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Immune phenotypic analysis of erythroid dysplasia in myelodysplastic syndromes. A report from the IMDSFlow working group

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ABSTRACT

Current recommendations for diagnosing myelodysplastic syndromes endorse flow cytometry as an informative tool. Most flow cytometry protocols focus on the analysis of progenitor cells and the evaluation of the maturing myelomonocytic lineage. However, one of the most frequently observed features of myelodysplastic syndromes is anemia, which may be associated with dyserythropoiesis. Therefore, analysis of changes in flow cytometry features of nucleated erythroid cells may complement current flow cytometry tools. The multicenter study within the IMDSFlow Working Group, reported herein, focused on defining flow cytometry parameters that enable discrimination of dyserythropoiesis associated with myelodysplastic syndromes from non-clonal cytopenias. Data from a learning cohort were compared between myelodysplasia and controls, and results were validated in a separate cohort. The learning cohort comprised 245 myelodysplasia cases, 290 pathological, and 142 normal controls; the validation cohort comprised 129 myelodysplasia cases, 153 pathological, and 49 normal controls. Multivariate logistic regression analysis performed in the learning cohort revealed that analysis of expression of CD36 and CD71 (expressed as coefficient of variation), in combination with CD71 fluorescence intensity and the percentage of CD117+ erythroid progenitors provided the best discrimination between myelodysplastic syndromes and non-clonal cytopenias (specificity 90%; 95% confidence interval: 84–94%). The high specificity of this marker set was confirmed in the validation cohort (92%; 95% confidence interval: 86–97%). This erythroid flow cytometry marker combination may improve the evaluation of cytopenic cases with suspected myelodysplasia, particularly when combined with flow cytometry assessment of the myelomonocytic lineage.
Introduction

Discriminating between cytopenia due to myelodysplastic syndromes (MDS) and cytopenia due to other (non-clonal) causes can be challenging, especially when dysplasia as assessed by cytomorphology does not fulfill the diagnostic criteria of MDS according to WHO, and when other MDS-associated features are absent (e.g., >15% ring sideroblasts (RS) and/or cytogenetic aberrations). Current recommendations for the diagnosis of MDS endorse flow cytometry (FC) as a valuable additional diagnostic tool. In this respect, it has been recommended to follow the guidelines set down by the International/European LeukemiaNet Working Group for FC in MDS (IMDSFlow).

Despite the fact that FC for MDS correlates with cytomorphology, the sensitivity of current validated FC scores for diagnosing MDS requires improvement. So far, most of the designed FC scores have comprised the analysis of the (im)mature myelomonocytic lineage with a median sensitivity of 75% for identifying MDS (median specificity, 94%; see Wouters et al 4). Since anemia is frequently observed in MDS, often accompanied by erythroid dysplasia, analysis of immunophenotypic changes of nucleated erythroid cells (NEC) may complement current FC analysis. Thus far, this has not been studied in great detail. Integration of results from analysis of the erythroid lineage to the primarily myelomonocytic and progenitor cell-based FC scores may improve sensitivity of FC analysis in MDS.

Incorporating erythroid markers in FC protocols requires knowledge of normal erythroid differentiation, and of potential aberrancies and pitfalls. The characteristic morphological stages of normal erythroid differentiation are reflected by their light scatter properties and by their differential expression of CD45, CD117, CD105, CD56, CD71 and/or CD235a. Some of the FC aberrancies that have been reported to reflect MDS-related dyserythropoiesis are: a) an increased number of NEC within total nucleated cells; b) an altered proportion of consecutive erythroid differentiation stages, such as an increased number of immature erythroid cells (CD117+ and/or CD105+) or, by contrast, a decrease in erythroid progenitors; c) an abnormal pattern of CD71 versus CD235a; d) a reduced expression of CD71 and/or CD56; and e) an overexpression of CD105. Most of these aberrancies are present in 70–80% of MDS cases.

However, a number of features may be shared across the spectrum of non-clonal cytopenias. The multicenter study reported herein focused on defining the erythroid FC parameters that enable distinction of dyserythropoiesis associated with MDS from non-clonal cytopenias. Hereto, data from a learning cohort were compared between MDS patients and controls, and the results were validated in a separate cohort.

Methods

MDS patients and controls

Nineteen centers (members of the IMDSFlow group) collected FC data on the erythroid lineage in low grade MDS cases (<5% blasts) and controls. Data were acquired from bone marrow samples taken from 1008 patients and healthy controls after informed consent in accordance with the Declaration of Helsinki; where required, local ethics committee approval was obtained. The learning cohort comprised 677 cases (18 centers, data collected between October 2012 and September 2013), and the validation cohort comprised 331 cases (9 centers, data collected between December 2013 and April 2014). Inclusion criterion for pathological controls was cytopenia not associated with MDS. In total, data on 374 MDS cases, 443 pathological and 191 normal controls were collected (specified in Tables 1 and 2). Information regarding age, gender, cytomorphology and cytogenetics was requested. One center with limited access to cytomorphology results only included MDS patients with typical features of MDS as the presence of more than 15% RS and/or MDS-associated cytogenetic anomalies. In 825/847 MDS cases, sub-classification according to the WHO-2008 was provided. The median contribution per center to the total study cohort was 47 cases (range 6–100); the median number of erythroid FC markers analyzed per case was 7 (range 1–9 of 10 proposed markers).

