI, 1.02–4.21, p = 0.043) and HCV (HR = 11.49; 95% CI, 2.74–48.25, p = 0.001).

Additional independent risk factors for survival were age and IPSS. Thus, shorter survival was associated with increasing age (age range 61.4–76.3 (HR = 2.51; 95% CI, 1.12–5.61, p = 0.026), age ≥76.3 (HR = 4.85; 95% CI, 1.75–13.41, p = 0.002)) and higher-risk IPSS categories (IPSS Intermediate-2 vs. low risk (HR = 4.76; 95% CI, 1.39–16.22, p = 0.013), IPSS high risk vs. low risk (HR = 11.34; 95% CI, 3.24–39.70, p < 0.001)).

When ruxolitinib was included in the regression analysis, the individual comorbidities significantly associated with survival were pulmonary disease (HR = 2.40; 95% CI, 1.29–4.47, p = 0.006), smoking (HR = 3.82; 95% CI, 2.02–7.24, p < 0.001) and HCV (HR = 9.86; 95% CI, 2.34–41.64, p = 0.002). Age 61.4 years and IPSS Intermediate-2 continued to have a negative impact on survival: age 61.4–76.3 (HR = 2.91; 95% CI, 1.24–6.85, p = 0.014), age ≥76.3 (HR = 4.23; 95% CI, 1.53–11.73, p = 0.006), IPSS Intermediate-2 vs. low risk (HR = 6.90; 95% CI, 1.95–24.37, p = 0.003) and IPSS high risk vs. low risk (HR = 15.97; 95% CI, 4.22–60.42, p < 0.001). Ruxolitinib treatment was significantly associated with better survival (HR = 0.13; 95% CI, 0.38–0.42, p = 0.001).
Figure 1. Kaplan-Meier analyses showing overall survival in patients with MF. (A) Univariate analysis for overall survival by hypertension. (B) Univariate analysis for overall survival by diabetes. (C) Univariate analysis for overall survival by dyslipidemia. (D) Univariate analysis for overall survival by pulmonary disease. (E) Univariate analysis for overall survival by smoking. (F) Univariate analysis for overall survival by renal dysfunction.
Table 3. Multivariate Cox’s proportional hazards models.

| Variable                     | Without Ruxolitinib | With Ruxolitinib |
|------------------------------|----------------------|------------------|
|                              | HR 95% CI p-Value    | HR 95% CI p-Value|
| Age (61.4–76.3 y) *          | 2.57 (1.12–5.61) 0.026 | 2.91 (1.24–6.85) 0.014 |
| Age (≥76.3 y) *              | 4.85 (1.75–13.41) 0.002 | 4.23 (1.53–11.73) 0.006 |
| Hypertension                 | 1.09 (0.52–2.24) 0.842 | 1.09 (0.52–2.24) 0.842 |
| Diabetes                     | 2.20 (0.98–4.94) 0.057 | 1.46 (0.61–3.49) 0.394 |
| Dyslipidemia                 | 1.40 (0.68–2.87) 0.365 | 1.07 (0.48–2.36) 0.872 |
| Renal dysfunction            | 2.08 (1.023–4.21) 0.043 | 1.69 (0.81–3.50) 0.159 |
| Pulmonary disease            | 2.69 (1.47–4.91) 0.001 | 2.40 (1.29–4.47) 0.006 |
| Smoking                      | 3.34 (1.85–6.04) <0.001 | 3.82 (2.02–7.24) <0.001 |
| HCV                          | 11.49 (2.74–48.25) 0.007 | 9.86 (2.34–41.64) 0.002 |
| IPSS Intermediate-1 **       | 2.53 (0.73–8.91) 0.142 | 2.94 (0.82–10.52) 0.096 |
| IPSS Intermediate-2 **       | 4.76 (1.40–16.22) 0.013 | 6.90 (1.95–24.37) 0.003 |
| IPSS High Risk **            | 11.34 (3.24–39.7) <0.001 | 15.97 (4.22–60.42) <0.001 |
| CALR                         | 0.97 (0.28–3.42) 0.966 | 0.82 (0.23–2.89) 0.763 |
| Ruxolitinib                  | -                    | 0.12 (0.04–0.43) 0.001 |

HR: hazard ratio. Other abbreviations are explained in Table 1. * Reference category: <61.4 years. ** Reference category: low risk.

