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Integrated Genomic and Transcriptomic Analysis Improves Disease Classification and Risk Stratification of MDS with Ring Sideroblasts



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ABSTRACT

Purpose: Ring sideroblasts (RS) define the low-risk myelodysplastic neoplasm (MDS) subgroup with RS but may also reflect erythroid dysplasia in higher risk myeloid neoplasm. The benign behavior of MDS with RS (MDS^{RS+}) is limited to *SF3B1*-mutated cases without additional high-risk genetic events, but one third of MDS^{RS+} carry no *SF3B1* mutation, suggesting that different molecular mechanisms may underlie RS formation. We integrated genomic and transcriptomic analyses to evaluate whether transcriptome profiles may improve current risk stratification.

Experimental Design: We studied a prospective cohort of MDS^{RS+} patients irrespective of World Health Organization (WHO) class with regard to somatic mutations, copy-number alterations, and bone marrow CD34⁺ cell transcriptomes to assess

whether transcriptome profiles add to prognostication and provide input on disease classification.

Results: *SF3B1*, *SRSF2*, or *TP53* multihit mutations were found in 89% of MDS^{RS+} cases, and each mutation category was associated with distinct clinical outcome, gene expression, and alternative splicing profiles. Unsupervised clustering analysis identified three clusters with distinct hemopoietic stem and progenitor (HSPC) composition, which only partially overlapped with mutation groups. IPSS-M and the transcriptome-defined proportion of megakaryocyte/erythroid progenitors (MEP) independently predicted survival in multivariable analysis.

Conclusions: These results provide essential input on the molecular basis of *SF3B1*-unmutated MDS^{RS+} and propose HSPC quantification as a prognostic marker in myeloid neoplasms with RS.

Introduction

Ring sideroblasts (RS) are erythroblasts with iron-loaded mitochondria visualized by Prussian blue staining (Perls' reaction) as a perinuclear ring of blue granules (1) and constitute a hallmark of myelodysplastic syndrome (MDS) with RS (MDS^{RS+}). Missense somatic mutations in the *SF3B1* gene, a core component of the splicing factor machinery, are identified in approximately 80% of MDS-ring sideroblast with single/multilineage dysplasia (MDS-RS-SLD/MLD; refs. 2, 3) according to the 2016 World Health Organization (WHO) classification (4). The identification of *SF3B1* mutations in MDS^{RS+} paved the way for characterization of the pathobiology of splicing factor mutated MDS (5–8) and iron metabolism dysregulation in RS (9, 10). On the basis of this evidence, the recent International Consensus Classification (ICC; ref. 11) as well as the WHO classification (12) recognize MDS with low blasts and *SF3B1* mutation as a distinct nosological entity because of its well-defined molecular, clinical and biological characteristics (13). However, according to the 2022 WHO criteria, the detection of ≥15% RS may substitute for *SF3B1* mutation, which makes the new systems diverge on wild-type *SF3B1* MDS with low blasts and ring sideroblasts classification. This is relevant because the benign behavior of MDS-RS is limited to *SF3B1*-mutated cases (14). Indeed, RS can also be a marker of erythroid dysplasia in bone marrow (BM) smears of MDS with excess blasts, myeloproliferative neoplasms (MPN), MDS/MPN and acute myeloid leukemia (AML) as well as nonclonal and transitory conditions (e.g., exposure to isoniazid, linezolid, ethanol, or lead poisoning; ref. 15).

Although a strong association between RS and *SF3B1* mutation has been confirmed throughout the spectrum of myeloid neoplasms, a

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Translational Relevance

MDS with ring sideroblasts (RS) and low myeloblast count is considered a lower-risk entity according to World Health Organization classifications, while the International Consensus Classification demands presence of *SF3B1* mutation. Notably, RS are also found in *SF3B1* wild-type (WT) myeloid neoplasms with or without increased blast, but molecular studies on these subtypes are lacking. We carried out a comprehensive evaluation of genomic and transcriptomic profiles of a prospective cohort of 129 MDS^{RS+} irrespective of classification. We identified *SF3B1*, *SRSF2*, or *TP53*^{multi-hit} mutations as distinct disease-modifying entities in 89% of MDS^{RS+}, and showed that outcome differed significantly between categories, highlighting the importance of identifying *TP53*^{multi-hit} status also in MDS^{RS+}. Transcriptome-defined hematopoietic progenitor signatures predicted survival independently from mutational categories and IPSS-M in multivariable analysis, proposing hematopoietic progenitor quantification as a valuable prognostic marker in myeloid neoplasms.

large proportion of cases (up to 10% in MDS/MPN-RS and thrombocytosis; MDS/MPN-RS-T, 20% in MDS-RS-SLD/MLD, 90% in AML with RS) carry no *SF3B1* mutation, suggesting that different molecular mechanisms may underlie RS formation (2, 16–20). The well-documented causative relationship between *SF3B1* mutation and RS formation in MDS-RS-SLD/MLD is elicited by ABCB7 downregulation and mitochondrial iron accumulation, but little is known about the molecular pathophysiology of *SF3B1*-unmutated MDS-RS-SLD/MLD, including whether ABCB7 plays a role also in RS formation in *SF3B1*-unmutated MDS-RS-SLD/MLD (9, 10). Furthermore, a transcriptomic signature enriched for genes specific for erythroid and megakaryocytic progenitors has been to be associated with *SF3B1*-mutated MDS-RS-SLD/MLD and favorable prognosis in MDS, while no evidence is available on *SF3B1*-unmutated MDS-RS-SLD/MLD transcriptomics and molecular determinants of its widely variable clinical course (2, 7, 14).

The recently published Molecular International Prognostic Scoring System (IPSS-M) identified different risk groups based on previous revised-IPSS criteria plus comprehensive targeted DNA-sequencing data (DNA-seq; ref. 21). An important question for this study was therefore to assess whether IPSS-M compensates for the questions raised above, and whether transcriptomic profiling adds a distinct new dimension to the understanding and prognostication of MDS^{RS+}.

We analyzed a large cohort of MDS^{RS+} irrespective of WHO categorization, and integrated somatic mutations, copy-number variation (CNV) and CD34⁺ BM mononuclear cells (MNC) transcriptome profiles at the time of diagnosis, assessing whether transcriptome profiles add to prognostication and provide input to disease classification.

Materials and Methods

Patients and samples

We studied a cohort of 129 MDS^{RS+} identified within a population of 834 myeloid neoplasms (682 MDS, 51 AML with myelodysplastic-related changes, 101 MDS/MPN) evaluated at Karolinska University Hospital in Stockholm between February 2004 and August 2020 for suspected MDS. Diagnostic procedures were performed according to the recommendations of the European LeukemiaNet and WHO

organization (4, 13). All cases were reevaluated by a central pathologist (B. Sander). MNCs from BM were collected at diagnosis and cryopreserved at Karolinska Institutet MDS Biobank following standard operating procedure. RS were deemed present if percentage of RS was $\geq 5\%$ of total nucleated erythroid cells in BM smears. Ten healthy individuals were included in the study as normal BM (NBM) controls for the comparative transcriptomic analysis. Diagnostic criteria were updated according to 2016 revision of WHO classification of hematopoietic tumors. Karolinska Institutet MDS Biobank was started on 2001 and received primary human BM samples from Stockholm area investigated for MDS or MDS/MPN. Considering the evolving classification of myeloid neoplasms over the last two decades, refractory anemia with excess blast in transformation (reclassified as AML since 2008) were also referred to the Biobank and therefore included into this study. For the same reason, cases with suspected MPN were not included in this study. Central pathology review according to the last WHO classification was carried out and diagnoses were confirmed at multiprofessional conference. Extensive clinical and survival information was available for all patients and retrieved from national electronic chart system. Cases that received any malignant hematologic-related treatment except from Erythropoiesis Stimulating Agents (ESA) prior biobanking were excluded from the study. Three additional MDS cases without RS (MDS^{RS-}) were also included into the study as control in to validate results from digital sorting (Supplementary Table S2). Cryopreserved MNCs were thawed in RPMI1640 Glutamax + 20% FBS + 100 U/mL DNase I (Merck) with a median cell viability of 92% (range 80–95). This study was approved by the local ethics committee at Karolinska Institutet (Stockholm, Sweden). All the procedures followed were in accordance with the Helsinki Declaration of 1975, regularly revised up to the last 2013 version. All samples were collected after written informed consent was obtained from patients and healthy donors. *SRSF2* and *TP53*^{MH} MDS^{RS+} overall survival (OS) was compared with historical MDS controls without RS from two recent large studies evaluating clinical characteristics of *SRSF2* and *TP53*^{MH}-mutated myeloid neoplasms (22, 23), to explore the outcome of *SF3B1*-unmutated MDS^{RS+} stratified by RS amount.