Sample preparation and antibody combinations

Flow cytometric analysis in MDS requires the removal of mature, enucleated erythrocytes through the use of lysis protocols. The vast majority of centers used ammonium chloride-based solutions, either home-made or commercial (e.g., PharmLyse; BD Biosciences, San Jose, CA); two centers used FACSLyse (BD Biosciences), and one other used VersaLyse (Beckman Coulter, Miami, FL). FACSLyse contains a fixative, whereas VersaLyse is recommended for use with a fixative when the sample contains anticoagulants other than EDTA. The duration of lysis and temperature varied among centers (5–25 minutes and 4–37°C, respectively), but most lysed for 10 minutes (n=10) at room temperature (n=16). Two centers reported the use of an additional fixative in their staining protocols, both in combination with an ammonium chloride-based lysing solution. Detailed information can be found in Online Supplementary Information. Most centers used the IMDSFlow-recommended stain-lyse-wash procedure; five centers performed stain-lyse-wash. Antibody combinations were similar between centers, but clones and fluorochromes differed. Most centers used a backbone of CD45 and CD34 and/or CD117, and added antibodies such as CD235a, CD71, CD36, and CD105. Examples of antibody combinations and panels have been described previously. Nuclear dyes were not routinely included in the panels, and only one center applied the live/dead stain 7-AAD. The flow cytometers used included: FACSCalibur (BD Biosciences; n=3); FACS CANTO-II (BD Biosciences; n=10); a combination of FACSCalibur and FACS CANTO-II (both BD Biosciences; n=2); and Navios (Beckman Coulter; n=4). Panels comprised 4-, 5-, 6- and/or 10-color FC; WinList7.0 (Verity Software House, Topsham, ME), Kaluza (Beckman Coulter), CellQuestPro, FACS DIVA (both BD Biosciences), and/or Infinicyt (Cytognos, Salamanca, Spain) software packages were used for data analysis.

Gating strategy and data collection

The gating strategy was discussed during the IMDSFlow meeting in 2011 and re-evaluated in 2012. All participants performed FC analysis of the erythroid lineage defined as CD45<sub>dim</sub>-to-negative and SSC<sub>low</sub>-to-intermediate. It is noteworthy that the initially proposed gating strategy (erythroid lineage defined by CD45 negativity) was altered to include early erythroid precursors that are within the CD45<sub>dim</sub> cell population. Six or more color panels enabled the inclusion of a myeloid-defining marker such as CD13 or CD33, and a more accurate separation of myeloid and erythroid progenitors. Moreover, to exclude platelets and
platelet aggregates, a combination of scatter properties and CD36<sup>high</sup>/CD71<sup>-dim</sup> was suggested. The final gating strategy was distributed among all centers (detailed information in Online Supplementary Information).

The following parameters were collected: the percentage of NEC within all nucleated cells; the expression pattern of CD71 versus CD235a; the percentage of CD71<sup>-dim</sup>/CD235a<sup>-dim</sup> cells within the CD71/CD235a pattern; CD71 and CD36 expression levels; the percentage of CD11<sup>-dim</sup> cells in the erythroid compartment; CD105 expression level and the percentage of CD105<sup>-dim</sup> cells in the erythroid compartment. Recent knowledge, such as the finding that CD71 and CD36 expression represented as CV is statistically more significant than when represented as mean fluorescence intensity (MFI), led to adjustments in the initially proposed protocol and, hence, reanalysis of the list mode data files by the individual centers. Gating strategies and analyzed parameters are shown in the Online Supplementary Information with FC plots of MDS in comparison to normal subjects.

Statistical analyses

Due to differences in sample processing, instrument settings, clones, and fluorochromes between centers, the expression levels of CD71, CD36, and CD105 varied. Therefore, the median expression levels of CD36, CD71, and CD105 in the subset of normal bone marrow samples were calculated for each individual center. Expression levels were then normalized against the median value for that particular marker for each center separately. Patient and control groups were compared using the Kruskal Wallis test for continuous data, and the Chi-square or the Fisher’s exact test for dichotomous data. Correlations between certain markers, and between markers and age, were analyzed using Spearman’s rank correlation coefficients. Comparing single parameters between MDS and control groups demonstrated that results overlapped (results section figure 2); hence, receiver-operator-curve (ROC) analyses did not yield applicable cut-offs. Therefore, cut-off values for aberrations were based on the 10th and/or 90th percentile of the data of pathological controls in the learning cohort. Multivariate logistic regression analyses were performed to determine the erythroid markers that discriminate between pathological controls and MDS; data were analyzed dichotomously. All variables that displayed a univariate difference of P<0.1 were included in a backward selection procedure based on the Likelihood Ratio score. Regression coefficients of the variables in the final model were used to define the weight of these markers in a descriptive score for dyserythropoiesis. Cut-off level of the score indicating MDS-associated erythroid aberrances was determined based on the total weight of these variables and a specificity of at least 90%. The sensitivity and specificity of the marker combination were calculated to illustrate predictive accuracy. The data were analyzed using SPSS 20.0 (IBM Corp, Armonk, NY), and GraphPad 6.0 software (La Jolla, CA). P-values <0.05 were considered significant.

<table>
<thead>
<tr>
<th>Table 1. Characteristics of MDS patients and controls in the learning and validation cohorts.</th>
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<tbody>
<tr>
<td>Learning cohort</td>
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<tr>
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</tr>
<tr>
<td>Normal</td>
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<td>n</td>
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<tr>
<td>age</td>
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<td>male/female</td>
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<tr>
<td>Pathological controls</td>
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<tr>
<td>n</td>
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<tr>
<td>age</td>
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<td>male/female</td>
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<tr>
<td>MDS</td>
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<td>n</td>
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<td>age</td>
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<td>male/female</td>
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<tr>
<td>Comparison of age normals vs. pathological controls</td>
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<tr>
<td>pathological controls vs. MDS</td>
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<tr>
<td>Comparison of gender</td>
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</tbody>
</table>

*Data are expressed as the median and range; abbreviations: MDS: myelodysplastic syndromes; n: number; n.s.: not significant.