4. Discussion

As expected, the majority of patients in our cohort had comorbidities that ultimately affected their fitness and ability to undergo MF therapy. Current methods of risk-stratifying patients with MF, including widely used prognostic models such as IPSS [15], DIPSS [16] and DIPSS Plus [17], do not take into account patient comorbidities, although they are known to impact survival in numerous malignancies. Until recently, clinical trials excluded patients with significant organ dysfunction and thus provided limited information on how such patients should be managed. The aim of our prospective study was to investigate the influence of individual comorbidities at diagnosis on the outcomes of MF patients.

Comorbidities are known to be associated with inferior survival among patients with MF [32–34]. The ACE-27 [31], which is specifically designed for patients with cancer, is the instrument most commonly used in previous studies of MF patients to measure the severity of their comorbidities [32–34]. In 2014, a retrospective study of 131 MF patients by Lekovic et al. [34] suggested ACE-27 could help predict patient survival. On the other hand, their multivariate model did not find age to be an important prognostic factor for survival. In the same year, Newberry et al. [33] reported that comorbidities had a significant negative impact on OS in PMF patients aged <65 years, but not in older patients. More recently, Bartozsko et al. [32] evaluated two comorbidity scales in a cohort of 309 patients with PPV-MF or PET-MF and PMF, and concluded that severe comorbidities according to ACE-27 (score ≥3) were associated with a reduced OS. By contrast, a high score on the hematopoietic cell transplantation comorbidity index (HCT-CI) [36] did not reach statistical significance. No differential effect of severe comorbidities on survival based on age was detected. Finally, Breccia et al. [37] showed that baseline comorbidities did not influence the probability of achieving spleen/symptom responses for MF patients receiving ruxolitinib treatment. It should be noted that the assessment of comorbidities differed among these studies, which precluded our reaching firm conclusions on the impact of this factor on survival in MF.

In our study, smoking, pulmonary disease, HCV and renal comorbidities were independent predictors of OS, whereas diabetes was only marginally associated with worse survival in MF patients. By contrast, hypertension and dyslipidemia did not have a significant impact on outcomes. As expected, older age and advanced IPSS were associated with poor survival. Of note, ruxolitinib treatment was associated with improved survival and with the impact of comorbidities on OS. Thus, when ruxolitinib exposure was included in the regression model, renal dysfunction was no longer associated with survival, while other comorbidities such as HCV and pulmonary conditions had a weaker influence on OS. These findings might be due to the well-known anti-inflammatory effect of ruxolitinib [38],
counteracting the inflammation process mediated by these comorbidities, as has been suggested by Hasselbalch [39,40].

Neither hypertension nor dyslipidemia significantly influenced outcomes, perhaps due to the effective management of these comorbidities (assuming the levels recorded were within the target range). Several recent studies on the impact of cardiovascular risk factors (CVRFs) such as diabetes, hypertension or dyslipidemia on cardiovascular complications have demonstrated significant improvement in outcomes achieved by early, effective control [41–43].

The risk of cardiovascular events is increased in MF patients [44,45], but it is unclear to what extent this is a direct complication of MF and what role is played by other CVRFs. The physiopathology of cardiovascular events is known to be associated with inflammatory disorders [46], but further research is needed to measure the impact of correct management of CVRFs and the effect of ruxolitinib on these events. Both CVRF control and ruxolitinib are believed to reduce levels of vascular inflammation [39,40] and could play a crucial role in alleviating or preventing inflammation-mediated complications. Therefore, close collaboration between the hematologist and other medical specialists in managing and controlling comorbid conditions is an important aspect of achieving optimal outcomes.

