Enumeration of CD34+ blasts by immunohistochemistry in bone marrow biopsies from MDS patients may have significant impact on final WHO classification

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Abstract

The percentage of blasts cells in the bone marrow (BM) of MDS patients is one of the key parameters for MDS classification and for the differential diagnosis with acute myeloid leukemia (AML). Currently, the gold standard to determine the blast percentage is conventional cytomorphology. To assess the possible impact of blast cell enumeration in BM biopsies from MDS patients on the final WHO classification using CD34 immunohistochemistry (IHC) a total of 156 BM samples from MDS and MDS-AML patients were studied and compared to blast counts by cytomorphology (CM). Eighty-nine BM aspirates were also studied by flow cytometry (FCM). Percentages of CD34+ blasts by IHC were determined blindly by two hematopathologists. Automated CD34-cell count was performed in 25 cases. Good overall agreement was found for CM and FCM with respect to critical blast thresholds (5%, 10%, 20%) (p < 0.05). However, in 17% of patients, CD34+ blast counts by IHC were higher as compared to CM with possible impact on MDS subclassification. In 7 of 21 AML patients, diagnosis was established on BM histology, while the blast percentage by CM was below the AML threshold. The assessment of CD34+ cells by IHC showed high interobserver agreement (Spearman R 0.95, p < 0.01), while automated CD34 counts were not optimal due to interference with other cellular and stromal elements. BM histology including CD34 IHC improves the diagnostic accuracy in MDS and AML. The quantification of blast cells should be based on the integration of all three methods for reliable disease classification and risk assessment.

Keywords Bone marrow biopsy · CD34 immunohistochemistry · Cytomorphology · Blast percentage · Flow cytometry

Introduction

The cytomorphological enumeration of blast percentages in bone marrow (BM) smears is a critical parameter for classification of myelodysplastic syndromes (MDSs) [1]. The blast percentage is also an important prognostic marker as reflected by the high intrinsic prognostic power of the WHO classification [2]. Blast percentage cut-offs at 5%, 10%, 20%, and 30% are included as prognostic factors in the International Prognostic Scoring System (IPSS) [3]. The further splitting of the lower blast range into two separate groups (0–2% vs >2–<5%) in the revised IPSS (R-IPSS) provided groups with very low risk versus low-risk features [4]. Indeed, patients with ≤2% BM blasts had lower risk of disease progression into acute myeloid leukemia (AML) and a more favorable outcome [4].

The International Working Group on Morphology of MDS proposed a consensus for the definition and enumeration of BM blasts, widely implemented in clinical practice [5]. Although the reliability of blast cell count with respect to the 5% threshold has been demonstrated [6–8], the reproducibility of counts within the lower blast range has been questioned. Font et al. found low agreement among cytologists when analyzing MDS patients with ≤2% BM blast counts [6].
Therefore, even when good-quality BM smears are available, the reproducible identification and quantification of blast cells might be a challenge. Moreover, blast counts in smears may not be representative due to hemodilution or low cellularity of smears that are often seen in hypoplastic MDS or in MDS with significant (≥ grade 2 WHO) fibrosis [9–12]. In addition, CD34+ cell clusters as identified by immunohistochemistry (IHC) that were reported as independent prognostic marker for progression to AML may be seen even in MDS without blast excess by cytomorphology (CM) [13, 14]. In follow-up for progression to AML may be seen even in MDS without blast excess by cytomorphology (CM) [13, 14]. In follow-up for progression to AML may be seen even in MDS without blast excess by cytomorphology (CM) [13, 14].

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In a significant number of cases [20–24]. However, it has been shown that a 2% cut-off for CD34+ cells by FCM reached higher diagnostic significance than 2% blasts by CM [25]. A recent multicenter quality assurance study stressed the need for a standardized approach for the enumeration of CD34+ myeloid blast cell count by FCM [26]. A series of publications from the International MDS Flow Cytometry Working and the ELN group provided guidelines for FCM application in the diagnostic work-up of MDS [23, 27–31]. However, as it has been stated in the revised WHO 2017, FCM results cannot replace the morphological differential count in the MDS subclassification [1, 32].

