**ORIGINAL ARTICLE**

**Involvement of MAF/SPP1 axis in the development of bone marrow fibrosis in PMF patients**

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Primary myelofibrosis (PMF) is a myeloproliferative neoplasm characterized by hyperplastic megakaryopoiesis and myelofibrosis. We recently described the upregulation of MAF (v-maf avian musculoaponeurotic fibrosarcoma oncogene homolog) in PMF CD34+ hematopoietic progenitor cells (HPCs) compared to healthy donor. Here we demonstrated that MAF is also upregulated in PMF compared with the essential thrombocytemia (ET) and polycytemia vera (PV) HPCs. MAF overexpression and knockdown experiments shed some light into the role of MAF in PMF pathogenesis, by demonstrating that MAF favors the megakaryocyte and monocyte/macrophage commitment of HPCs and leads to the increased expression of proinflammatory and profibrotic mediators. Among them, we focused our further studies on SPP1 and LGALS3. We assessed SPP1 and LGALS3 protein levels in 115 PMF, 47 ET and 24 PV patients plasma samples and we found that SPP1 plasma levels are significantly higher in PMF compared with ET and PV patients. Furthermore, in vitro assays demonstrated that SPP1 promotes fibroblasts and mesenchymal stromal cells proliferation and collagen production. Strikingly, clinical correlation analyses uncovered that higher SPP1 plasma levels in PMF patients correlate with a more severe fibrosis degree and a shorter overall survival. Collectively our data unveil that MAF overexpression contributes to PMF pathogenesis by driving the deranged production of the profibrotic mediator SPP1.

**INTRODUCTION**

Primary myelofibrosis (PMF) is a Philadelphia-negative (Ph−) myeloproliferative neoplasm (MPN). Ph− MPNs include polycythemia vera (PV) and essential thrombocytemia (ET) besides PMF and share a myeloproliferative phenotype with clonal hematopoietic stem/progenitor cells expansion. On the basis of the WHO classification of Ph− MPNs, the primary criteria for differential diagnosis of PMF from ET and PV are the presence of hyperplastic and dysplastic megakaryocytes with maturation defects and the bone marrow (BM) fibrosis, that results from reticulin and/or collagen fibers deposition.1 The malignant hematopoietic cells, mainly megakaryocytes and monocytes, are endowed of a proinflammatory and profibrotic phenotype in PMF patients.2 The altered interplay between these cells and the BM stromal cells in turn drives the development of myelofibrosis, osteosclerosis and extramedullary hematopoiesis, that are all features of PMF pathogenesis.3

Great efforts to unravel the genomic landscape of MPNs have provided an increasing knowledge of the genomic lesions responsible for the uncontrolled clonal proliferation, with the identification of ‘driver’ mutations (JAK2, MPL and CALR) that directly or indirectly lead to a deranged JAK/STAT signaling.4–7 Nevertheless, none of the genomic lesions described to date is able to completely recapitulate all the pathological features of PMF in vivo. Therefore, further efforts should be focused on elucidating the molecular mechanisms behind the skewed crosstalk between the malignant hematopoietic cells and BM microenvironment to develop novel therapeutic options to dampen the proinflammatory and profibrotic phenotype of malignant hematopoietic cells in PMF patients.2

To shed light on the molecular mechanisms underlying PMF pathogenesis, we recently compared the gene expression profiles of CD34+ hematopoietic progenitor cells (HPCs) from PMF patients and healthy donors (HDs).8 This study provided many insights into the molecular mechanisms potentially relevant for PMF pathogenesis; notably we found the strong upregulation of the transcription factor MAF (v-maf avian musculoaponeurotic fibrosarcoma oncogene homolog) in CD34+ HPCs from PMF patients.8

MAF belongs to the large-Maf family of transcription factors that act as homo- or heterodimers by binding to the Maf-responsive
elements.\textsuperscript{9,10} c-Maf gene gives rise to two splicing isoforms, MAF variant 1 (RefSeq NM_005360.3) and MAF variant 2 (RefSeq NM_001031804.1). The proteins encoded differ for their carboxy-terminal portion. Translocations involving c-Maf locus are present in 5% of multiple myeloma patients, whereas an additional 50% of cases overexpresses MAF through an unknown mechanism.\textsuperscript{11} In both these conditions, the oncogenic activity of MAF is linked to its overexpression.