It has long been understood that comorbidities are a significant element in the evolution of cancer patients and that any evaluation should take this circumstance into account [30]. The cumulative burden of morbidity, rather than individual, disease-specific effects, is normally considered a risk marker for mortality [26–34]. To our knowledge, the present study is the first to consider the impact of individual comorbidities on patients with MF.

Our analysis has several limitations that should be acknowledged. First, the restricted sample size and event number per comorbidity limited the power to detect comorbidity effects. However, the major comorbidities were well-represented in the cohort. Second, our study considered all-cause and non-MF-related mortality versus other specific disease mortality. This approach was taken because it is difficult to ascertain disease-related deaths in an elderly population such as that of MF patients. Third, because the study was designed before the 2016 WHO classification, we did not make distinction between prefibrotic MF and overt MF [1]. Finally, the impact of comorbidities may change due to variations in the medical care provided.

5. Conclusions
In summary, our study findings suggest that individual comorbidities may significantly influence the OS of MF patients. Therefore, it is essential to take individual comorbidities into account in forecasting survival and optimizing treatment management for MF patients. Treatment with ruxolitinib seems to reduce the deleterious effect of specific comorbidities, which could partially explain its association with improved survival in MF patients. However, further studies are needed to confirm these findings.

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Appendix A

The Membership of the Group Español de Enfermedades Mieloproliferativas Filadelfia Negativas (GEMFIN): Albo C., Hospital Alvaro Cunqueiro, Vigo; Alonso José M., Complejo Asistencial de Palencia; Alonso-Dominguez J.M., Hospital Universitario Fundación Jiménez Díaz; Angona Figueras A., Senín A., Hospital del Mar, Barcelona; Boqué C. Hospital Durán i Reynalds, Hospital de Llobregat; Caballero G., Hospital Miguel Servet, Zaragoza; Cáceres A., Carrera M.D., Hospital Arnau de Vilanova, Valencia; Calvo J.M., Hospital Nuestra Señora de Sonsoles, Ávila; Cuevas B., Hospital Universitario de Burgos; Durán M.A., Hospital Son Espases, Palma de Mallorca; Ferrer-Marin F., Hospital Morales Meseguer, Murcia; Fox M.L., Hospital Universitario Vall d’Hebron, Barcelona; García-Delgado R., García-Fortes M., Hospital Virgen de la Victoria, Málaga; García-Gutiérrez V., Hospital Ramón y Cajal, Madrid; García-Hernández C., Hospital General de Alicante; Gómez-Casares M.T., Stuckey Ruth, Hospital Dr Negrín, Las Palmas de Gran Canarias; Guerra J.M., Hospital de Son Llatzer, Palma de Mallorca; Hernández-Rivas J.A., Hospital Universitario Infanta Leonor, Madrid; López-Abadía E., Conesa V., Hospital General Universitario de Elche; Magro E., Hospital Príncipe de Asturias, Alcalá de Henares; Mata M.I., Hospital de la Costa del Sol, Marbella; Mora E., Hospital Universitari i Politècnic, La Fe, Valencia; Moretó A., Del Orbe R, Hospital de Cruces, Barakalde; Martínez C., Hospital de Sant Pau, Barcelona; Martínez-López J., Hospital 12 de Octubre, Madrid; Murillo I., Hospital General de San Jorge, Huesca; Noya-Pereira M.S., Hospital Universitario de A Coruña; Palomino A., Hospital Clínic, Barcelona; Pastor-Galán I., Hospital Clínico, Valencia; Pérez-Encinas M., Hospital Clínico Universitario, Santiago de Compostela; Pérez-López R., Hospital Virgen de la Arrixaca, Murcia; Ramos de León Y., Gordillo M., Hospital Doctor Molina Orosa, Arrecife; Raya J.M., Hospital Universitario de Canarias, Tenerife; Sagüés M., Hospital Josep Trueta-ICO, Girona; Xicoy B., Hospital Germans Trias i Pujol, Badalona.
### Appendix B