<table>
<thead>
<tr>
<th>Table 2. Subcategories of MDS and pathological controls in the learning and validation cohort.</th>
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<tbody>
<tr>
<td>Learning cohort</td>
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<tr>
<td>MDS subcategories</td>
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<tr>
<td>RCUD</td>
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<tr>
<td>RARS</td>
</tr>
<tr>
<td>RARS&lt;sup&gt;-t&lt;/sup&gt;</td>
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<tr>
<td>RCMD</td>
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<tr>
<td>del(5q)</td>
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<tr>
<td>MDS-U</td>
</tr>
<tr>
<td>other</td>
</tr>
<tr>
<td>not specified</td>
</tr>
<tr>
<td>Subcategories of pathological controls</td>
</tr>
<tr>
<td>iron deficiency anemia</td>
</tr>
<tr>
<td>anemia in chronic disease&lt;sup&gt;†&lt;/sup&gt;</td>
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<tr>
<td>vitamin B12/folic acid deficiencies</td>
</tr>
<tr>
<td>anemia in auto-immune diseases&lt;sup&gt;†&lt;/sup&gt;</td>
</tr>
<tr>
<td>anemia due to renal failure</td>
</tr>
<tr>
<td>anemia other</td>
</tr>
<tr>
<td>cytopenia associated with marrow infiltration</td>
</tr>
<tr>
<td>cytopenia induced by chemotherapy or medication or post-SCT</td>
</tr>
<tr>
<td>TIP or neutropenia or</td>
</tr>
<tr>
<td>auto immune cytopenia NOS</td>
</tr>
<tr>
<td>reactive conditions or</td>
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<tr>
<td>cytopenia induced by infections</td>
</tr>
<tr>
<td>normal bone marrow, peripheral cytopenia</td>
</tr>
<tr>
<td>other than defined subcategories</td>
</tr>
<tr>
<td>inconclusive</td>
</tr>
<tr>
<td>non clonal cytopenia NOS</td>
</tr>
<tr>
<td>ET, PV, primary myelofibrosis</td>
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<tr>
<td>PNH</td>
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<tr>
<td>AA</td>
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</tbody>
</table>

Values between brackets represent the relative distribution within MDS or pathological control subgroups; <sup>†</sup>other than defined subcategories; AA: aplastic anemia; ET: essential thrombocythemia; MDS: myelodysplastic syndromes; MDS-U: MDS unclassifiable; NOS: not otherwise specified; PNH: paroxysmal nocturnal hemoglobinuria; PV: polycythemia vera; RARS: refractory anemia with ring sideroblasts; RARS<sup>-t</sup>: RARS and thrombocytosis; RCMD: refractory cytopenia with multilineage dysplasia; RCUD: refractory cytopenia with unilineage dysplasia; del(5q): MDS with isolated del(5q). Comparison of the distribution of MDS subsets among the learning and validation cohorts did not differ (χ² test; P<0.51); the distribution of subsets of pathological controls did not differ (P>0.001). The subcategories “anemia in chronic disease” comprises iron incorporation disorders; bowel diseases, diabetes, etc., “anemia in auto-immune diseases” comprises AIHA, AITP, Rheumatoid Arthritis, SLE, etc.; “anemia other” comprises, among others, cases of normocytic anemia, anemia unexplained.
Results

Flow cytometric analysis of the erythroid lineage in normal bone marrow samples: comparison of results from participating centers

Discrepancies in erythroid analysis between centers (and samples) can occur at several levels: a) sample quality (e.g., hemodilution); b) sample preparation (e.g., lysing procedure); c) data acquisition (e.g., acquisition rate and threshold of forward light scatter); and d) degree of adherence to the proposed gating strategy. Therefore, we first compared the FC results for normal bone marrow samples (learning cohort) between centers in terms of each defined marker. The percentage of NEC was highly diverse among centers; yet, it seemed to be independent of the lysing method applied (Online Supplementary Figure S1). Similarly, the percentage of CD71dim cells differed largely between centers.

Furthermore, two centers reported higher percentages of CD117+ erythroid progenitors (up to 50% within the NEC) than the other centers (<15%). Results for one center could be explained by their stringent lysing procedures (i.e., 15 minutes at 37°C) which removed more mature (orthochromatic and polychromatic) erythroblasts resulting in a relative increase in early progenitors (data not shown). To circumvent the issues regarding differences in percentages of erythroid (sub)populations between centers, the percentages of NEC, CD117+ and CD105+ erythroid progenitors were also normalized as described for antigen expression levels (see Material and Methods statistics section); these are further referred to as relative percentages, i.e., relative to the median percentage in normal bone marrow samples (Online Supplementary Figure S2). CD71dim cells were rarely seen in normal controls; therefore, results for the percentage CD71dim could not be normalized.