Targeted capture DNA-seq

Genomic DNA was extracted from MNC following standard protocols for human tissue in all cases and controls and submitted for DNA-seq. A validated targeted sequencing panel was used to evaluate 152 genes recurrently mutated in MDS, as well as 1,118 genome-wide SNP probes for copy-number analysis, as previously reported (21, 23). Libraries were sequenced with paired-end Illumina HiSeq with a 100 or 125 bp read length, with a median coverage of $1,241 \times$ per sample (range, $338\text{--}1,956 \times$). Raw sequence data were aligned to the human genome (GRCh37).

RNA sequencing

CD34⁺ cells (median 0.4×10^6 , range $0.1\text{--}0.5 \times 10^6$) were isolated from the MNCs using AUTO-MACS with double-separation option (Miltenyi Biotec) and submitted for RNA extraction for all cases and controls. RNA was extracted with RNeasy Microkit (Qiagen) and treated with DNase, according to the manufacturer's instructions. RNA integrity number was estimated using Agilent RNA 6000 Pico (Agilent Technologies) and was greater than 6.5 for all the samples (median 8.2). The RNA-sequencing (RNA-seq) libraries were prepared from total RNA using SMARTer Stranded Total RNA-Seq Kit v2 Pico Input Mammalian with enzymatic ribosomal depletion (Takara Bio). Libraries were sequenced using the Novaseq 6,000 with paired-end 150-bp configuration. The reads were processed with nf-core/

rnaseq pipeline (ref. 24; version 1.0), using human genome GRCh37 for STAR alignment. Uniquely mapped read pairs were counted using featureCounts (ref. 25; version 1.6.3) and used for gene expression analysis. Differential expression analysis was performed with DESeq2 (26) version 1.26. Genes with raw read counts <50 in less than 10% of samples were filtered out. Raw read counts were normalized by DESeq2 by applying variance stabilizing transformation and used for the downstream unsupervised clustering analysis after logarithmic transformation. The significance of differential expression among groups was calculated using likelihood ratio test including sequencing batch in the reduced model. Genes with false discovery rate P_{adj} (FDR) < 0.001 were considered as significantly differentially expressed. ClusterProfiler (27), Reactome PA (28) version 3.14, and fgsea (29) version 3 R packages were used for Gene Ontology (GO), pathway and gene-set enrichment analyses. Differential splicing analysis was assessed using rMATS turbo (ref. 30; version 4.1) on BAM alignments, comparing disease subgroups against NBM controls. Splicing events with FDR < 0.0001 and absolute delta-PSI > 0.1 were considered statistically significant.

Flow cytometry and HSPC sorting

MNCs isolated from cryopreserved BM samples were washed twice with PBS (Thermo Fisher Scientific) with 5% FBS before staining. Cells were sorted on a FACS ARIA II Fusion (Becton Dickinson) at the WIRM FACS facility at the Karolinska Institutet, Flemingsberg (Huddinge, Sweden). All experiments included fluorescent-minus-one (FMO) and single-stained controls. A previously described strategy was used to define the HSPC populations (31).

Clonal hierarchy analysis

The bioinformatic analysis of hierarchical rank of detected mutations was carried out with both DPCLust (22, 32) and Pyclone (33). Mutations belonging to the largest clone were tagged as dominant and distinguished from other secondary mutations harbored by smaller subclones. Single-cell-derived hematopoietic stem and progenitor colonies from *SF3B1*-*SRSF2* comutated patients were obtained as previously described (22). DNA from each single-cell-derived colony underwent droplet digital PCR (ddPCR) using mutation-specific primers, according to the manufacturer's instructions, as previously described (34).

Unsupervised clustering analysis

Consensus clustering with partitioning around medoids algorithm and binary distance measures was performed using genetic features, the latter considered as binary variable by means of ConsensusClusterPlus (version 1.50; refs. 35–37). The same methods with K-means algorithm and Euclidean distance was used to carry out unsupervised clustering analysis of scaled normalized gene expression counts (7).

Digital sorting on RNA-seq CD34⁺ MNC

To further explore composition of bulk CD34⁺ MNC, CIBERSORTx algorithm was used to provide a digital estimation of hematopoietic stem and progenitor cells (HSPC; ref. 38). Publicly available single-cell RNA-seq data representative of HSPC purified from CD34⁺ BM MNCs (39, 40) were used to create a signature matrix and quantify hematopoietic precursors using the following parameters: $\kappa = 13$, q -value = 10^{-5} , number of barcode genes = 3–50, minimum expression = 1, replicates = 30, sampling = 0.5. Only genes shared between the single cell and MDS^{RS+} bulk RNA-seq data were selected. Normalized gene expression counts were therefore used to quantify hemopoietic precursors with the following settings: absolute mode, B-mode batch correction and 500 permutations. Only outputs with Monte Carlo

sampling empirical $P < 0.05$ for the deconvolution were considered for further analysis.

Statistical analysis

Categorical variables were compared using χ^2 test or Fisher exact test, in accordance with variable characteristics. Numerical variables were compared using Mann–Whitney *U*, Kruskal–Wallis one-way ANOVA or Spearman correlation test, in accordance with variable characteristics. For multiple test correction, *P* values were adjusted by the Benjamini–Hochberg FDR.

Survival was computed from study enrollment to the event of interest (e.g., death, AML progression) and accounted for right censoring at the time of disease-modifying treatments, including intensive chemotherapy and allogeneic stem cell transplantation. Survival probabilities were estimated using Kaplan–Meier methodology, and comparisons of survival across subgroups were conducted using the two-sided log-rank test. Multivariable models of OS were performed with Cox proportional hazards regression and optimized on the results of Akaike information criterion stepwise selection. HRs and 95% confidence intervals (CI) were reported for covariates, along with *P* values from the Wald test. All statistical analyses were performed with R 3.6.2 (<https://www.r-project.org>) software.

Data availability

Deidentified individual participant data are available in the Swedish National Data Service repository at <https://doi.org/10.48723/zt59-8x04> upon request.

Results

Mutational landscape of MDS^{RS+}

A total of 129 MDS^{RS+} cases were included in the study. Median age at sampling was 74 years (range 20–88), male/female ratio was 1.2, median OS was 5.5 years (interquartile range 0.9–9.5). Ninety-five patients (74%) had MDS without blast excess (EB), 23 (18%) MDS with EB, 9 (7%) MDS/MPN, and 2 AML. Detailed clinical and molecular features of MDS^{RS+} are reported in Table 1.

Overall, 392 somatic mutations were identified in the MDS^{RS+} population. *SF3B1* was found to be the most recurrent mutated gene (67%) followed by *TET2* (37%), *DNMT3A* (19%), *SRSF2* (15%), and *TP53* (15%). MDS^{RS+} were found to have ≥ 1 , ≥ 2 or ≥ 3 recurrent somatic mutations in 97.7%, 79.2%, and 54.6% of cases, respectively. A total of 138 chromosomal alterations [96 unbalanced, 15 balanced aberrations and 27 copy-neutral loss of heterozygosity (LOH)] were identified in 54 (41.5%) MDS^{RS+} cases. Two MDS/MPN-RS-T cases did not harbor any clonal marker. NBM controls were all negative for clonal hematopoiesis with a tumor cell fraction cutoff of 1%. Multihit gene alterations (defined as having ≥ 2 distinct mutations in the same gene or one mutation associated with chromosomal alterations of the gene locus) were assessed by combining CNV analysis with somatic mutation profiling. *TET2* and *TP53* were the most recurrent genes with multi-hit (MH) mutations, with a prevalence of 19 of 48 (48%) *TET2*-mutated and 13 of 19 (68%) *TP53*-mutated cases, respectively.