#### Table A1. Definitions of comorbidities.

| Comorbidity     | Definition                                                                 |
|-----------------|---------------------------------------------------------------------------|
| Hypertension    | Prior medical diagnosis of hypertension.                                  |
| Diabetes        | Diabetes Mellitus requiring treatment with insulin or hypoglycemic agents at the time of initiation of conditioning or HbA1c > 7%. |
| Dyslipidemia    | Prior medical diagnosis of dyslipidemia.                                  |
| Cardiovascular  | Coronary artery disease, congestive heart failure, history of myocardial infarction, or left ventricular ejection fraction ≤ 50%, or prior diagnosis of cerebrovascular disease (diagnostic imaging tests is required). |
| Pulmonary       | DLCo1 or FEV1 < 80%. Restrictive Lung Disease or Chronic obstructive pulmonary disease with dyspnea. |
| Renal           | eGFRc < 60 mL/min.                                                         |
| Hepatic         | Serum bilirubin ≥ 1.5 times ULN; ALT or AST ≥ 2.5 times ULN, or chronic hepatitis. |
| HIV             | Prior medical diagnosis of HIV by using serological and molecular test.    |
| HBV             | Prior medical diagnosis of active HBV infection by using serological and molecular test. |
| HCV             | Prior medical diagnosis of active HCV infection by using serological and molecular test. |
| Other neoplasm  | Malignancy unrelated to the Myelofibrosis which has been treated at any point in the patient’s history, excluding non-melanoma skin cancer Someone who has smoked greater than 100 cigarettes (including hand rolled cigarettes, cigars, cigarillos etc) in their lifetime and has smoked in the last 180 days. |
| Smoking         | Restricted Lung Disease or Chronic obstructive pulmonary disease with dyspnea. |

ALT, alanine transaminase; AST, aspartate transaminase; DLCo, diffusion capacity of the lungs for carbon monoxide; FEV1, forced expiratory volume in the first second; eGFR, estimated glomerular filtration rate; HIV, human immunodeficiency virus; HBV, Hepatitis B virus; HCV, Hepatitis C virus; ULN, upper limit of normal.

#### References

1. Arber, D.A.; Orazi, A.; Hasserjian, R.; Thiele, J.; Borowitz, M.J.; Le Beau, M.M.; Bloomfield, C.D.; Cazzola, M.; Vardiman, J.W. The 2016 Revision to the World Health Organization Classification of Myeloid Neoplasms and Acute Leukemia. *Blood* 2016, 127, 2391–2405. [CrossRef] [PubMed]

2. Barosi, G.; Mesa, R.A.; Thiele, J.; Cervantes, F.; Campbell, P.J.; Verstovsek, S.; Dupriez, B.; Levine, R.L.; Passamonti, F.; Gotlib, J.; et al. Proposed Criteria for the Diagnosis of Post-Polycythemia Vera and Post-Essential Thrombocythemia Myelofibrosis: A Consensus Statement from the International Working Group for Myelofibrosis Research and Treatment. *Leukemia* 2008, 22, 437–438. [CrossRef] [PubMed]

3. Cerquozzi, S.; Tefferi, A. Blast Transformation and Fibrotic Progression in Polycythemia Vera and Essential Thrombocythemia: A Literature Review of Incidence and Risk Factors. *Blood Cancer J.* 2015, 5, e366. [CrossRef] [PubMed]

4. Swerdlow, S.H.; Campo, E.; Harris, N.L.; Jaffe, E.S.; Pileri, S.A.; Stein, H.; Thiele, J. (Eds.) *WHO Classification of Tumours of the Haematopoietic and Lymphoid Tissues*, 4th ed.; IARC: Lyon, France, 2016; ISBN -13 9789283244943.