Figure 1. Flow cytometric profiles of normal erythroid differentiation. Early erythroid precursors are defined as CD45−/SSC−/CD34+/CD117+/CD105−/CD235a−, proerythroblasts as CD45−/SSC−/CD34+/CD117+/CD105−/CD235a−, basophillic erythroblasts as CD45−/SSC−/CD34+/CD117+/CD105−/CD235a−, polychromatic erythroblasts as CD45−/SSC+/CD34+/CD117+/CD105−/CD235a− and orthochromatic erythroblasts CD45−/SSC+/CD34+/CD117−/CD105−/CD235a−. Indicated colors reflecting erythroid subsets are not visible in the CD71 vs. CD235a plot (Fig 1F). Herein, pink colored cells represent the total erythroid lineage in this plot. Mature erythrocytes (CD45−/CD36−/CD71−/CD235a−) can be seen in improperly lysed cell preparations (Figure 1F). Reticulocytes are not covered in these graphs, but they may appear as CD71−CD235a− in non-lysed cell preparations. Myeloid progenitors are CD34+/CD117+/HLA-DR+/CD105− (Figure 1C and D); these cells have slightly higher CD45 expression than erythroid precursors; moreover, in contrast to myeloid progenitors erythroid cells do not express HLA-DR (adapted from references 14-16).
Erythroid aberrancies that may discriminate between MDS and pathological controls

Next, all FC-erythroid parameters in the learning cohort were compared between MDS and controls. The results from the learning cohort are summarized in Figure 2 (P-values in Online Supplementary Table S1). The relative percentage of NEC within the total nucleated cell population was significantly higher in MDS than in the pathological and normal controls (P<0.001). Similarly, the CD71-CD235a differentiation pattern was more frequently considered aberrant in MDS (65%, 109/167 cases) than in pathological and normal controls (18% (44/254) and 3.7% (5/154) of cases, respectively (P<0.001)). To objectify the evaluation of this pattern, we analyzed its components separately. CD71 expression was analyzed in terms of MFI, CV, and the presence of a subpopulation with reduced CD71 expression (CD71dim). The relative CD71 MFI was significantly reduced in MDS, whereas the relative CV for CD71 and the percentage of CD71dim cells were significantly higher than those in both control groups (P<0.001); no significant differences were observed between the pathological controls and normal bone marrow samples. The expression of CD235a largely depends on the success of removing mature erythrocytes from a sample. Moreover, membrane fragments of lysed erythrocytes may stick to other cells in the analysis sample, mimicking positivity. Hence, this parameter was considered too unreliable for evaluation when considered individually.

The percentage of immature erythroid progenitors can also affect the appearance of the differentiation pattern. Analysis of the relative percentage of CD117+ (and CD105+) erythroid progenitor cells revealed a broader

Figure 2. Distribution of erythroid markers analyzed by flow cytometry among MDS patients and controls within the learning cohort. Results of the analysis of indicated markers of the erythroid lineage are plotted along the X-axes: relative (rel.) percentages of nucleated erythroid cells (NEC); rel. mean fluorescence intensity (MFI) for CD36, CD71, and CD105; rel. coefficient of variation (CV) of CD36 and CD71; and rel. percentages of CD117+ and CD105+ erythroid progenitors. Relative frequencies (as percentage of the MDS or control cohort for a particular marker) are depicted along the Y-axes. Dotted lines represent results for normal bone marrow (NBM) samples, dashed lines pathological controls (PC) and solid lines MDS cases. P-values of comparison between groups are depicted: **: <0.001, *: <0.05, ns: not significant (Kruskal Wallis test). Grey boxes indicate reference ranges for the analyzed markers as defined by 10th and 90th percentiles of pathological controls. Scatterplots of results for the markers (depicted here as frequency histograms) that were selected as FC-markers for erythroid dysplasia from the multivariate analysis are depicted in Online Supplementary Figure S3.
range of these cells in MDS, although not significantly different from the control groups.

Relative expression of CD105 was either increased or decreased in MDS. Nonetheless, CD105 expression did not discriminate between MDS and pathological controls.

Similar to that for CD71, the relative MFI of CD36 was significantly lower and the relative CV for CD36 was significantly higher in MDS than in the control groups.

To summarize, the markers that showed a significantly different distribution in MDS as compared to controls were: the relative percentage of NEC and the percentage of CD71dim cells (increased in MDS); the relative MFI of CD71 and CD36 (decreased in MDS); and the relative CV for CD71 and CD36 (increased in MDS); Figure 2.

Selection of a combination of erythroid FC aberrancies that distinguish MDS from pathological controls

To be applicable in FC analysis of a single patient in daily practice, cut-offs for the identification of MDS-associated changes of all potential aberrancies were defined (10th and 90th percentiles of the pathological controls, Online Supplementary Table S2A) and compared between MDS and controls (Table 3). All parameters differed between groups at a P-value of <0.001, and thus could have been considered for the multivariate logistic regression analysis. However, due to large differences between centers, regarding the percentages of NEC and CD71dim cells (likely due to technical variation as shown for normal controls), these parameters were not entered in the multivariate analysis. Besides, irrespective of the finding that data for CD105 significantly discriminated between subgroups (Table 3), this marker was not included. Entering CD105 data would have reduced the power of the (multicenter) analysis, since data on CD105 were only available in a limited number of centers (5/18) and cases. Multivariate logistic regression analysis was based on 119 MDS cases and 153 pathological controls that had available data for all parameters entered in the test. CD36 CV was identified as the best discriminator between MDS and pathological controls in combination with the CV of CD71, the MFI of CD71 and the percentage of CD117+ erythroid cells (Table 4). These four markers were used to define a FC-erythroid dysplasia score in which aberrancies were considered in a weighted manner: four points for increase in CD36 CV; three points for increase in CD71 CV; two points for decreased CD71 MFI; and two points in the case of decreased or increased percentage of CD117+ erythroid cells (reference ranges are summarized in Table 5). A cutoff of ≥5 points resulted in the identification of MDS-associated erythroid aberrancies by FC at a specificity of 90% (95% CI: 84–94%) and a sensitivity of 83% (95% CI: 24–42). Results for the selected markers and the application of the FC-erythroid dysplasia score in the learning cohort are displayed in Table 6 and Figure 3A. In daily practice, a numerical way of counting aberrancies would be more convenient. This involves the definition of a new cut-off; i.e., ≥2 aberrant markers (Figure 3C). Note that, the exception to this numerical score is that the combination of aberrancies in CD71 MFI and percentage of CD117+ alone is not sufficient to conclude dyserythropoiesis by FC (<5 points in the weighted score). The latter was seen in only one pathological control and three MDS cases.