Unsupervised analysis based on consensus clustering of genetic alternations was used to identify subsets with homogenous genetic profiles. We identified three major subgroups within MDS^{RS+}, each driven by *SF3B1*, *SRSF2*, or *TP53*^{MH} mutation, respectively (Fig. 1A; Table 1). Patients (11%) negative for *SF3B1*, *SRSF2*, and *TP53*^{MH} mutation showed mutational patterns concordant with a recent publication and were classified as MDS^{RS+} not otherwise specified (NOS; Supplementary Fig. S1; ref. 20). *TP53*^{MH} never occurred with *SF3B1* or *SRSF2*, whereas the combination *SF3B1*-*SRSF2*

Table 1. Clinical and hematologic features of the MDS with RS (MDS^{RS+}), stratified by genetic classification.

	Total (N = 129)	<i>SF3B1</i> ^{mutated} N = 82 (64%)	<i>SRSF2</i> ^{mutated} N = 15 (11%)	<i>TP53</i> ^{MH} N = 13 (10%)	NOS N = 19 (15%)	P
Age (years)						0.426
Mean (range)	72.7 (20–88)	73.3 (44–88)	69.6 (53–83)	69 (20–86)	72.7 (20–88)	—
Gender						0.067
Female	58 (45.0%)	39 (47.6%)	2 (13.3%)	7 (53.8%)	10 (52.6%)	—
Male	71 (55.0%)	43 (52.4%)	13 (86.7%)	6 (46.2%)	9 (47.4%)	—
Diagnosis according to the WHO 2016 classification						<0.001
MDS RS MLD	62 (48.1%)	45 (54.9%)	7 (46.7%)	4 (30.8%)	6 (31.6%)	—
MDS RS SLD	23 (17.8%)	23 (28%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	—
MDS MPN RS T	7 (5.4%)	2 (2.4%)	0 (0.0%)	0 (0.0%)	5 (26.3%)	—
MDS MLD	8 (6.2%)	0 (0.0%)	4 (26.7%)	1 (7.7%)	3 (15.8%)	—
MDS EB 1	11 (8.5%)	6 (7.3%)	0 (0.0%)	3 (23.1%)	2 (10.5%)	—
MDS EB 2	12 (9.3%)	5 (6.1%)	2 (13.3%)	5 (38.5%)	0 (0.0%)	—
AML MRC	2 (1.6%)	0 (0.0%)	1 (6.7%)	0 (0.0%)	1 (5.3%)	—
MDS del5q	1 (0.8%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	1 (5.3%)	—
MDS UNS	1 (0.8%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	1 (5.3%)	—
CMML 2	1 (0.8%)	1 (1.2%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	—
MDS MPN	1 (0.8%)	0 (0.0%)	1 (6.7%)	0 (0.0%)	0 (0.0%)	—
WBC (×10 ⁹ /L)						0.369
Mean (range)	6.4 (1.4–34.6)	6.2 (1.4–32.8)	6.9 (1.7–34.6)	5.1 (1.9–8.9)	7.6 (1.5–18.2)	—
ANC (×10 ⁹ /L)						0.263
Mean (range)	3.7 (0.3–21.1)	3.6 (0.3–18.5)	3.6 (0.3–18.5)	2.7 (0.8–5.6)	4.8 (0.3–12.2)	—
Monocytes (×10 ⁹ /L)						0.419
Mean (range)	0.7 (0–10)	0.6 (0–4.9)	1.1 (0.1–10)	0.5 (0–1)	1.0 (0.1–4.7)	—
HB (g/dL)						0.012
Mean (range)	10.0 (6–14.6)	10 (6–13.6)	10.8 (8–14.2)	8.8 (6–10.9)	9.8 (7–14.6)	—
Platelets (×10 ⁹ /L)						<0.001
Mean (range)	258 (23–1234)	272.3 (57–715)	144.9 (25–261)	152.4 (23–389)	357.7 (36–1234)	—
BM Blasts (%)						<0.001
Mean (range)	3.6 (0.5–20)	2.9 (0.5–19)	5.3 (1.5–18)	6.5 (1–15)	3.6 (1–20)	—
BM RS (%)						<0.001
Mean (range)	37.1 (5–88)	43.8 (9–88)	24.2 (6–73)	25.2 (6–52)	26.8 (5–75)	—
BM Cellularity (%)						0.051
Mean (range)	63 (10–100)	60.2 (20–100)	66 (30–100)	74.6 (50–100)	64.8 (10–100)	—
BM Erythroid cell (%)						0.166
Mean (range)	31.9 (3–83)	33.1 (7.5–83)	25.7 (8.5–43)	36.3 (13.5–65)	28.2 (3–80.5)	—
IPSS-M						<0.001
Very Low	37 (29%)	32 (39%)	2 (13%)	0	3 (16%)	—
Low	51 (39%)	36 (44%)	4 (27%)	2 (15%)	9 (48%)	—
Intermediate low	8 (6%)	3 (4%)	3 (20%)	1 (8%)	1 (5%)	—
Intermediate high	5 (4%)	1 (1%)	3 (20%)	1 (8%)	0	—
High	15 (12%)	8 (10%)	2 (13%)	0 (%)	5 (26%)	—
Very high	13 (10%)	2 (2%)	1 (7%)	9 (69%)	1 (5%)	—
mOS						<0.0001
Median (IQR)	5.5 years (1.7–9.5)	7.6 years (4–9.5)	3.3 years (1–5.9)	0.7 years (0.3–1.7)	4.4 years (1.6–4.9)	—
Total SNV hits						0.034
Mean (range)	3 (0–11)	2.8 (1–6)	4.2 (1–11)	2.7 (1–5)	2.9 (0–9)	—
Total CNV hits						<0.001
Mean (range)	1.1 (0–11)	0.5 (0–8)	0.3 (0–2)	6.2 (2–11)	0.6 (0–2)	—

Abbreviations: AML MRC, acute myeloid neoplasms with myelodysplastic related changes; ANC, absolute neutrophil count; CMML 2, chronic myelomonocytic neoplasms type 2; HB, hemoglobin; IQR, interquartile range; MDS MPN RS T, MDS/myeloproliferative neoplasms with RS and thrombocytosis; MH, multi hit mutated; MLD, multi lineage dysplasia; mOS, median overall survival; NOS, not otherwise specified; SLD, single lineage dysplasia; SNV, single-nucleotide variant; UNS, unspecified; WBC, white blood cell.

was found in 4 cases (3%). *SF3B1*-*SRSF2* comutated cases were classified as NOS in downstream analysis and underwent additional clonal hierarchy studies. The analysis of mutations hierarchical ranks indicated that *SF3B1* and *SRSF2* mutations were comutated at the cellular level in two cases. Single-cell-derived hemopoietic and progenitor colony genotyping experiment confirmed coexistence of *SF3B1*^{K666T}/*SRSF2*^{P95H} and *SF3B1*^{K626S}/*SRSF2*^{P95H} mutations in these

two cases. The other two comutated cases harbor *SF3B1*^{K700E} mutation in the dominant clone, with a small secondary clone harboring a less common *SRSF2* mutation outside the P95 hotspot (Supplementary Fig. S2A and S2B). The latter two cases had gene-expression and alternative splicing profiles similar but not identical to the *SF3B1*-only mutated cases, while the two cases with true concomitant *SF3B1* and *SRSF2* mutation showed a gene-expression and alternative splicing