As MAF is upregulated in PMF HPCs compared with both the HD-derived and the ET/PV patients-derived counterparts, here we investigated the role of MAF in PMF pathogenesis. Collectively, our data unveil that the upregulation of MAF in PMF patients drives the expression of proinflammatory and profibrotic genes in the HPCs progeny, therefore contributing to establish the deranged BM microenvironment, that is a hallmark of PMF.

**MATERIALS AND METHODS**

Ethic statements

Human CD34\(^+\) cells were purified upon donor’s informed written consent from umbilical cord blood (CB) samples, collected after normal deliveries, according to the institutional guidelines for discarded material. Monocytes of HDs and PMF patients, and PMF CD34\(^+\) cells were purified from peripheral blood (PB) samples. Plasma samples were collected from 186 patients with MPNs (115 PMF, 24 PV and 47 ET) and 60 HDs. BM mesenchymal stromal cells (MSC) were collected from 5 HDs and 5 PMF patients. All subjects provided informed written consent, and the study was performed under the AGIMM (AIRC-Gruppo Italiano Malattie Mieloproliferative) protocol (Comitato Etico Area Vasta dell’Azienda Ospedaliero-Universitaria Careggi, approval date: 22 April, 2011, approval number # 2011/0014777).

MAF expression analysis in PMF and HD CD34\(^+\) cells

The raw Affymetrix data previously published\textsuperscript{8} for CD34\(^+\) cells isolated from PMF patients (n = 38) and HDs (n = 30; n = 14 from BM and n = 16 from PB) were downloaded from the GEO repository (Series GSE85190). PMF patients were further characterized by the JAK2, MPL or CALR mutation carried and the BM fibrosis degree (Supplementary Table S1). The raw Affymetrix data for CD34\(^+\) cells from PV (n = 24) and ET patients (n = 27; R2 and RM, manuscript submitted) were also included for MAF expression analysis.

The probe level data were normalized and converted into expression values by using the robust multiarray average procedure in Partek GS, 6.6 Software Package (http://www.partek.com).

The comparative analyses of MAF expression in CD34\(^+\) cells from MPN patients and HDs were performed by using the Kruskal–Wallis H-test when comparing more than two independent groups. Next, the differences between sample pairs were assessed by the Mann–Whitney U test.

Statistical analysis

The statistics used for in vitro experiments data analysis were based on two-tailed Student’s t-tests for averages comparison in paired samples. Data were analyzed by using Microsoft Excel (Microsoft Office, 2008 release) and are reported as mean ± standard error of the mean (s.e.m.). \(P < 0.05\) was considered significant. Statistical analyses for clinical correlation were performed using the IBM Statistical Package for Social Sciences (SPSS) statistics v23. Comparisons between groups were performed by using the Mann–Whitney U test, and the chosen level of significance was \(P < 0.05\) with a two-tailed test. Correlation between two variables were examined by simple regression analysis. Patients with plasma levels of SPP1 and LGALS3 greater than the median value reported in normal plasma were considered high level patients. The cumulative probability of overall survival (OS) was estimated using the Kaplan–Meier method. Differences in OS among the groups were compared by using a log-rank test in univariate analysis.

**RESULTS**

MAF expression levels in CD34\(^+\) cells from MPN patients

Microarray data for human primary CD34\(^+\) HPCs from PMF patients (n = 38 from PB; n = 20 JAK2-mutated, n = 3 MPL-mutated, n = 9 CALR-mutated and n = 6 triple-negative) and HDs (n = 14 from BM and n = 16 from PB) were obtained from a data set that we previously published.\textsuperscript{5} The MAF expression data in BM-derived CD34\(^+\) cells from ET (n = 27) and PV (n = 24) patients were collected from an unpublished data set (RZ and RM, manuscript submitted).