Correlation between erythroid markers and age

The incidence of MDS increases with age; hence, erythroid markers that are significantly correlated with age may be less suitable for discriminating between MDS and controls. Since we observed significant differences between the groups regarding age (Table 1); correlations between FC results for erythroid markers and age were evaluated for normal bone marrow samples. Only CD105 MFI, a variable that was not included in the multivariate analysis, demonstrated a moderate-to-good inverse correlation with age, i.e., CD105 expression decreased with increasing age (Spearman's Rho -0.55, P<0.001, n=47, Online Supplementary Table S4 and Online Supplementary Figure S4).

Validation of FC aberrancies in the erythroid lineage in MDS and pathological controls

The value of the defined antigenic combinations was tested in an independent cohort. Nine centers provided

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Table 3. Aberrancies in FC markers of the erythroid lineage between MDS and controls within the learning cohort.

<table>
<thead>
<tr>
<th></th>
<th>NBM</th>
<th>PC</th>
<th>MDS</th>
<th>P MDS vs. PC</th>
<th>P MDS vs. NBM</th>
<th>P PC vs. NBM</th>
</tr>
</thead>
<tbody>
<tr>
<td>rel. NEC</td>
<td>2.9</td>
<td>10.1</td>
<td>32.1</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>0.013</td>
</tr>
<tr>
<td>pattern CD71 vs. CD235a</td>
<td>3.7</td>
<td>17.3</td>
<td>64.9</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>%CD71dim</td>
<td>0.8</td>
<td>10.0</td>
<td>31.5</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>rel. MFI of CD71</td>
<td>4.0</td>
<td>10.0</td>
<td>27.8</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>0.045</td>
</tr>
<tr>
<td>rel.CV of CD71</td>
<td>4.5</td>
<td>10.3</td>
<td>45.5</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>0.152</td>
</tr>
<tr>
<td>rel. MFI of CD36</td>
<td>0.8</td>
<td>10.3</td>
<td>25.8</td>
<td>0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>rel. CV of CD36</td>
<td>0.0</td>
<td>10.2</td>
<td>30.1</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>0.001</td>
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<tr>
<td>rel. %CD117 progenitors</td>
<td>8.2</td>
<td>19.4</td>
<td>33.8</td>
<td>0.005</td>
<td>&lt;0.001</td>
<td>0.008</td>
</tr>
<tr>
<td>rel. %CD105 progenitors</td>
<td>6.7</td>
<td>28.8</td>
<td>48.4</td>
<td>0.001</td>
<td>&lt;0.001</td>
<td>0.092</td>
</tr>
<tr>
<td>rel. MFI of CD105</td>
<td>29.8</td>
<td>22.5</td>
<td>58.7</td>
<td>&lt;0.001</td>
<td>0.004</td>
<td>0.395</td>
</tr>
</tbody>
</table>

After applying cut-offs as defined in the set of pathological controls, the results were expressed as ‘0’ and ‘1’ for within and beyond reference range(s), respectively (ranges as displayed in Online Supplementary Table S2A). Percentages of subjects with aberrancy are displayed for normal bone marrow (NB); pathological controls (PC) and MDS cases. Results were compared among subgroups using Fisher’s Exact test. P-values are depicted (P). CV: coefficient of variation; dim: diminished; MFI: mean fluorescence intensity; NEC: nucleated erythroid cells; P: P-value; rel.: relative.
data for this validation cohort, the results are depicted in Figure 4 and Online Supplementary Table S1. Similar to results in the learning cohort, the relative CVs of CD36 and CD71 were significantly increased in MDS as compared to controls, whereas CD36 MFI was significantly decreased. Since the distribution of subcategories was similar in the MDS learning and validation cohorts, we compared FC results between the two MDS cohorts. This revealed that the increase in CD71 CV and the decrease in CD71 MFI were significantly less evident in the MDS validation cohort than in the learning cohort (t-test, P<0.001). Results for CD36 CV and the percentage of CD117+ erythroid cells did not differ between both MDS cohorts (P=0.134 and 0.116, respectively).

Reference ranges, as defined in the learning cohort, were applied to evaluate the data from the validation cohort,

Table 4. Results of multivariate logistic regression analysis in learning cohort.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>odds ratio</th>
<th>95% CI</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>rel. CD36 CV</td>
<td>3.7</td>
<td>1.6 – 8.5</td>
<td>0.003</td>
</tr>
<tr>
<td>rel. CD71 CV</td>
<td>3.2</td>
<td>1.6 – 6.4</td>
<td>0.001</td>
</tr>
<tr>
<td>rel. CD71 MFI</td>
<td>2.2</td>
<td>1.1 – 4.5</td>
<td>0.033</td>
</tr>
<tr>
<td>rel. %CD117</td>
<td>1.7</td>
<td>0.92 – 3.2</td>
<td>0.084</td>
</tr>
</tbody>
</table>

Markers entered in the analysis were relative CD36 MFI, CD36 CV, CD71 MFI, and CD71 CV, and the relative percentage of CD117+ erythroid cells (%CD117). 272/335 cases were available for analysis of which 153 pathological controls and 119 MDS cases in the learning cohort (P<0.001). CI: confidence interval; CV: coefficient of variation; MFI: mean fluorescence intensity; P: P-value; rel.: relative.
and then to calculate the weighted FC-erythroid dysplasia score. This resulted in a specificity of 92% (95% CI: 86–97%) and a sensitivity of 24% (95% CI: 15–34%) for identifying MDS-associated erythroid aberrancies by FC (Figure 3B and Online Supplementary Table S5). In most cases, the numerical score could have been applied (cut-off ≥2 aberrancies; Figure 3D) with the same result regarding presence of erythroid dysplasia; only one MDS had decreased CD71 MFI in combination with an altered CD117⁺ percentage.