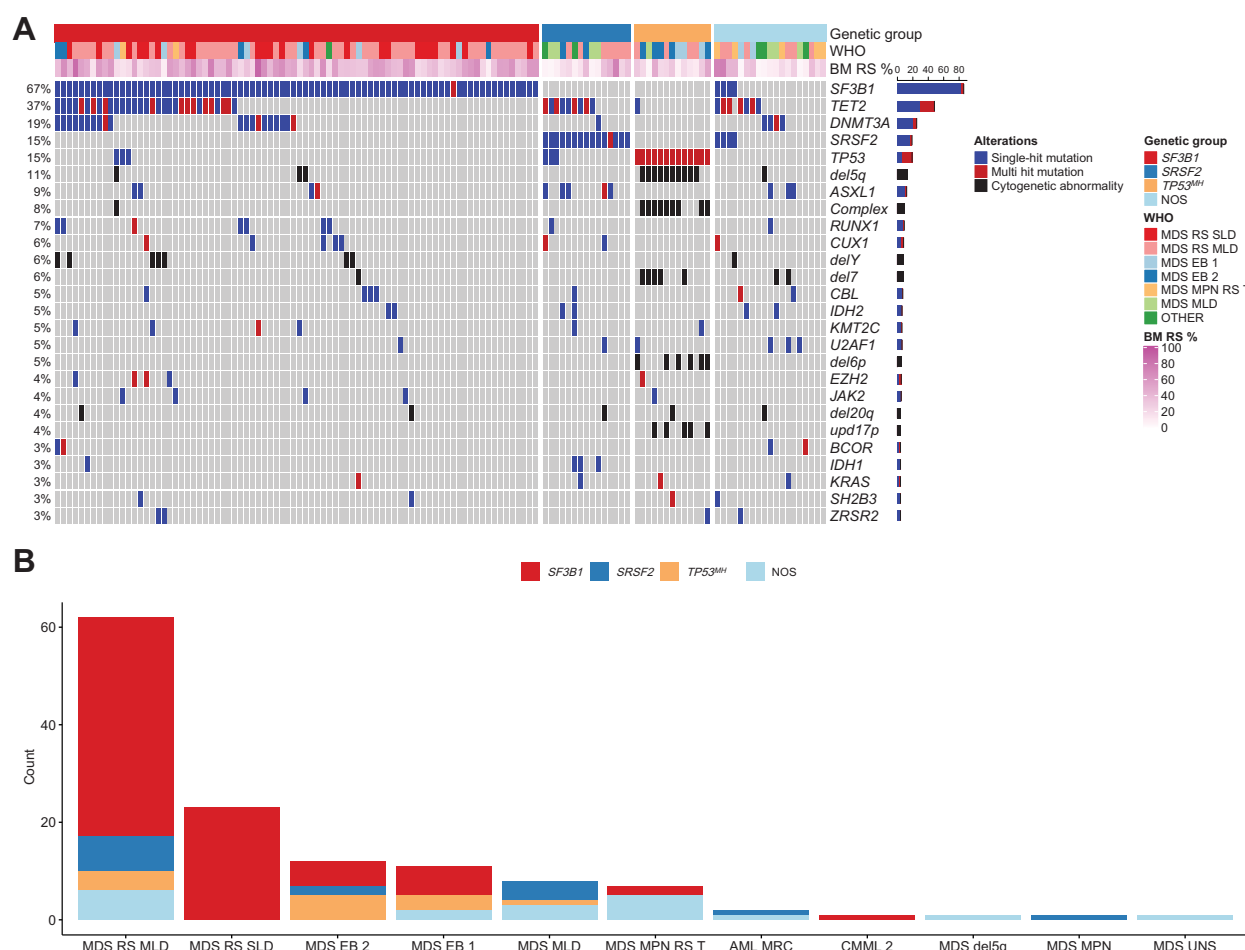


Figure 1.

Mutational landscape of MDS^{RS+}. **A**, Mutation profile displaying clusters identified by unsupervised clustering analysis in MDS^{RS+}. Genetic aberrations and somatic mutations with prevalence >2% are shown. Genes having ≥2 distinct mutations or one mutation associated with chromosomal alterations of the same gene locus are labeled as multihit and distinguished by single-mutation. Genetic subgroup, WHO subtypes, and BM RS percentage (BM RS %) are shown above the heatmap. **B**, Distribution of genetic classes across WHO categories of the MDS^{RS+}. Complex, complex karyotype; del, deletion; mh, multihit; upd, uniparental disomy.

profiles with mixed features in-between single-mutated *SF3B1* and *SRSF2* cases. Given the current limited applicability into clinical practice of hierarchical clonal evaluation and transcriptomic profiling, we cautiously classified these four cases in the NOS subgroup, considering the limited power of any clinical outcome analysis carried out on 4 patients.

Association between mutation profile and clinical features

Mutations in *DNMT3A*, *TET2*, *ASXL1* (usually referred to as DTA mutations) and loss of chromosome Y (-Y) – the most recurrent genetic aberrations in age-related clonal hematopoiesis (37, 41) – cooccurred with *SF3B1* or *SRSF2* in 53% and 75% of cases, respectively. Notably, *TP53*^{MH} cases were depleted for DTA mutations or -Y (OR = 0.07, *P* < 0.001) but enriched for del5q (OR = 94.7, *P* < 0.001), del7q/-7 (OR = 32.9, *P* < 0.001), and complex karyotype (OR = 29.9, *P* < 0.001). No significant difference in age was found among the genetic subgroups.

The four genetic categories (i.e., *SF3B1*, *SRSF2*, *TP53*^{MH}, and NOS) were variably distributed in WHO subcategories (Fig. 1B). MDS^{RS+}

SF3B1^{mutated} accounted for 100% of MDS-RS-SLD (*P* = 0.01), whereas MDS^{RS+} *TP53*^{MH} were enriched in MDS-EB type 2 (MDS-EB-2, *P* = 0.01), but also observed in four cases with myeloblasts < 5% and RS ≥ 15%. When exploring genotype-phenotype associations irrespective of WHO classification, *TP53* (both single and double) and *RUNX1* mutations were found to be the strongest predicting factors of a blastic phenotype (BM blast count ≥ 5%, OR = 6.6 and 6.8, *P* < 0.001 and 0.01, respectively). Thrombocytosis was associated with *JAK2* (OR = 25, *P* < 0.001), but no other independent molecular predictor of proliferative phenotype (e.g., leukocytosis, monocytosis) was identified in this cohort. In *SRSF2* and *TP53*^{MH}-mutated MDS, no survival difference was found when comparing cases with high (≥ 15%), low (5%–14%), RS and historical controls without RS (Supplementary Fig. S3).

Transcriptomic landscape

RNA-Seq analysis of CD34⁺ BM MNC was carried out in all 129 MDS^{RS+} and 10 NBM controls. Median sequencing coverage per patient was 67 million reads (Supplementary Tables S1 and S2). Overall, transcription analysis revealed a continuous spectrum of

changes in transcriptome profiles (Supplementary Results; Supplementary Figs. S4–S7). Unsupervised consensus cluster analysis revealed three stable clusters (Fig. 2A; Supplementary Figures S8–S10). Gene-set enrichment analysis (GSEA) of differentially expressed genes (DEG) among the three groups detected a specific enrichment of genes expressed by immature myeloid progenitors (IMP) in the first cluster (hereafter referred to as IMP group), and erythroblast-megakaryocyte (EMK) precursors in the second cluster (EMK group). These results were in line with a previous transcriptome study in MDS (7). A third cluster was identified displaying intermediate molecular features between EMK and IMP, with a strong upregulation of genes involved in mature myelopoiesis. We referred to this third group as intermediate (INT) group (Fig. 2B; Supplementary Table S3).

Relationship among clinical features, genetic, and transcriptome profiles

Compared with the EMK group, IMP cases had lower BM RS (median 22% vs. 40%, $P < 0.001$; Fig. 2C) and higher BM blast (median 5% vs. 1.5%, $P = 0.002$; Fig. 2D). IMP group was enriched in MDS-EB-2 and MDS with multilineage dysplasia (MDS-MLD, 50% and 62%, $P < 0.001$ and 0.4, respectively) and underrepresented in MDS-RS-SLD/MLD and MDS/MPN-RS-T (21% and 0, respectively, both $P = 0.02$; Fig. 2E). Consistently with transcriptomic analysis, INT signature was associated with an intermediate morphologic phenotype between EMK and IMP groups. No other significant differences were found in age, peripheral blood counts, serum ferritin, serum erythropoietin or BM myeloid/erythroid ratio across the three groups.

Subsequently, we explored the relationship between genomic and transcriptomic classification in the context of MDS^{RS+}. Genetic class-defining allelic burden (median 0.35%; range, 0.02–0.48) did not differ across the genetic and transcriptomic subgroups (Fig. 2F). EMK-like signature was associated with *SF3B1* mutation (OR 2.7, $P = 0.01$; Fig. 2G and H) and, interestingly, INT signature was associated with *TP53*^{MH} mutation (OR 3.5, $P = 0.02$). IMP signature was not restricted to any of the genetic subgroups but was specifically associated with *ASXL1* (OR = 11, $P < 0.001$), *del7q/-7* (OR = 8.9, $P = 0.004$), and *RUNX1* (OR = 6.4, $P = 0.01$) mutations.