As we previously reported,\textsuperscript{8} MAF was strongly upregulated in PMF patients compared with HDs (fold change (FC) = 8.69, \(P = 4.25 \times 10^{-12}\); Figure 1a). Both the MAF splicing variants, namely MAF variant 1 (hereafter reported as MAFVAR1, RefSeq NM_005360.3) and MAF variant 2 (MAFVAR2, RefSeq NM_001031804.1), were upregulated in PMF versus HD CD34\(^+\) cells (Supplementary Results and Supplementary Figure S1). In addition, MAF was remarkably upregulated in PMF CD34\(^+\) cells compared with ET (FC = 2.61, \(P = 1.62 \times 10^{-5}\)) and PV counterparts (FC = 2.99, \(P = 1.07 \times 10^{-5}\); Figure 1a). On the contrary, no significant correlation between the mutation carried (JAK2, MPL, CALR) and the expression levels of MAF in PMF patients was found (Figure 1b). Although the expression of MAF appeared to be slightly lower in triple-negative compared with JAK2-, MPL- and CALR-mutated PMF patients (Figure 1b), this difference was not statistically significant (\(P = 0.093\)). Furthermore, the same analysis restricted to the JAK2V617F-positive PMF and ET patients did not highlight any correlation between the JAK2V617F allele burden and the MAF expression levels. Indeed, CD34\(^+\) cells from PMF patients with different allele burden displayed superimposable MAF expression levels (\(P > 0.05\)). No differences were detected between JAK2V617 homozygous and heterozygous PMF patients (\(P = 0.3837\)). Overlapping MAF expression levels were similarly detected in ET patients regardless of their JAK2V617F allele burden (\(P > 0.05\)) (Figure 1c).

Effects of STAT5 activation on MAF expression

We asked whether the JAK/STAT signaling could underlie the overexpression of MAF in MPNs. As detected by Western Blot (Supplementary Figure S2A), MAF protein level resulted increased in JAK2- and MPL- mutant cells (Supplementary Results and Supplementary Figure S4). In contrast, no differences were detected in ET patients regardless of their JAK2V617 homozygous and heterozygous PMF patients (\(P = 0.3837\)). Overlapping MAF expression levels were similarly detected in ET patients regardless of their JAK2V617F allele burden (\(P > 0.05\)) (Figure 1c).

Role of MAF in HPCs lineage commitment and differentiation

To understand whether MAF could be relevant for PMF pathogenesis, we studied the effects of its overexpression on the lineage commitment and differentiation of HD-derived CD34\(^+\) HPCs. Collectively, our data demonstrated that MAF overexpression enhanced the monocyte/macrophage and megakaryocyte differentiation of HD HPCs (Supplementary Results, Supplementary Table S2 and Supplementary Figure S3). In line with these observations, MAF knockdown in PMF HPCs restrained their megakaryocyte and monocyte/macrophage differentiation (Supplementary Results and Supplementary Figure S4).

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To gain insights into whether and how the upregulation of MAF detected in HPCs from PMF patients (Figure 1) could be relevant for PMF pathogenesis, in a set of three independent experiments CB-derived CD34+ HPCs transduced with the LMAFVAR1IDN, LMAFVAR2IDN and LXIDN retroviral vectors were profiled by Affymetrix U219 Array (Supplementary Results; Supplementary Figures S5 and S6A–C).

A list of 729 differentially expressed genes (DEGs) in both LMAFVAR1IDN- and LMAFVAR2IDN-transduced samples compared to the controls was identified (Supplementary Tables S3 and S4).

Microarray data unveiled that several genes known to promote fibrosis (for example, ENPP2,12 and ADAM9,13 Supplementary Figure S6A) and inflammation (for example, TNF, CCL2,14 and CCL3,15 Supplementary Figure S6B) were remarkably upregulated upon MAF overexpression. Noteworthy, many of them encoded for secreted proteins (Supplementary Figure S6C).

Owing to their putative role in inflammation and fibrosis, which are key mechanisms in PMF pathogenesis, a subset of

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Figure 1. Expression levels of MAF in CD34+ cells from MPN patients. Expression levels of MAF in CD34+ cells from MPN patients and HDs. Gene expression levels were measured by microarray analysis performed by Affymetrix platform as detailed in Norfo et al.8 (please, see the 'Materials and Methods' section). MAF expression levels are reported as Robust Multiarray Analysis (RMA)-normalized log2 signals, which were obtained by using the Partek GS software. Boxes represent the interquartile range that contains 50% of the subjects, the horizontal line in the box marks the median, and bars show the range of values. Data in a are representative of 38 PMF, 27 ET, 24 PV and 30 HD samples (14 from BM and 16 from PB). Only PMF (n = 38) and HD (n = 30) samples are shown in b, where PMF samples are classified into JAK2-mutated (n = 20), MPL-mutated (n = 3), CALR-mutated (n = 9) and triple-negative (n = 6), based on the mutational status. Only JAK2V617F-positive PMF (n = 20) and ET (n = 16) samples are shown in c, where samples are classified based on the JAK2V617F allele burden. BM, bone marrow; ET, essential thrombocythemia; MPN, myeloproliferative neoplasm; PMF, primary myelo fibrosis; PV, polycythemia vera; HD, healthy donor; PB, peripheral blood; n, number of samples.
transcripts coding for inflammation and fibrosis mediators was selected for the qRT-PCR validation (Supplementary Table S5). TaqMan qRT-PCR was performed in LMAFVAR1DN-, LMAFVAR2DN- and LXIDN-transduced cells from an independent set of five experiments and allowed us to validate the expression of 7 out of 8 genes (87.5%) (Supplementary Figures S6D and E).