Table 5. Reference ranges of FC parameters incorporated in the FC-erythroid dysplasia score (learning cohort).

<table>
<thead>
<tr>
<th>Reference ranges</th>
<th># of PC cases*</th>
<th># of NBM cases†</th>
</tr>
</thead>
<tbody>
<tr>
<td>relative CV of CD36 &lt;145%</td>
<td>175</td>
<td>92</td>
</tr>
<tr>
<td>relative CV of CD71 &lt;133%</td>
<td>177</td>
<td>86</td>
</tr>
<tr>
<td>relative MFI of CD71 &gt;46%</td>
<td>250</td>
<td>126</td>
</tr>
<tr>
<td>relative %CD117⁺ erythroid cells 37–212%</td>
<td>182</td>
<td>122</td>
</tr>
</tbody>
</table>

Reference ranges were determined in the learning cohort. Reference ranges represent values relative to median values for the analyzed markers in the erythroid compartment of normal bone marrow (NBM) subjects (learning cohort). These values represent 10th and/or 90th percentiles as determined in the set of pathological controls (PC) within the learning cohort. *number of PC cases that were available to calculate cut-off values (10th and 90th percentiles); †number of NBM cases that were available to calculate median values. CV: coefficient of variation; MFI: mean fluorescence intensity.
Discussion

Analysis of erythroid dysplasia is rarely included in current FC protocols for MDS, since the significance of FC data from the erythroid lineage is, to a large extent, still under debate. Here, we reported the results of a multicenter study within the IMDSFlow group, which focused on defining erythroid parameters that enable discrimination of dyserythropoiesis associated with MDS from non-clonal cytopenia. The majority of erythroid FC markers that are recommended for evaluation of dysplasia according to ELNet guidelines were significantly different between MDS and controls. Analysis of the presence of aberrancies in the erythroid markers CD71 and CD36 (expressed as the CV), together with the MFI of CD71 and an abnormal percentage of CD117+ erythroid progenitor cells, provided the best discrimination between MDS and non-clonal cytopenia. A weighted score based on these four parameters yielded a specificity of 90% (95% CI: 84–94%) in the learning cohort and 92% (95% CI: 86–97%) in the validation cohort. Sensitivity of the weighted score was 33% (95% CI: 24–42%) and 24% (95% CI: 15–34%) in the learning and validation cohorts, respectively. The latter lower sensitivity could be explained by a less evidently increased CD71 CV and decreased CD71 MFI in the MDS validation cohort compared to the learning cohort. Hence, fewer MDS cases scored CD71 CV and/or CD71 MFI as aberrant (Online Supplementary Table S3). Notably, these scores only reflect the presence of FC-erythroid dysplasia, not the likelihood of an MDS diagnosis.

Tenth and 90th percentiles in the validation cohort’s control cases slightly differed from the learning cohort (Online Supplementary Table S2B). Yet, application of these cut-offs in the validation cohort resulted in comparable specificity: 91% (95% CI: 84–96%; sensitivity: 27% (95% CI: 18–37%). In general, cut-off values are most reliable when defined by standardized analyses of control samples in a single center. Notably, no consensus has been reached as to whether percentiles, standard deviations or log differences should be applied as cut-offs for any of the MDS-associated aberrances.

Specificity of the defined markers for identification of MDS-associated erythroid changes is considered to be more important than their general diagnostic value for

Table 6. Results of FC aberrancies in the erythroid lineage and the FC-erythroid dysplasia score among MDS cases and controls within the learning cohort.