A number of previous studies compared the percentages of blasts by CM to the assessment of the blast compartment by FCM. In this study, we assessed the percentage of CD34+ blasts by IHC in BM biopsies from MDS patients and compared the results to the blast counts obtained by conventional CM and, in a subset of patients, also to the enumeration of CD34+ myeloid blast cells by FCM. By using three different methods in a simultaneous setting, we demonstrate that BM histology with CD34 IHC staining (mouse MoAb CD34 QBend10, RTU) was performed on FFPE BM sections using a Ventana BenchMark automated slide stainer. The fraction of CD34-positive blast cells was assessed in randomly selected fields based on a count of ≥500 hematopoietic cells using ×60 lens and a ×10 eyepiece. A CD34+ cluster was defined as a group of ≥3 positive cells as previously described [13].

Bone marrow histology and cytology

CD34 IHC staining (mouse MoAb CD34 QBend10, RTU) was performed on FFPE BM sections using a Ventana BenchMark automated slide stainer. The fraction of CD34-positive blast cells was assessed in randomly selected fields based on a count of ≥500 hematopoietic cells using ×60 lens and a ×10 eyepiece. A CD34+ cluster was defined as a group of ≥3 positive cells as previously described [13].

Multiparameter flow cytometry

The BM aspirates were processed within 2 h from sampling using a stain and lyse/wash technique [34]. Data acquisition was performed using FACS Calibur (BD) flow cytometer (4-color panel) and FACS Canto II (BD) (8-color panel) for samples collected between 2004 and 2007 and 2008 and 2016, respectively. The applied panels are shown in Supplementary Tables 3–4. The percentage of myeloid CD34+ cells was compared to clots from aspirates, strongly supporting the use of the former at initial investigation for correct disease classification.

Methods

Study design

This retrospective study was performed using archived BM samples (BM trephines/clots from aspirates together with ≥2 BM smears and ≥1 imprint from BM trephines) from patients with MDS (n = 135) and AML with myelodysplasia-related changes (n = 21), obtained at initial diagnosis (between 2004 and 2016). Material for FCM was available in 89 of 135 (66%) MDS patients. BM trephines from AML patients were referred from outside hospitals without material for FCM. All samples were reviewed and reclassified by two experienced hematopathologists (LS, AP) according to the WHO 2017 classification [16]. Baseline characteristics of the study cohort are shown in Table 1 (detailed data is provided in Supplemental Tables 1–2). This study was performed in line with the principles of the Helsinki Declaration. Approval was granted by the Swedish Ethical Review Authority (Dnr 2009–00394).

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calculated from all nucleated BM cells after excluding debris and doublets. In the 4-color panel, CD34+ B cell precursors were excluded as CD34+ cells with very low SSC [24]. In the 8-color panel, CD34+ myeloid blast cells were identified by the CD45dim CD34+ CD117+ CD33+13dim HLA-DR+ phenotype (tube 1) and CD45dim CD34+ CD117+ CD33+13dim HLA-DR+ phenotype (tube 2). Following ELN guidelines [30], aberrant myeloid blast populations were identified by loss or overexpression of myeloid-associated antigens, aberrant expression of lymphoid markers, and monocytic differentiation by the expression of CD11b, CD64, and CD36.

Statistical analyses

The Spearman rank correlation coefficient and Pearson test were used to determine the correlation between blast cell percentages as assessed by the three methods. The chi-square and Fisher’s exact tests were used to assess the correlation between concordant and discordant cases with respect to the IPSS cytogenetic subgroup. The non-parametric Mann Whitney test was used to assess the effect of marrow fibrosis on blast percentage for the comparison between CM and IHC. All p values are two-tailed and considered statistically significant when < 0.05. The statistical analyses were performed using Statview (Abacus Concepts Inc., Berkeley, CA, USA).