Expression of IL-8, THBS1, CCL2, PLAUR, MMP9, LGALS3 and SPP1 in culture supernatants from MAF-overexpressing cells and in plasma from MPN patients

The secreted proteins coded by the 7 qRT-PCR-validated genes were therefore assessed by means of an enzyme-linked immunosorbent assay (ELISA) (a) in the culture supernatants from LMAFVAR1DN-, LMAFVAR2DN- and LXIDN-transduced cells,
collected at 3 days post-purification and (b) in plasma samples of PMF patients \((n = 30)\) and HDs \((n = 10)\) (Supplementary Table S6). As shown in Figure 2, subpanels i, all the proteins selected displayed significantly higher expression levels in both the MAF-overexpressing samples compared with the controls. In addition, IL-8, CCL2, PLAUR, MMP9, LGALS3 and SPP1 displayed significantly higher plasma levels in PMF patients than in HDs \((P < 0.05; \text{Figures 2a–f, subpanels ii}), whereas no difference was detected for THBS1 (Figure 2gii).

As the increase of SPP1 and LGALS3 in the plasma of PMF patients had never been reported before, we further analyzed SPP1 and LGALS3 plasma levels in 85 additional PMF patients and 60 HDs, together with 47 ET and 24 PV patients (Supplementary Table S6). As shown in Figure 3a, even though increased also in ET and PV patients versus HDs, SPP1 plasma levels were significantly higher in PMF versus ET \((P = 0.0139)\) and PV patients \((P = 0.0453)\). On the contrary, LGALS3 plasma levels were increased in all the MPN patients versus HDs \((P < 0.00001)\), with no remarkable differences among PMF, PV and ET (Figure 3b).

Assessment of the mechanistic relation between MAF and SPP1/ LGALS3 expression

To shed some light on the mechanistic connection between the increased MAF expression and the overproduction of SPP1 and LGALS3, we assessed by Luciferase reporter assay the ability of MAF to transactivate SPP1 and LGALS3 expression. The proximal promoter regions of SPP1 and LGALS3 genes were cloned upstream of the Firefly Luciferase gene. Luciferase assay demonstrated that MAF is able to transactivate SPP1 promoter, as increasing doses of MAF-coding vector caused a dose-dependent induction in the luciferase activity (Figure 4aii). Conversely, MAF did not significantly affect LGALS3 promoter-driven luciferase expression (Figure 4aii).

Overall, these data indicated that MAF is able to directly transactivate the expression of SPP1, but not of LGALS3.

Next, we wondered which cell types could be responsible for SPP1 and LGALS3 secretion. Our data strongly suggested that monocytes and megakaryocytes mainly account for SPP1 and LGALS3 production (Supplementary Results and Supplementary Figures S7A and B). Indeed, SPP1 and LGALS3 expression levels were downregulated upon MAF knockdown in monocytes, whereas only SPP1 was significantly downregulated in megakaryocytes (Figures 4bi and ii). Noteworthy, SPP1 expression was remarkably reduced upon MAF knockdown in PMF CD34+ cells (Figure 4biii).

Overall, these data reveal that MAF is able to transactivate SPP1 expression in monocytes and megakaryocytes. Of note, the downregulation of SPP1 was also observed in PMF-derived HPCs and monocytes after ruxolitinib treatment (Supplementary Results and Supplementary Figures S7C and D).

Effects of recombinant human SPP1 and LGALS3 on inflammation

To investigate the role of MAF-induced SPP1 and LGALS3 in inflammation, the expression of proinflammatory mediators (namely, IL-8, IL-6, IL-1β and TNF-α) in HDs-derived monocytes treated with recombinant human (rh)SPP1 or rhLGALS3 was analyzed. qRT-PCR data showed that both SPP1 and LGALS3 enhanced the expression of proinflammatory mediators in monocytes (Figure 4c).