<table>
<thead>
<tr>
<th>Categories</th>
<th>Increased CV of CD36</th>
<th>Increased CV of CD71</th>
<th>Decreased relative MFI of CD71</th>
<th>Increased % of CD117+ erythroid progenitors</th>
<th>FC-erythroid dysplasia score ≥5</th>
<th># of cases in flow score*</th>
</tr>
</thead>
<tbody>
<tr>
<td>normal controls</td>
<td>0</td>
<td>5</td>
<td>4</td>
<td>8</td>
<td>3</td>
<td>3/79</td>
</tr>
<tr>
<td>pathological controls</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>19</td>
<td>10</td>
<td>15/153</td>
</tr>
<tr>
<td>MDS</td>
<td>30</td>
<td>46</td>
<td>28</td>
<td>34</td>
<td>33</td>
<td>39/119</td>
</tr>
<tr>
<td>MDS subcategories</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RCUD</td>
<td>25</td>
<td>31</td>
<td>26</td>
<td>41</td>
<td>33</td>
<td>5/15</td>
</tr>
<tr>
<td>RARS(-t)</td>
<td>60</td>
<td>64</td>
<td>33</td>
<td>31</td>
<td>57</td>
<td>8/14</td>
</tr>
<tr>
<td>RCMD</td>
<td>28</td>
<td>44</td>
<td>25</td>
<td>32</td>
<td>30</td>
<td>24/79</td>
</tr>
<tr>
<td>del(5q)</td>
<td>10</td>
<td>30</td>
<td>38</td>
<td>63</td>
<td>13</td>
<td>1/8</td>
</tr>
<tr>
<td>MDS NOS</td>
<td>–</td>
<td>80</td>
<td>40</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Pathological control subcategories</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>iron deficiency anemia</td>
<td>11</td>
<td>0</td>
<td>0</td>
<td>11</td>
<td>6</td>
<td>1/18</td>
</tr>
<tr>
<td>anemia in chronic disease†</td>
<td>7</td>
<td>10</td>
<td>3</td>
<td>23</td>
<td>10</td>
<td>3/29</td>
</tr>
<tr>
<td>vitamin B12/folic acid deficiencies</td>
<td>0</td>
<td>17</td>
<td>0</td>
<td>25</td>
<td>0</td>
<td>0/6</td>
</tr>
<tr>
<td>anemia in auto-immune diseases†</td>
<td>0</td>
<td>0</td>
<td>20</td>
<td>0</td>
<td>0</td>
<td>0/5</td>
</tr>
<tr>
<td>anemia due to renal failure</td>
<td>25</td>
<td>0</td>
<td>0</td>
<td>60</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>anemia other†</td>
<td>0</td>
<td>22</td>
<td>11</td>
<td>0</td>
<td>0</td>
<td>0/5</td>
</tr>
<tr>
<td>cytopenia associated with marrow infiltration</td>
<td>0</td>
<td>20</td>
<td>16</td>
<td>40</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>cytopenia induced by chemotherapy or medication or post-SCT</td>
<td>10</td>
<td>0</td>
<td>24</td>
<td>9</td>
<td>0</td>
<td>0/7</td>
</tr>
<tr>
<td>ITP or neutropenia or auto immune cytopenia NOS</td>
<td>17</td>
<td>9</td>
<td>12</td>
<td>9</td>
<td>14</td>
<td>3/21</td>
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<tr>
<td>reactive conditions</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>cytopenia induced by infections</td>
<td>19</td>
<td>39</td>
<td>24</td>
<td>19</td>
<td>28</td>
<td>5/18</td>
</tr>
<tr>
<td>normal bone marrow</td>
<td>0</td>
<td>20</td>
<td>33</td>
<td>67</td>
<td>20</td>
<td>1/5</td>
</tr>
<tr>
<td>(peripheral cytopenia NOS)</td>
<td>5</td>
<td>0</td>
<td>4</td>
<td>16</td>
<td>0</td>
<td>0/20</td>
</tr>
<tr>
<td>other than defined subcategories NOS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>inconclusive</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>18</td>
<td>0</td>
<td>0/11</td>
</tr>
</tbody>
</table>

Displayed numbers correspond to the percentage of cases per subgroup that were beyond the reference ranges (Table 5). The coefficient of variation (CV) for CD71 and CD36 were tested against the 90th percentile; the expression level (mean fluorescence intensity: MFI) of CD71 was against the 10th percentile; and the percentage of CD117+ erythroid progenitor cells was tested against both the 10th and 90th percentiles. Only data of subsets with five or more cases are depicted. Diagonally marked cells represent data not available or reliable (i.e., missing or only small data sets (<5 cases)). Note: “number of cases with a FC-erythroid dysplasia score of ≥2 per total number of cases in which all parameters were available that comprise the score; the subcategories “anemia in chronic disease” comprises iron incorporation disorders, bowel diseases, diabetes, etc., “anemia in auto-immune diseases” comprises AIHA, AITP, Rheumatoid Arthritis, SLE, etc. and “anemia other” comprises, among others, cases of normocytic anemia, anemia unexplained; NOS: not otherwise specified.
Flow cytometric analysis of erythrodysplasia in MDS

MDS. The specificity may be optimized by increasing the cut-off from ≥5 to ≥6 points (specificity 96% in both cohorts), at the cost, however, of a decrease in sensitivity (24% and 14%, in the learning and validation cohorts, respectively).

To simplify interpretation of results from erythroid analysis, a numerical way of counting aberrancies was tested; a cut-off of ≥2 aberrant markers led to comparable specificity and sensitivity as for the weighted score. However, it must be taken into account that the sole combination of CD71 MFI and percentage of CD117- erythroid progenitors is not sufficient to indicate MDS-associated changes in the erythroid lineage. Cremers et al. compared the analysis of the set of four FC-parameters to erythroid dysplasia as assessed by cytomorphology.29 They demonstrated that FC correlated well with cytomorphology, albeit at a lower sensitivity (low/int-1 risk MDS, 64% vs. 84%, respectively); controls showed 11% and 10% of dysplasia by FC and cytomorphology, respectively. The findings presented herein confirmed results from a recent study27 that reported a significant increase in CD71 CV and CD36 CV to be highly suggestive for MDS. Yet, discrimination between MDS and controls based on CV values was less clear in the current dataset. Mathis et al.28 stated that the difference in CD71 CV between MDS and controls was less pronounced after erythrocyte lysis; however, this was not the case for CD36 CV. Since all data in the present study were obtained after erythrocyte lysis, it might explain the observed differences. It may seem paradoxical to use erythrocyte lysing procedures when the focus is on analysis of the erythroid lineage; but lyse-stain-wash is the recommended protocol for processing samples for FC in MDS.3 Despite IMDSFlow recommendations, methodological variation between centers may have led to differences in results as demonstrated in normal controls. Harmonization, or even standardization, of methods may narrow differences and improve validity of conclusions from multicenter studies, as has been demonstrated with the Euroflow consortium.30 Hence, grouping of data per center, in case of multicenter studies, as has been demonstrated with the Euroflow consortium.30 Hence, grouping of data per center, may improve the evaluation of results. The findings presented in this study were obtained after erythrocyte lysis, it might explain the observed differences. It may seem paradoxical to use erythrocyte lysing procedures when the focus is on analysis of the erythroid lineage; but lyse-stain-wash is the recommended protocol for processing samples for FC in MDS.3 Despite IMDSFlow recommendations, methodological variation between centers may have led to differences in results as demonstrated in normal controls. Hence, grouping of data per technical procedure could have been informative from a practical perspective; yet, the power of the analyses within and between numerous subgroups of centers would have been strongly limited by sample sizes. Notably, in daily practice, FC results in subjects suspected of MDS should preferably be compared with a center’s own cohort of control samples. Despite these technical considerations, our data confirm the robustness of the evaluation of an increase in CD71 and CD36 CV on erythroid cells.28