However, none of the three transcriptomic signatures was highly specific for any mutation, suggesting that transcriptomic heterogeneity is influenced but not fully explained by genetic abnormalities. Therefore, we carried out supervised DEG analysis across the previously described genetically defined subgroups to get insight into the molecular pathophysiology of MDS^{RS+}.

Genetically defined MDS^{RS+} subgroups display distinct gene expression and alternative splicing profiles

DEG analysis revealed distinct signatures among *SF3B1*, *SRSF2*, and *TP53*^{MH} mutated MDS^{RS+} (Fig. 3A). *ABCB7* – an iron mitochondrial transporter known to be downregulated in MDS-RS-SLD/MLD (6, 9, 10) – was found selectively downregulated in *SF3B1*^{mut} MDS^{RS+} (Fig. 3A; Supplementary Fig. S11). In addition, *SF3B1*^{mut} MDS^{RS+} also displayed significant downregulation of DNA/RNA polymerases (e.g., *DNAJC3*, *PARP2*, *RTF1*) and phosphatases involved in cellular signaling (e.g., *PTPN11*, *ENOPH1*), together with upregulation of genes involved in protein translation. Conversely, MDS^{RS+} harboring *TP53*^{MH} mutation showed strong downregulation of genes involved in protein translation and upregulation of DNA/RNA polymerases and phosphatases, as compared with other MDS^{RS+} and NBM controls. *ABCB7*, phosphatases, DNA/RNA polymerases and protein

translation gene expression was not significantly altered in *SRSF2*^{mut} MDS^{RS+}.

Alternative splicing analysis found that splicing factor mutated MDS^{RS+} had a significantly higher number of alternative splicing events than *TP53*^{MH} subgroup when compared with NBM controls (Fig. 3B; Supplementary Fig. S12). Consistently with previous reports (6, 9), *SF3B1*^{mut} MDS^{RS+} had a specific increase of A3' splice site events, whereas *SRSF2*^{mut} had a substantial decrease in exon skipping events. Missplicing events affecting *ABCB7* and *TMEM14C*, already described in MDS-RS-SLD/MLD (6, 42), were confirmed to be restricted to the *SF3B1*^{mut} subgroup, together with other genes involved in DNA homeostasis (*CHD1L* and *PDS5A*) and in protein translation (*MAT2B* and *TCOF1*). Results from alternative splicing analysis applied to the transcriptomic stratification groups mostly reflected different distribution of splicing factor mutations across the groups; redundancy with genotyping data limited the translational application of these findings in following analyses. Overall, distinct patterns of differentially expressed genes and aberrant splicing events were found in the genetically defined subgroups, providing additional evidence that MDS^{RS+} genetic classification unveils distinct disease entities.

CD34⁺ transcriptome profile reflects HSPC composition in MDS^{RS+}

Considering that GSEA and pathway analyses were significant for transmembrane signaling activity involved in hematopoietic precursors differentiation (as previously shown in Fig. 2B; Supplementary Figs. S6 and S7), and transcriptomic heterogeneity was only partly explained by genetic abnormalities (Fig. 2G), we hypothesized that the BM CD34⁺ transcriptome may reflect composition in hemopoietic stem and progenitor cells (HSPC). To explore this hypothesis and better characterize the relationship between gene and hemopoietic precursor signatures, we used single-cell transcriptomic-based deconvolution to dissect distinct HSPCs in the BM CD34⁺ transcriptome. This technique allowed accurate HSPC quantification by integrating quantitative evaluation of a restricted gene-set specific for each HSPC subtype. Granulocyte/macrophage progenitors (GMP), megakaryocyte/erythroid progenitors (MEP), and hemopoietic stem cells (HSC) were the most represented estimated cell subtypes and accounted for 39%, 35%, and 6% of the total imputed CD34⁺ cell population in MDS^{RS+} (Fig. 4A; Supplementary Results and Supplementary Figs. S13–S17). Among the most represented subpopulations, cells with MEP-like signature showed the highest variability across the transcriptomic groups (median value was 45%, 37%, and 13% in EMK, INT, and IMP group, respectively, $P < 0.0001$; Fig. 4B), with IMP cases displaying very low value of MEP compared with other MDS^{RS+} and NBM controls. Results from RNA deconvolution were validated using multiparameter flow cytometry ($R = 0.9$, $P = 0.0001$; Fig. 4C; Supplementary Fig. S15) in a subset of five cases already included into this study. MEP was also the only estimated population that retained statistical significance when multinomial logistic regression model was built to predict results from transcriptome unsupervised clustering analysis ($P < 0.001$). Cases with BM blasts $\geq 10\%$ displayed significantly low MEP levels compared with other cases, with very high variance especially in cases without blast excess ($< 5\%$, Supplementary Fig. S18). Importantly, MEP-like signature was independent from myeloid/erythroid ratio evaluated on total BM cellularity. These results suggest that MDS^{RS+} transcriptome reflect heterogeneity of the HSPC compartment, simplistically summarized as a decreasing gradient in MEP-like signature across the EMK, INT, and IMP subgroups and not mimicked by standard morphologic features.

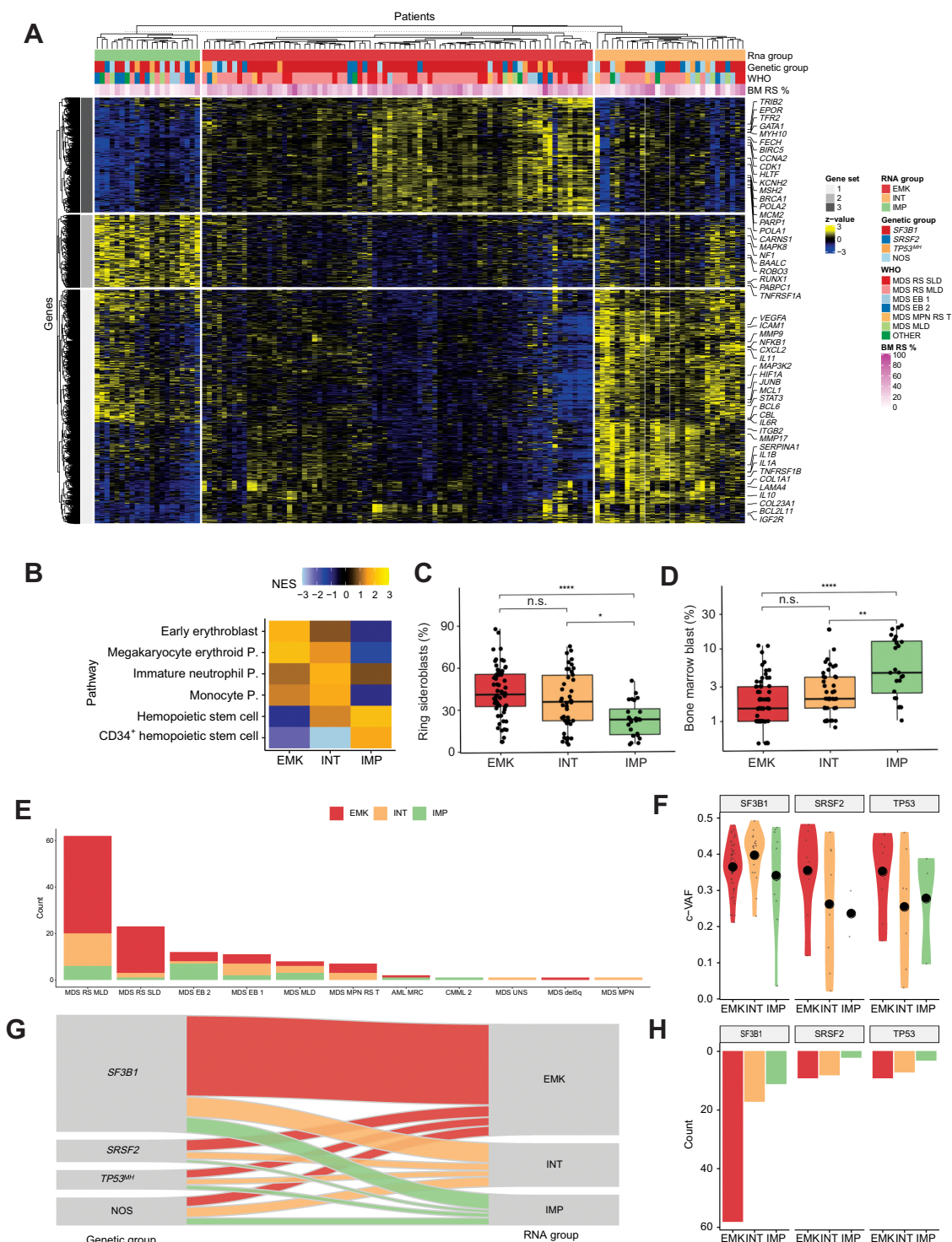


Figure 2.