Effects of rhSPP1 and rhLGALS3 on human fibroblasts and MSCs

Owing to the key role of BM fibrosis in PMF pathogenesis, we next moved to unravel the effects of SPP1 and LGALS3, whose increased plasma levels in PMF patients had never been reported before, on fibroblasts proliferation and collagen production. First, we treated normal human dermal fibroblasts with either rhSPP1 or rhLGALS3. A sample treated with rhTGF-β1 was included as positive control.

The treatment with rhSPP1 or rhLGALS3 in culture promoted fibroblasts growth, as demonstrated by the increased cell counts \((P < 0.05; \text{Figure 5a})\) and cell density (Supplementary Figure S8A) in rhSPP1- and rhLGALS3-treated fibroblasts at 24 and 48 h of culture. In addition, the increase in cell count detected upon treatment with rhSPP1 or rhLGALS3 was reversed by the simultaneous treatment with SPP1 (Figure 5bi) or LGALS3 (Figure 5bii) neutralizing antibody, respectively.

At last, we tried to elucidate whether rhSPP1 and rhLGALS3 could stimulate the fibroblasts to express collagen type III.

Figure 3. Detection of SPP1 and LGALS3 in the plasma of MPN patients by ELISA. Protein levels of SPP1 (a) and LGALS3 (b) were measured by ELISA in plasma samples derived from PMF \((n = 115)\), ET \((n = 47)\) and PV \((n = 24)\) patients and HDs \((n = 60)\). Protein levels are expressed as ng/mL. Boxes represent the interquartile range that contains 50% of the subjects, the horizontal line in the box marks the median, and the bars show the range of values. Comparisons between two groups were performed by using the Mann–Whitney U test, and the chosen level of significance was \(P < 0.05\) with a two-tailed test. ELISA, enzyme-linked immunosorbent assay; ET, essential thrombocythemia; HDs, healthy donors; n, number of samples; PMF, primary myelofibrosis; PV, polycythemia vera.
Figure 4. Role of MAF in the regulation of SPP1 and LGALS3 expression. (a) MAFvar1 and MAFvar2-dependent transactivation of human SPP1 promoter-driven (i) and LGALS3 promoter-driven (ii) luciferase expression. The amounts (in ng) of co-transfected plasmids are reported. Renilla-normalized Firefly luciferase levels (mean ± s.e.m.; n = 5 for SPP1 promoter and n = 5 for LGALS3 promoter, respectively) are normalized by setting as = 1 the pXP1-SPP1(−853/+22)/LXIDN empty vector-transfected and the pXP1-LGALS3(−1720/+35)/LXIDN empty vector-transfected samples, respectively. Error bars represent s.e.m. *P < 0.05 compared to pXP1-SPP1(−853/+22)/LXIDN empty vector-transfected (i) or pXP1- pXP1-LGALS3(−1720/+35)/LXIDN empty vector-transfected (ii) samples, respectively. (b) Expression levels of MAFvar1, MAFvar2, SPP1 and LGALS3 upon MAF knockdown in HD-derived (i) megakaryocytes (n = 6) and (ii) monocytes (n = 6) as well as (iii) in PMF patients-derived CD34+ cells (n = 5) were measured by qRT-PCR 24 h post-nucleofection. Expression levels are reported as mean RQ ± s.e.m. respect to the NegCTRsiRNA sample, set as calibrator. *P < 0.05; **P < 0.01; ***P < 0.001 versus NegCTRsiRNA. (c) Expression levels of proinflammatory cytokines IL-6, TNF-α and IL-1β in HDs monocytes are measured by qRT-PCR after 24 h of culture in the presence or absence of rhSPP1 or rhLGALS3 and are reported as RQ (mean ± s.e.m.; n = 6) respect to the untreated control samples, set as calibrator. *P < 0.05 versus untreated control. IL-1β, interleukin-1β; IL-6, interleukin-6; IL-8, interleukin-8; n, number of experiments; rh, recombinant human; RQ, relative quantity; s.e.m., standard error of the mean; TNF-α, tumor necrosis factor-alpha.
and collagen type I (COL1A1), the main components of reticulin and collagen fibers, respectively. qRT-PCR data showed that treatment with rhSPP1 and rhLGALS3 increased the expression of COL1A1 (Figure 5c), but not of COL3A1 (data not shown).