5. Moulard, O.; Mehta, J.; Fryzek, J.; Oliwares, R.; Iqbal, U.; Mesa, R.A. Epidemiology of Myelofibrosis, Essential Thrombocythemia, and Polycythemia Vera in the European Union. *Eur. J. Haematol.* 2014, 92, 289–297. [CrossRef] [PubMed]

6. Srour, S.A.; Devesa, S.S.; Morton, L.M.; Check, D.P.; Curtis, R.E.; Linet, M.S.; Dores, G.M. Incidence and Patient Survival of Myeloproliferative Neoplasm and Myelodysplastic/Myeloproliferative Neoplasms in the United States, 2001–2012. *Br. J. Haematol.* 2016, 174, 382–396. [CrossRef] [PubMed]

7. Tefferi, A.; Laslo, T.L.; Finke, C.M.; Knudson, R.A.; Ketterling, R.; Hanson, C.H.; Maffioli, M.; Caramazza, D.; Passamonti, F.; Paradani, A. CALR vs JAK2 vs MPL-Mutated or Triple-Negative Myelofibrosis: Clinical, Cytogenetic and Molecular Comparisons. *Leukemia* 2014, 28, 1472–1477. [CrossRef]

8. Vannucchi, A.M.; Laslo, T.L.; Guiglielmelli, P.; Biamonte, F.; Paradani, A.; Pereira, A.; Finke, C.; Score, J.; Gangat, N.; Mannarelli, C.; et al. Mutations and Prognosis in Primary Myelofibrosis. *Leukemia* 2013, 27, 1861–1869. [CrossRef]

9. Klampfl, T.; Gisslinger, H.; Harutyunyan, A.S.; Nivarthi, H.; Rumi, E.; Milosevic, J.D.; Them, N.C.C.; Berg, T.; Gisslinger, B.; Pietra, D.; et al. Somatic Mutations of Calreticulin in Myeloproliferative Neoplasms. *N. Engl. J. Med.* 2013, 369, 2379–2390. [CrossRef]

10. Tefferi, A. Pathogenesis of Myelofibrosis with Myeloid Metaplasia. *J. Clin. Oncol.* 2005, 23, 8520–8530. [CrossRef]

11. Tefferi, A. Primary Myelofibrosis: 2017 Update on Diagnosis, Risk-Stratification, and Management. *Am. J. Hematol.* 2016, 91, 1262–1271. [CrossRef]
33. Newberry, K.J.; Naqvi, K.; Nguyen, K.T.; Cardenas-Turanzas, M.; Florencia Tanaka, M.; Pierce, S.; Verstovsek, S. Comorbidities Predict Worse Prognosis in Patients with Primary Myelofibrosis. Cancer 2014, 120, 2996–3002. [CrossRef] [PubMed]

34. Lekovic, D.; Gotic, M.; Perunicic-Jovanovic, M.; Vidovic, A.; Bogdanovic, A.; Jankovic, G.; Cokic, V.; Milic, N. Contribution of Comorbidities and Grade of Bone Marrow Fibrosis to the Prognosis of Survival in Patients with Primary Myelofibrosis. Med. Oncol. 2014, 31, 1–6. [CrossRef] [PubMed]

35. Emanuel, R.M.; Dueck, A.C.; Geyer, H.L.; Kiladjian, J.J.; Slot, S.; Zweegman, S.; te Boekhorst, P.A.W.; Commandeur, S.; Schouten, H.C.; Sackmann, F.; et al. Myeloproliferative Neoplasm (MPN) Symptom Assessment Form Total Symptom Score: Prospective International Assessment of an Abbreviated Symptom Burden Scoring System among Patients with MPNs. J. Clin. Oncol. 2012, 30, 4098–4103. [CrossRef] [PubMed]