Transcriptomic landscape of MDS^{RS+}. **A**, Heat map showing expression levels of differentially expressed genes in MDS^{RS+} across the three transcriptomic groups. Rows represent genes and samples are depicted in columns. Covariates display gene expression-based group, genetic group, WHO subtypes and BM RS %. **B**, GSEA on DEGs among the three clusters. Normalized enrichment score (NES) was computed considering the comparison between each group and NBM used as controls. P, progenitor. **C–D**, Comparison of BM RS (C) and blasts (D) distributions across the three groups derived from gene expression unsupervised clustering analysis. **E**, Frequency distribution of transcriptomic classes across WHO categories of the MDS^{RS+} cohort. **F**, Distribution of variant allele frequency (VAF) corrected by CNV/LOH (c-VAF) of *SF3B1*, *SRSF2*, and *TP53*^{MH}, stratified by gene expression groups. Specifically, in case of deletion or LOH, VAF was halved to making it comparable with mutation without CNV/LOH. In case of multiple *TP53* mutations, *TP53*^{MH} c-VAF referred to the second-ranked *TP53* mutation VAF. **G**, Sankey diagram showing the relationship between genomic and transcriptomic classification. Color labels distinguish cases according to the three gene expression groups. **H**, Absolute frequency distribution of *SF3B1*, *SRSF2*, and *TP53*^{MH} mutation, stratified by gene expression group.

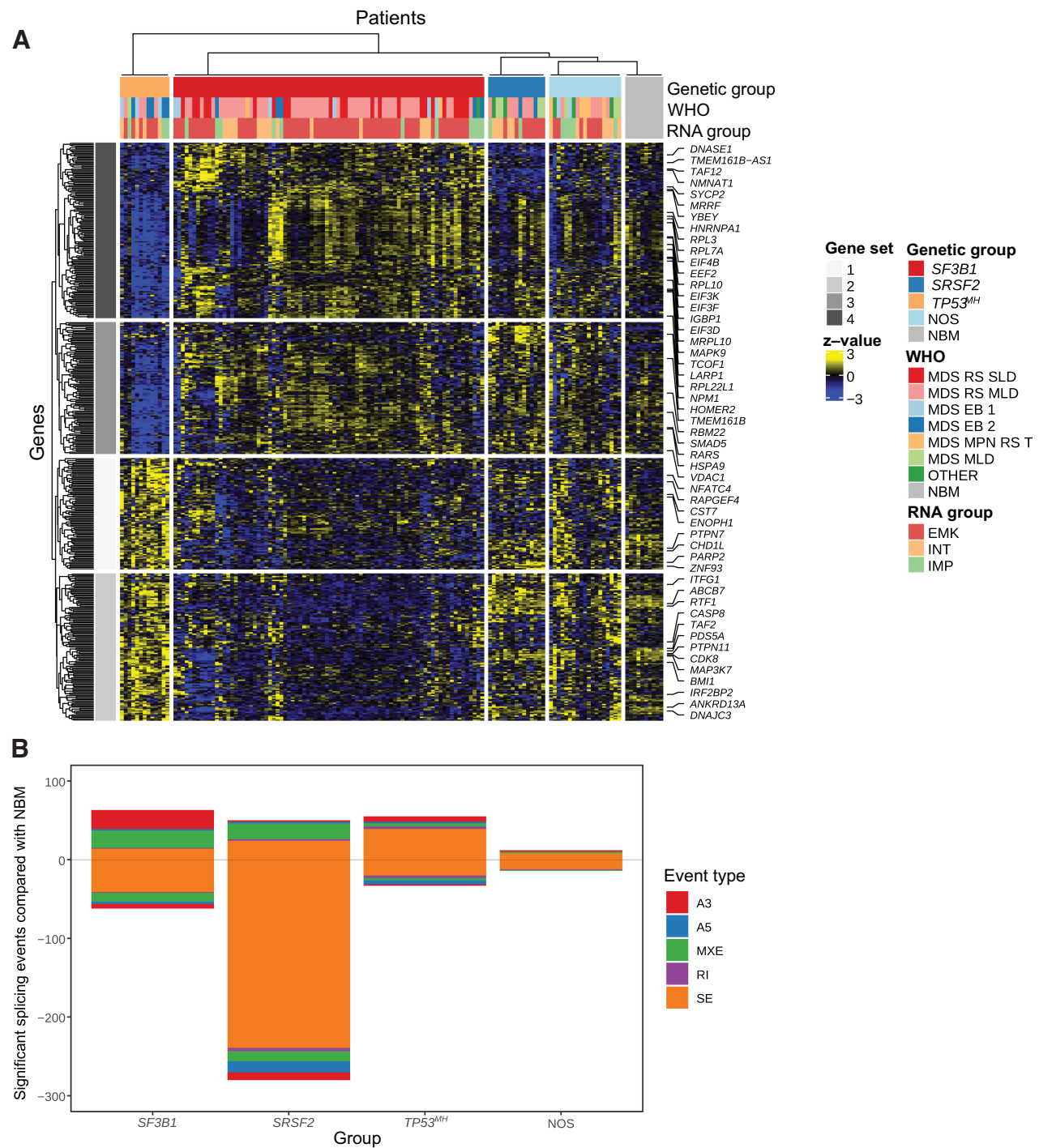


Figure 3. Supervised transcriptome analysis according to MDS^{RS+} genomic classification. **A**, Heat map showing differentially expressed genes among *SF3B1*, *SRSF2*, and *TP53^{MH}* MDS^{RS+} compared with NBM controls. Rows represent genes and columns represent samples. Genetic, WHO, and transcriptomic subtypes are shown as covariates. **B**, Types and numbers of splicing alterations significantly associated with each genetic group. Bars in the top half indicate alternative splicing events that were enriched in MDS^{RS+}, bars in the bottom half indicate alternative splicing events enriched in the healthy donors used as controls, stratified by splicing alterations subtypes.

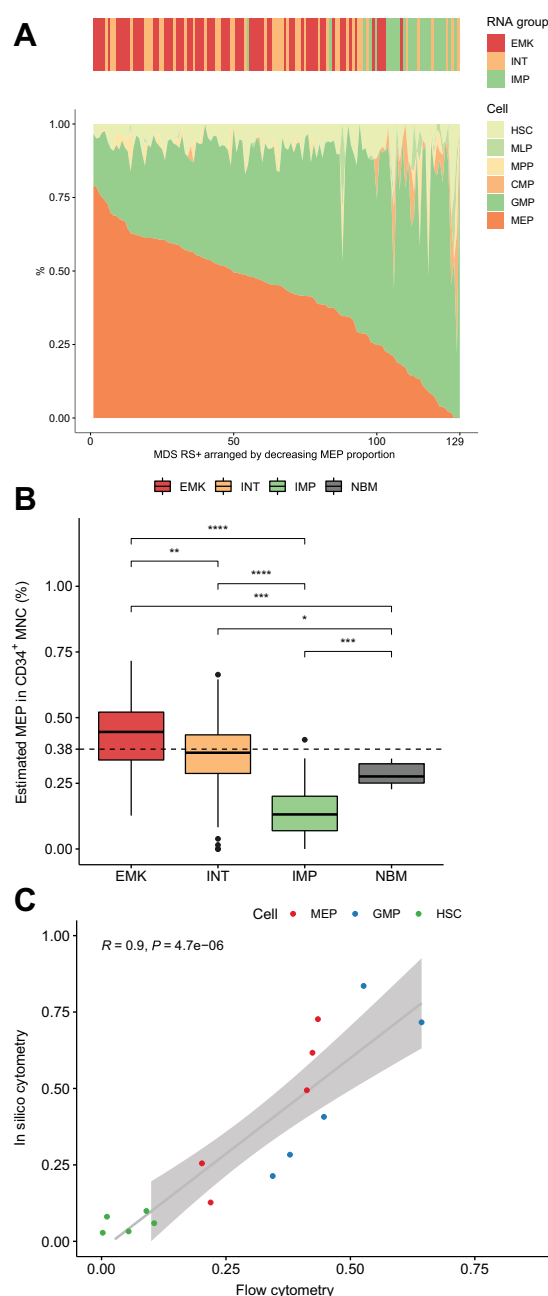


Figure 4.