Given the role of MSCs in BM remodeling, we also investigated the effects of rhSPP1 and rhLGALS3 on HD (Supplementary Figures S8B-D) and PMF BM MSCs (Figure 6). The treatment with rhSPP1 and rhLGALS3 increased both HD (Supplementary Figure S8B) and PMF MSCs proliferation (Figure 6a). This effect was reverted by treatment with SPP1 (Supplementary Figure S8Ci and Figure 6bi) or LGALS3 neutralizing antibodies (Supplementary Figure S8Cii and Figure 6bii).

In addition, rhSPP1 and rhLGALS3 induced COL1A1 expression in both HDs (Supplementary Figure S8D) and PMF MSCs.
whereas no modulation of COL3A1 mRNA level was observed (data not shown).

Collectively, these results suggest that SPP1 and LGALS3 could promote fibroblast and MSC growth and collagen I production and therefore could be involved in BM fibrosis in PMF patients. Noteworthy, we also unveiled that the production of SPP1 was higher in PMF versus HD MSCs. Strikingly, the treatment with PMF monocytes-conditioned medium strongly increased the SPP1 protein release by PMF MSCs (Supplementary Results and Supplementary Figures S8E and F).

Correlations between MAF, SPP1 and LGALS3 expression and clinical features of PMF patients

As our data collectively supported the idea of a profibrotic role for MAF/SPP1 axis, we further analyzed the expression levels of MAF in CD34+ cells from PMF patients according to their BM fibrosis grade. Strikingly, we uncovered that MAF expression levels in CD34+ cells were significantly higher in overt fibrotic (MF-2/MF-3) compared with prefibrotic (MF-0/MF-1) PMF (Figure 7a).

We next unraveled the correlations between SPP1 and LGALS3 plasma levels and clinical features of PMF patients. We first compared hematological, clinical and molecular characteristics of PMF patients according to the SPP1/LGALS3 expression levels in the plasma in comparison with HDs. SPP1 and LGALS3 plasma levels were significantly greater in PMF patients than in HDs (median = 70.3 ng/ml, range = 10.3–392.4 ng/ml in PMF patients versus median = 42.2, range = 4.4–87.4 ng/ml in HDs for SPP1, P < 0.00001 and median = 8.8 ng/ml, range = 1.6–36.2 ng/ml in PMF patients versus median 5.4 ng/ml, range = 1.5–12.8 ng/ml in HDs for LGALS3, P < 0.00001).

Figure 6. Effects of SPP1 and LGALS3 on PMF mesenchymal stromal cells. PMF-derived MSCs were cultured in the absence or presence of 500 ng/ml rhSPP1, 1000 ng/ml rhLGALS3 or 5 ng/ml rhTGF-β1 (used as positive control) for 48 h. (a) Cell proliferation was monitored by Trypan Blue exclusion assay after 48 h of treatment. Results are shown as number of cells, starting from 1.6 × 10⁶ cells per sample. Values are reported as mean ± standard error of the mean (s.e.m.). The results come from three independent experiments. * P < 0.05; ** P < 0.01 versus untreated control. (b) Effect of an anti-SPP1 (i) and anti-LGALS3 (ii) neutralizing antibody on MSCs cell proliferation. PMF MSCs were treated with rhSPP1 (i) or rhLGALS3 (ii) in the absence or presence of anti-SPP1 neutralizing antibody (2.5 μg/ml, bi) or anti-LGALS3 neutralizing antibody (10 μg/ml, bii) or an isotype-matched antibody for 48 h. Cell proliferation was evaluated by Trypan Blue exclusion assay. For each sample the fold of increase in cell counts after 48 h of culture was normalized to the number of cells plated at t₀. Values are reported as mean ± s.e.m. (n = 5). * P < 0.05. (c) Expression levels of the fibrotic marker COL1A1 in PMF MSCs were measured by qRT-PCR after 48 h of culture with rhSPP1, rhLGALS3 or rhTGF-β1 and are reported as RQ (mean ± s.e.m.; n = 5) respect to the untreated control sample, set as calibrator. * P < 0.05; ** P < 0.01. COL1A1, collagen type 1 alpha 1; MSCs, mesenchymal stromal cells; rh, recombinant human; n, number of experiments.

(Figure 6c), whereas no modulation of COL3A1 mRNA level was observed (data not shown).

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On the basis of SPP1 and LGALS3 plasma levels in HDs, we defined high SPP1 ($n = 88$ patients), low SPP1 ($n = 27$), high LGALS3 ($n = 96$) and low LGALS3 ($n = 19$) levels those patients with SPP1/LGALS3 levels higher or lower, respectively, than the median value in HDs (42.2 ng/ml for SPP1 and 5.4 ng/ml for LGALS3).