Another discriminatory marker was the percentage of erythroid progenitors defined as CD117-. A potential marker for future inclusion in erythroid data analysis by FC is CD105. It has been demonstrated (in normal and pathological controls) that CD105 is lost before carboxylic anhydrase is expressed, which suggests that the majority of CD105+ erythroid progenitor cells are not subject to ammonium chloride-based lysing protocols.16 This confirms the robustness of the percentage of erythroid progenitors (CD117+ and/or CD105+) as a marker for erythroid dysplasia. Notably, hemodilution impacts the analysis of the erythroid compartment as it may result in a lack or paucity of erythroid progenitors, similar to what is seen in the myelomonocytic compartments. In heavily hemodiluted samples, the erythroid lineage should be considered non-evaluable.

Application of CD105 may overcome the potential error of assigning CD117+ myeloid progenitors as erythroid progenitors, especially when combined with CD117 and an additional myeloid marker such as CD33. Furthermore, CD105 overexpression was confirmed in some cases of MDS in our dataset.15,20 However, we also observed a decreased expression in MDS. Notably, CD105 expression was negatively correlated with age in normal controls. Future studies in larger data sets may elucidate whether CD105 is truly valuable in analysis of erythroid dysplasia in MDS.

New insights may improve the impact of FC in the diagnosis of MDS. A recent report showed that increased expression of CD44 on all maturation stages of erythroid cells was associated with MDS, irrespective of presence or absence of morphologic dyserythropoiesis.28 In addition, decreased expression of the major coxsackie-adenovirus receptor (CAR) was demonstrated in dysplastic CD105+ erythroid progenitors.32,33 The diagnostic value of the here-in presented parameter combination is limited. Yet, ultimately, the analysis of the erythroid and myelomonocytic lineages and hematogones should be combined. Further validation should reveal the power of the herein defined erythroid markers. Results from a prospective clinical trial in low/int-1 risk MDS demonstrated that the addition of proposed erythroid FC-markers to the more widely acknowledged analysis of the myelomonocytic lineage increased the sensitivity of MDS-FC from 68 to 80% (specificity only slightly decreased from 98% to 95%).29 In addition, it would be relevant to elucidate the value of the combination of myeloid and erythroid FC markers in indeterminate cases according to cytomorphology. MDS-FC of the myelomonocytic lineage has shown to be of negative predictive value in these cases.34,35

In view of emerging knowledge from next generation sequencing analysis, future research may also concern the analysis of the relation between gene modifications/mutations and FC findings. Data comparing cytogenetic aberrations and FC have already demonstrated distinct profiles.36,57 Parallel mutational data in the current cohort are not available, but it would be of interest to compare, for instance, the immunophenotypic profile of nucleated erythroid cells in relation to the presence of a SF3B1 mutation since there is a relation between this mutation and the occurrence of ring sideroblasts.38 Cytomorphology reports dysplastic features in erythropoiesis in non-MDS cases, such as reactive conditions.59,62 Moreover, patients with cytopenia due to marrow infiltration may demonstrate FC aberrancies associated with MDS. MDS may even coincide with the other malignancy in these patients.39,40 A subset of patients with reactive marrow or marrow infiltration in our dataset indeed showed multiple erythroid aberrancies (5/23 and 2/9, respectively; Online Supplementary Table S5). Follow-up analysis after several months may exclude or confirm MDS in these cases.34,35 This stresses that FC analysis in suspected MDS, though proven very specific, should always be part of an integrated diagnostic approach rather than a solitary diagnostic tool.41 In line with this, FC may attribute to the diagnostic work-up in cases that show clonal hematopoiesis of indeterminate potential (CHIP), particularly when patients present with cytopenia and have indeterminate cytomorphology and/or non-informative cytogenetics. To summarize, we identified significant aberrancies with respect to the FC markers recommended by
IMDSFlow for analysis of the erythroid lineage in MDS. The best indicators of dysplastic changes associated with MDS were an increased CV of CD56 and CD71, a decreased MFI of CD71 in combination with decreased or increased percentages of erythroid progenitors (CD117+). Application of the defined marker set demonstrated high specificity. Future studies should assess the contribution of the selected erythroid markers to the evaluation of the myeloid progenitors, the maturing myelomonocytic lineage, and hemogones in current FC protocols in MDS. This will be implemented in an upcoming multicenter data collection exercise within IMDSFlow.

Acknowledgments

The authors would like to thank Claudia Cabi and Kelly Scheuten (VU University Medical Center Amsterdam, The Netherlands), Jeroen Lauf (Elisabethinen Hospital, Linz, Austria), Fratke Bellos (MIL Munich Leukemia Laboratory, Munich, Germany), Hans Veersna (Radboud University Medical Center, Nijmegen, The Netherlands), Rik Broinmans and Andre Bijkerk (Erasmus Medical Center, Rotterdam, The Netherlands) for their contribution to data collection, although they did not take part in the IMDSFlow working group conferences. Contributors to The Netherlands are all members of the “Flow Cytometry in MDS” working group within the Dutch Society of Cytometry.

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