Digital sorting analysis carried out on bulk CD34⁺ MNC transcriptome from MDS^{RS+} cases. **A**, Digital sorting analysis, gated for HSPCs. Mature cells were gated out to focus only on HSPC compartment (results including mature cells are shown in Supplementary Fig. S13); CMP, common myeloid progenitors; GMP, granulocyte-monocyte progenitors; HSC, hematopoietic stem cells; MLP, multi-lymphoid progenitors; MPP, multipotent progenitors; MEP, megakaryocyte-erythroid progenitors). Cases (x-axis) are sorted by decreasing estimated MEP abundance, showing enriched IMP signature in cases with MEP depletion on right side of the plots. **B**, Relative distribution of estimated MEP in the three transcriptomic groups and (NBM). **C**, Scatter plot comparing results from multiparameter flow (x-axis) and *in silico* cytometry (y-axis). MEP, GMP, and HSC were quantified in the CD34⁺ BM MNCs by means of the two techniques, showing a highly concordance [correlation line and 95% confidence interval (CI) are represented in dark and light gray, respectively, Pearson correlation coefficient = 0.9, $P < 0.001$].

Clinical relevance of combined genomic-transcriptomic profiling

Finally, we explored the predictive value of multiomics characterization on patient outcome. In univariate analysis, age, blast, complete blood count, cytogenetic risk, and IPSS-M score were associated with distinct OS and EFS, together with genetic and transcriptomic classification and estimated MEP proportion (Fig. 5A and B; Supplementary Fig. S19A and S19B, and Supplementary Tables S4 and S5).

Given the reduced applicability in clinical practice of continuous variables as a prognostic factor, we applied maximally selected rank statistics method and found that 38% was the most significant MEP cut-off level in survival univariate analysis (both OS and EFS). MEP $\geq 38\%$ predicted EMK profile with the accuracy of 70% (positive predictive value 64%, negative predictive value 81%) and was consistently associated with longer OS and EFS (Supplementary Fig. S19C and Fig. 5C, respectively; and Supplementary Fig. S20).

Prognostic effect of genetic classification and MEP was also confirmed when only MDS-RS-SLD/MLD were considered (Fig. 5D–F; Supplementary Fig. S19D–S19F). MEP depletion was found to be an independent prognostic factor when the new IPSS-M was included in the multivariable analysis (Fig. 5G–I; Supplementary Figs. S19G–S19I and S21). Similar results were found when we restricted our analysis to the MDS with low blast and SF3B1 mutation, as defined according to the 2022 WHO and ICC new classifications (Supplementary Fig. S22).

Discussion

In this study we explored the clinical relevance of combined genomic/transcriptomic profiling in MDS^{RS+}. To study the association of a RS phenotype with other clinical features in an unbiased manner, we included any MDS cases with ring sideroblast (i.e., $\geq 5\%$) without limitation to MDS specific subclasses as defined by recent classifications of myeloid neoplasm. Mutations in *SF3B1*, *SRSF2* or *TP53^{MH}* were found to be the most relevant drivers in 89% of all MDS^{RS+}. Unsupervised clustering of genetic alterations confirmed that these three genotypes cooccur very rarely. Indeed, *TP53^{MH}* never cooccurred with *SF3B1* or *SRSF2* mutation, prompting future studies evaluating RNA splicing impairment in *TP53^{MH}*-mutant cells. Coexistence of *SF3B1*/*SRSF2* mutation was very rare (3%) and, when cooccurring in the same cell, involved a permissive combination of less common alleles with reduced effects on RNA splicing (43).

The high allelic burden of *SF3B1*, *SRSF2*, and *TP53^{MH}* mutations suggests that mutations in these three genes may play a driving role in pathogenesis of MDS^{RS+}, while comutation patterns contribute to further shape the clinical phenotype of the disease, as suggested by previous reports on myeloid neoplasms (19, 22). Survival analysis based on the resulting genetic stratification confirmed that the benign behavior of MDS-LB-RS is limited to case with *SF3B1* mutation but without concomitant high-risk cytogenetic aberrations or mutations, arguing against the accuracy of RS as a surrogate marker for *SF3B1* mutation in the recent WHO diagnostic criteria (12). In addition, *TP53^{MH}* and *SRSF2* MDS^{RS+} survival was similar to historical controls (i.e., MDS and MDS/MPN) with RS $< 5\%$ (22, 23), and no survival difference was found between having RS between 5% and 14% or $\geq 15\%$ in *SF3B1^{wt}* cases (as it was already known for MDS^{*SF3B1*+}). Our results support the classification of *SF3B1^{wt}* MDS-LB-RS cases within MDS-NOS irrespective of the number of RS, and confirm that the definition of MDS-LB-RS should be only limited to the cases with *SF3B1* mutation, as proposed by new ICC 2022 classification of myeloid neoplasms (11). Further studies evaluating clinical