Patients with higher SPP1 plasma levels suffered more frequently of constitutional symptoms (94.9%) compared with patients with normal SPP1 levels (67.1%; $P = 0.001$).

Further clinical correlations results are available in the Supplementary Results section. The levels of SPP1 were progressively higher depending on fibrosis grade, increasing from fibrosis grade 0–1 (MF-0/1) (median = 51.8 ng/ml, range = 12.1–220.5 ng/ml, $n = 54$) to grade 2 (MF-2) (median = 96.0 ng/ml, range = 10.3–392.4 ng/ml, $n = 43$) and grade 3 (MF-3) (median = 103.5 ng/ml, range = 18.6–365.9 ng/ml, $n = 18$) ($P = 0.0016$ in MF-2 versus MF-0/1 and $P = 0.0021$ in MF-3 versus MF-0/1, Figure 7b). Furthermore, higher plasma levels of SPP1 were more frequently detected in the high molecular risk (HMR) category (median = 113.8 ng/ml, range = 22.2–392.4 ng/ml) in comparison with the low molecular risk group (median = 57.2 ng/ml, range = 10.3–364.7 ng/ml) ($P = 0.014$); no difference was observed for LGALS3 levels according to HMR status. We then evaluated the impact of SSP1 and LGAL3 plasma levels on OS. In univariate analysis, patients with higher SPP1 plasma levels displayed significantly shorter OS compared to those with SSP1 level in the normal range (HR: 31, 95% CI: 1.0–183.7, $P = 0.006$; Figure 7c) together with the IPSS variables evaluated at the time of plasma collection ($P < 0.001$). In multivariable Cox proportional hazard regression analysis, higher SPP1 plasma levels remained...
significant for survival (HR: 12.7, 95% CI:1.36–119.2; P = 0.026) together with all the IPSS categories (HR: 3.7, 95% CI: 2.36–5.72; P < 0.0001), therefore representing an independent prognostically adverse variable for survival in PMF.

DISCUSSION

As we have recently unveiled the upregulation of MAF in CD34+ cells from PMF patients compared with their normal counterparts, in this study we further unraveled the role of MAF in PMF pathogenesis.

We uncovered that MAF is also remarkably upregulated in PMF CD34+ cells in comparison with ET and PV CD34+ cells. Notably, MAF expression levels in CD34+ cells from PMF patients are independent of the driver mutation they carry. We demonstrated that MAF promotes the HPCs commitment toward the monocyte/macrophage and megakaryocyte lineages, in line with previous reports. Of note, both megakaryocytes and monocytes are key players in PMF pathogenesis. Gene expression profile of MAF-overexpressing cells provided further insights into the role of MAF in PMF pathogenesis, as they unveiled the enhanced expression of genes related to inflammation and fibrosis.

PMF, as well as ET and PV, is associated with a chronic nonresolving inflammation. This long-lasting inflammatory state is fueled by the continuous release of proinflammatory and microenvironment-regulating cytokines and chemokines, which are mainly produced by megakaryocytes, monocytes and other hematopoietic cells. Notably, elevated levels of circulating IL-8, IL-2, IL-12 and IL-15 have been introduced as independent prognostic criteria for PMF risk stratification according to the Dynamic International Prognostic Scoring System (DIPSS) plus. Similarly, increased plasma levels of proinflammatory cytokines and chemokines, such as TNFα, IL-6 and CCL4, have been detected in PMF patients and their reduction has been related to the clinical benefits (that is, the resolution of splenomegaly and constitutional symptoms) of ruxolitinib treatment.

Our study unveils that the upregulation of MAF in CD34+ cells from PMF patients is at least in part responsible for the induction of a proinflammatory state and for the increased expression of proinflammatory mediators in their progeny. Some of these molecules (for example, IL-8, CCL2, PLAUR, SPP1 and LGALS3) showed higher plasma levels in PMF patients compared with HDS. Although CCL2, IL-8 and PLAUR have been already reported to be increased in PMF patients in comparison with HDS, no data were available for SPP1 and LGALS3, respectively. However, both these molecules have been associated with other inflammatory disorders. The upregulation of SPP1 is a hallmark of inflammatory neuropathologies such as multiple sclerosis, Parkinson’s and Alzheimer’s disease. Similarly, LGALS3 promotes inflammation by stimulating macrophage activation and migration and LGALS3 levels are associated with neutrophils accumulation in patients with chronic obstructive pulmonary disease.