36. Sorror, M.L.; Storer, B.; Storb, R.F. Validation of the Hematopoietic Cell Transplantation-Specific Comorbidity Index (HCT-CI) in Single and Multiple Institutions: Limitations and Inferences. Biol. Blood Marrow Transplant. 2009, 15, 757–758. [CrossRef]

37. Breccia, M.; Bartoletti, D.; Bonifacio, M.; Palumbo, G.A.; Polverelli, N.; Abruzzese, E.; Bergamaschi, M.; Tieghi, A.; Tirielli, M.; Iurlo, A.; et al. Impact of Comorbidities and Body Mass Index in Patients with Myelofibrosis Treated with Ruxolitinib. Ann. Hematol. 2019, 98, 889–896. [CrossRef]

38. Song, H.T.; Cui, Y.; Zhang, L.L.; Cao, G.; Li, L.; Li, G.; Jia, X.J. Ruxolitinib Attenuates Intimal Hyperplasia via Inhibiting JAK2/STAT3 Signaling Pathway Activation Induced by PDGF-BB in Vascular Smooth Muscle Cells. Microvasc. Res. 2020, 132, 104060. [CrossRef]

39. Hasselbalch, H.C. Perspectives on the Impact of JAK-Inhibitor Therapy upon Inflammation-Mediated Comorbidities in Myelofibrosis and Related Neoplasms. Expert Rev. Hematol. 2014, 7, 203–216. [CrossRef]

40. Bjørn, M.E.; Hasselbalch, H.C. The Impact of Ruxolitinib Treatment on Inflammation-Mediated Comorbidities in Myelofibrosis and Related Neoplasms. Clin. Case Rep. 2015, 3, 499–503. [CrossRef]

41. Lee, J.H.; Kim, S.H.; Kang, S.H.; Cho, J.H.; Cho, Y.; Oh, I.Y.; Yoon, C.H.; Lee, H.Y.; Youn, T.J.; Chae, I.H.; et al. Blood Pressure Control and Cardiovascular Outcomes: Real-World Implications of the 2017 ACC/AHA Hypertension Guideline. Sci. Rep. 2018, 8. [CrossRef]

42. Laiteerapong, N.; Ham, S.A.; Gao, Y.; Moffet, H.H.; Liu, J.Y.; Huang, E.S.; Karter, A.J. The Legacy Effect in Type 2 Diabetes: Impact of Early Glycemic Control on Future Complications (the Diabetes & Aging Study). Diabetes Care 2019, 42, 416–426. [CrossRef] [PubMed]

43. Storey, B.C.; Staplin, N.; Haynes, R.; Reith, C.; Emberson, J.; Herrington, W.G.; Wheeler, D.C.; Walker, R.; Fellström, B.; Wanner, C.; et al. Lowering LDL Cholesterol Reduces Cardiovascular Risk Independently of Presence of Inflammation. Kidney Int. 2018, 93, 1000–1007. [CrossRef] [PubMed]

44. Haybar, H.; Khodadi, E.; Shahjahani, M.; Saki, N. Cardiovascular Events: A Challenge in JAK2-Positive Myeloproliferative Neoplasms. Cardiovasc. Hematol. Disord. Targets 2018, 17, 161–166. [CrossRef] [PubMed]

45. Frederiksen, H.; Szepligeti, S.; Bak, M.; Ghanima, W.; Hasselbalch, H.C.; Christiansen, C.F. Vascular Diseases in Patients with Chronic Myeloproliferative Neoplasms—Impact of Comorbidity. Clin. Epidemiol. 2019, 11, 955–967. [CrossRef] [PubMed]

46. Golia, E.; Limongelli, G.; Natale, F.; Fimiani, F.; Maddaloni, V.; Pariggiano, I.; Bianchi, R.; Crisci, M.; D’Acierno, L.; Giordano, R.; et al. Inflammation and Cardiovascular Disease: From Pathogenesis to Therapeutic Target. Curr. Atheroscler. Rep. 2014, 16. [CrossRef] [PubMed]