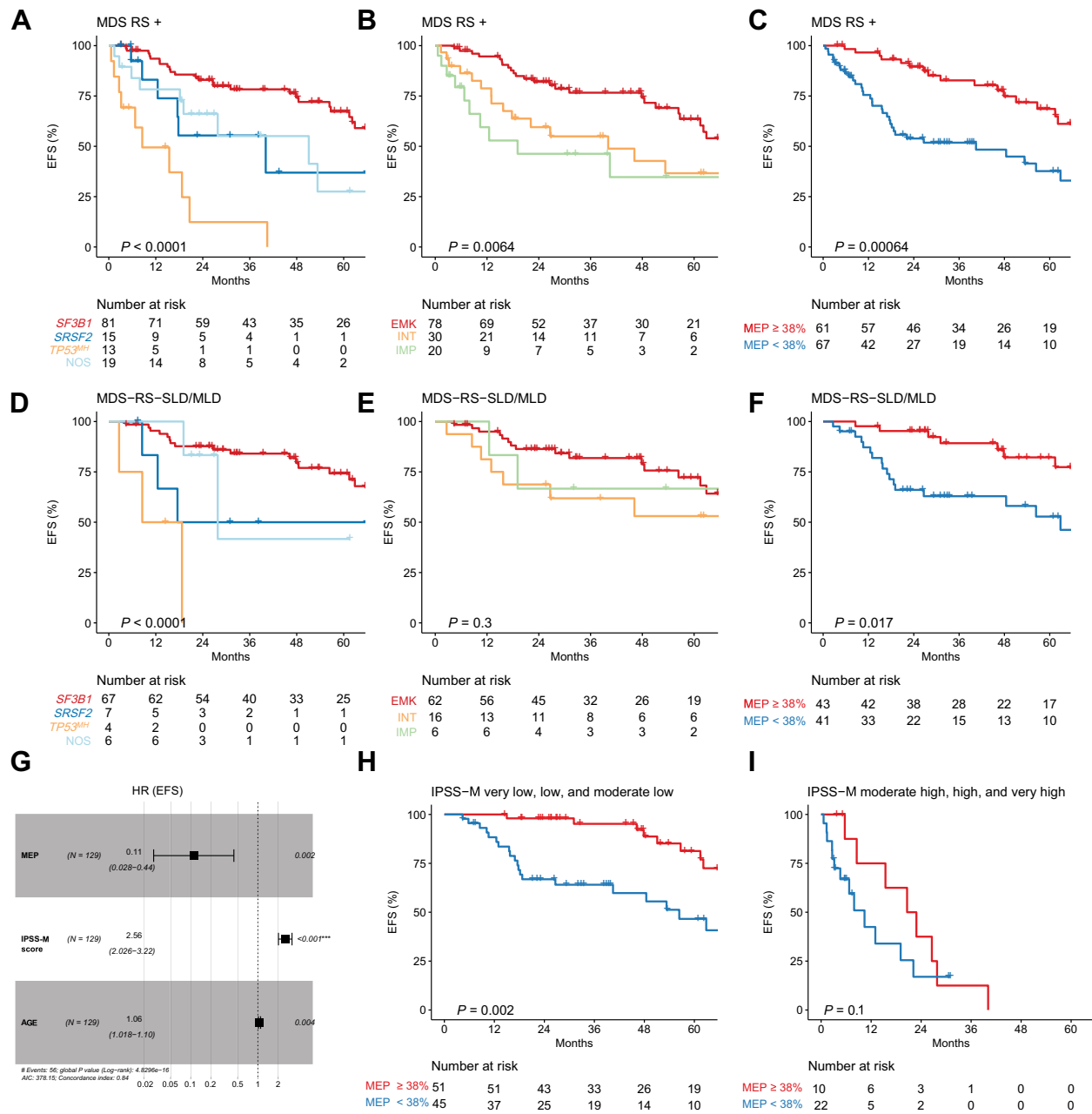


Figure 5.

Prognostic effect of genomic and transcriptomic analyses on MDS^{RS+} outcome. EFS stratified by genomic (A), transcriptomic classification (B), and estimated MEP percentage (C) in all MDS^{RS+} (A-C) and MDS-RS-SLD/MLD only (D-F). Multivariable Cox proportional hazard model for EFS in all MDS^{RS+} including age, IPSS-M score, and estimated MEP percentage as continuous variables (G). EFS stratified by estimated MEP percentage and IPSS-M risk category (H and I) (full representation of the six IPSS-M categories shown in Supplementary Fig. S21A).

characteristics and outcome of the rare cases with concomitant *SF3B1*/*SRSF2* mutations are warranted, giving the relevant implication on prognostic stratification and the current lack of evidences on this rare molecular condition.

Transcriptome analysis of CD34⁺ enriched BM cells revealed that transcriptomic heterogeneity was mainly led by genes involved in inflammation, transmembrane signaling and extracellular matrix organization, highlighting the interaction between bone marrow

microenvironment and HSPC in MDS^{RS+}. Consistent with a previous study (7), we identified a gene-expression signature characterized by upregulation of genes involved in erythrocyte-megakaryocytic differentiation (i.e., EMK signature). Compared with previous report – that distinguished only two groups – we found one additional stratification in the non-EMK group, where we distinguished IMP from INT signature. Both INT and IMP signatures included genes upregulated in myeloid progenitors, while

INT signature displayed a specific upregulation of genes involved in interleukin signaling and inflammation.

Given the differential expression pattern involving genes related to hematopoietic precursor differentiation, we hypothesized that signatures captured by bulk CD34⁺ MNC RNA-seq reflected a distinct HSPC composition. This hypothesis was confirmed by digital sorting and showed that EMK group had higher proportion of MEP as compared IMP/INT cases and NBM controls that was independent from myeloid/erythroid ratio evaluated on total BM cellularity. To translate these findings into a clinically useful tool, we explored the prognostic value of MEP proportion on survival analysis and found that low MEP predicted poor survival, with 38% threshold as the optimal cut-off value. Multivariable survival analysis showed that MEP proportion and IPSS-M score independently contributed to survival prediction. These results are consistent with previous studies evaluating CD34⁺ BM MNCs by flow cytometry analysis (44) and gene array (45) reporting MEP decrease in high-risk MDS, and were found to be significant even when the IPSS-M risk score was taken into account, in particular in low/moderate-low risk classes, where clinical decision making may be a challenge. Overall, our results suggest that MEP (Lin[−]CD34⁺CD38⁺CD123[−]CD45RA[−]) quantification by diagnostic flow cytometry analysis on BM aspirate at diagnosis should be explored as a surrogate marker for high-risk transcriptomic profile in larger prospective trials aiming to improve MDS prognostication.

Furthermore, results from differential gene expression and alternative splicing analyses supported distinction of *SF3B1*^{mut} from *SRSF2*^{mut} and *TP53*^{MH} MDS^{RS+} and suggests that RS development has a distinct pathobiology depending on the genetic driver. Indeed, *SF3B1*^{mut} and *SRSF2*^{mut} MDS^{RS+} displayed distinct iron and mitochondrial homeostasis regulators among differentially expressed and differentially spliced genes. Impaired expression of *ABCB7* and *TMEM14C* was found to be restricted to *SF3B1*^{mut} MDS^{RS+} (42), whereas *CASP8* and *ESWRI* – both known to be involved in mitochondrial homeostasis (46, 47) – were specifically misspliced in *SRSF2*^{mut} cases. Conversely, *TP53*^{MH} MDS^{RS+} exhibited a selective dysregulation of genes involved in protein synthesis, suggesting that mitochondrial iron overload in these cases might be consequent to defective iron incorporation into hemoglobin rather than increased mitochondrial iron uptake. Further studies to explore the interconnection between clonal hematopoiesis pathophysiology and iron metabolism may elucidate the molecular mechanisms leading to impaired iron cellular balance in distinct genetically-defined MDS^{RS+} subtypes.

In conclusion, integrated genomic-transcriptomic analysis supports the distinction of *SF3B1*^{mut}, *SRSF2*^{mut}, *TP53*^{MH}, and NOS MDS^{RS+} and introduces HSPC transcriptomes as a novel and independent prognostic variable. Our results provide essential input on the molecular basis of *SF3B1*-unmutated MDS^{RS+}, and support *SF3B1*^{mut} MDS^{RS+} as a stand-alone category in the myeloid neoplasms classifications.

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Authors' Disclosures

E. Bernard reports personal fees from Pfizer and grants from Edward P Evans Foundation outside the submitted work. S.E. Reinsbach reports grants from Knut and Alice Wallenberg Foundation during the conduct of the study. E. Papaemmanuil reports grants from MDS Foundation outside the submitted work; is a founder, equity holders and holds fiduciary roles in Isabl Inc.; is also a Scientific Advisor for Tensixteen Bio and accordingly has shares in the company; and has received travel funding from Illumina. E. Hellström-Lindberg reports grants from The Swedish Wallenberg Foundation during the conduct of the study. No disclosures were reported by the other authors.

Authors' Contributions

G. Todisco: Conceptualization, data curation, formal analysis, investigation, visualization, methodology, writing—original draft, project administration, writing—review and editing. **M. Creignou:** Data curation, formal analysis. **E. Bernard:** Resources, data curation, formal analysis, methodology. **A. Björklund:** Data curation, formal analysis, investigation. **P.L. Moura:** Data curation, formal analysis, investigation. **B. Tesi:** Data curation, formal analysis, investigation. **T. Mortera Blanco:** Data curation, formal analysis, investigation. **B. Sander:** Data curation, formal analysis, investigation. **M. Jansson:** Data curation, formal analysis, investigation. **G. Walldin:** Data curation. **I. Barbosa:** Data curation, formal analysis, investigation. **S.E. Reinsbach:** Data curation, formal analysis, investigation. **I.J. Hofman:** Data curation, formal analysis, investigation. **C. Nilsson:** Data curation, formal analysis. **T. Yoshizato:** Data curation, formal analysis, investigation. **M. Dimitriou:** Data curation, formal analysis, investigation. **D. Chang:** Data curation. **S. Olafsdottir:** Data curation. **S. Venckute Larsson:** Data curation. **M. Tobinsson:** Data curation, formal analysis. **L. Malcovati:** Supervision, funding acquisition, methodology. **P. Woll:** Supervision, methodology. **S.W. Jacobsen:** Supervision, methodology. **E. Papaemmanuil:** Supervision, methodology. **E. Hellström-Lindberg:** Conceptualization, resources, funding acquisition, methodology, writing—original draft, project administration, writing—review and editing.

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Note

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