As SPP1 and LGALS3 increased plasma levels in PMF patients had never been reported before, we further unraveled their role in PMF-related inflammation by demonstrating that SPP1 and LGALS3 enhance the expression of proinflammatory mediators (namely, IL-8, IL-6, IL-1β and TNF-α) in monocytes. No correlation between SPP1/LGALS3 plasma levels and LDH levels, neutrophil or monocyte counts was found in PMF patients, suggesting that the increased SPP1/LGALS3 plasma levels did not exclusively reflect the ongoing chronic inflammation in PMF.

Interestingly, even though several data demonstrate that MPN-associated oncogenes can induce a proinflammatory response by both malignant and non-malignant cells, emerging evidence suggests that chronic inflammation itself may foster clonal evolution in MPNs by increasing the risk of oxidative DNA damage and consequently promoting the emergence of novel genomic lesions. Therefore, the question of whether the MPN-associated genonic lesions boost inflammation or vice versa still remains unanswered. As JAK/STAT signaling is involved in both malignant hematopoiesis and inflammation, we cannot exclude that chronic inflammation in the BM rather than JAK/MPL/CALR-driven oncogenic signaling could underlie the upregulation of MAF in CD34+ cells. However, our data demonstrate that MAF contributes to enhance the proinflammatory microenvironment that is a major feature of PMF and, to a lesser extent, of the other MPNs.

Concerning the development of BM fibrosis and stromal changes characterizing PMF, the most acknowledged theory is that they result from an overproduction of growth factors by malignant hematopoietic cells, mainly monocytes and megakaryocytes. The proinflammatory/profibrotic cytokines and growth factors released (for example, TGFβ, PDGF, bFGF and VEGF) in turn act on stromal cells leading to the development of BM fibrosis, osteosclerosis and other pathological features of PMF.

Here we demonstrate that the upregulation of MAF in malignant HPCs from PMF patients contributes to enhance the expression of several profibrotic mediators, such as CCL2, IL-8, MMP9, LGALS3 and SPP1 in their progeny. Indeed, all the latter molecules show significantly increased plasma levels in PMF patients in comparison with HDS. We showed that SPP1 is a direct transcriptional target of MAF, whereas the mechanism of LGALS3 overproduction upon MAF overexpression still remains unclear. Interestingly, both LGALS3 and SPP1 have been previously linked to the development of liver fibrosis. In line with these data, we demonstrated that SPP1 promotes fibroblasts and MSCs proliferation and collagen production. In addition, in line with the increasing evidences suggesting that stromal cells may foster key pathophysiological features of PMF, we showed that the production of SPP1 is significantly higher in PMF MSCs compared with their normal counterparts and, interestingly, their crosstalk with PMF monocytes strongly enhances the SPP1 production by MSCs.

Our study showed that SPP1 plasma levels in PMF patients have a prognostic value as they were able to distinguish pre-fibrotic (MF-0/MF-1) and overt fibrotic (MF-2/MF-3) PMF patients. Indeed, higher SPP1 plasma levels correlate with a more severe symptoms) of ruxolitinib treatment. In the present study, we showed for the first time an increased expression of LGALS3 and SPP1 in MPN patients’ plasma compared to HD; however, while LGALS3 could be considered a putative serum marker of MPN, SPP1 increased expression is more remarkable in PMF patients, in which it is related to a greater degree of BM fibrosis and shorter OS.
As a whole, in the present study we demonstrated that MAF–SPP1 axis is involved in the development of BM fibrosis with potential prognostic value in PMF patients.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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AUTHOR CONTRIBUTIONS

AMV and RM designed the study; PG enrolled patients and analyzed clinical correlations; GCF, CM, TF, NB, BM, LE, SiSa, contributed patients cell samples and performed molecular characterization; VR, FP, AR enrolled patients; MA performed MSC experiments; SeRo and SS performed microarray and WB experiments; CR and ZP performed real-time PCR assays; SaRu and EB performed ELISA assays; GB and LT performed luciferase reporter assays; RZ, RN, VP and ET performed data analysis; SaRu and EB performed gene overexpression experiments; EB, SF, AMV and RM wrote the manuscript.

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Supplementary Information accompanies this paper on the Leukemia website (http://www.nature.com/leu)