Results

Baseline clinical and morphological characteristics

Of 135 MDS cases, most were either MDS-MLD (47%) or MDS-EB (39%), while other subtypes were less frequent (Table 1). In 129 patients with available data, the distribution of IPSS-R cytogenetic risk groups was as follows: very good/good (n = 64), intermediate (n = 13), and poor/very poor (n = 52). Among AML with myelodysplasia-related changes (AML-MRC), 17 of 19 patients with available data had poor or very poor cytogenetics including del(5q) with complex karyotypes in 15 patients. BM showed median cellularity of 64% and 68% (range 10–100%) in MDS and AML patients, respectively; 14 of 135 (10%) MDS and 3 of 21 (14%) AML patients had ≤30% BM cellularity. Moderate or severe (≥ grade 2) marrow fibrosis was seen in 15% of samples in both groups. The majority of the specimen from MDS patients

| Table 1 | Baseline characteristics of the study cohort |
|---------|---------------------------------------------|
|         | MDS (n = 135) No. (%) | AML with myelodysplasia related changes (n = 21) No. (%) |
| Median age, years (range) | 71 (2–89) | 73 (52–83) |
| Sex, male/female | 77/58 | 11/10 |
| WHO 2017 |        |        |
| • MDS del5q | 6 (4%) |        |
| • MDS-RS-MLD | 7 (5%) |        |
| • MDS-SLD | 3 (2%) |        |
| • MDS-MLD | 64 (47%) |        |
| • MDS-EB | 52 (39%) |        |
| o MDS-EB-1 | 33 (24%) |        |
| o MDS-EB2 | 19 (14%) |        |
| • Refractory cytopenia of childhood (RCC) | 3 (2%) |        |
| BM biopsy, specimen type |        |        |
| • BM trephine | 96 (71%) | 21 |
| • Aspiration biopsy | 39 (28%) | 0 |
| Cellularity, median (range) | 64% (20–100%) | 68% (range 10–100%) |
| Marrow fibrosis |        |        |
| • Grade 0–1 | 115 (85%) | 18 (97%) |
| • Grade 2–3 | 20 (15%) | 3 (15%) |
| CD34+ clusters |        |        |
| • Present | 30 (22%) | NA |
| IPSS-R cytogenetic risk group |        |        |
| • Very good/good | 64 (47%) | 1 (5%) |
| • Intermediate | 13 (10%) | 1 (5%) |
| • Poor/very poor | 52 (39%) | 17 (80%) |
| • Missing data | 6 (4%) | 2 (10%) |

MDS-RS-MLD MDS with ring sideroblasts and multilineage dysplasia, MDS-SLD MDS with single lineage dysplasia, MDS-MLD MDS with multilineage dysplasia, MDS-EB MDS with excess blasts, NA not applicable
were BM trephines (96 of 135 cases). All AML patients had adequate BM trephines, while BM aspirate for FCM was not available for this group.

Characteristics of CD34 immunostaining and interobserver reproducibility

BM trephines \((n = 117)\) and clots from aspirates \((n = 39)\) were retrospectively reviewed by two experienced hematopathologists (LS and AP) in a blinded fashion for the percentage of CD34+ cells. The individual results of CD34 IHC in MDS and AML-MRC patients are provided in Suppl. Tables 1-2. CD34-IHC showed \(\geq 1\%\) cells consistent with blasts in 89 of 135 (66%) MDS; all AML-MRC cases were CD34-positive. Uneven distribution of CD34-positive blast cells and CD34-positive clusters were frequently seen in MDS-EB subtypes (25 of 53 cases, 47%) and occasionally in MDS-MLD (5 of 63 cases, 8%) (Fig. 1). Of note, four of MDS-MLD cases (<5% blasts by CM) had also an overall increase of CD34-blasts cells by IHC (>5%). In addition, strong aberrant CD34 expression in megakaryocytes was frequently observed, particularly in MDS-EB subtypes as illustrated in Fig. 2.

The blinded assessment of CD34+ blast cells by both reviewers in a subset \((n = 26)\) of patients (7 MDS-MLD, 14 MDS-EB-1/2, 5 AML) showed high interobserver correlation (Suppl. Figure 1, Spearman R 0.95, \(p < 0.01\)). Notably, both reviewers found higher percentage of CD34+ blast cells by IHC as compared to CM with respect to WHO blast thresholds in 7 of 21 (33%) MDS cases, many of which showed uneven distribution of CD34-positive blast cells in the BM biopsies. The five AML cases were diagnosed on the basis of BM histology and CD34 IHC while blast percentages by CM were below the AML threshold.

Comparison of the blast count by cytomorphology to the percentage of CD34-positive blast cells by immunohistochemistry and flow cytometry

Based on the blast count performed on BM smears and imprints in comparison to the percentage of CD34+ cells by either IHC or FCM, the cases were stratified into three groups according to WHO blast thresholds. Of 132 cases, 84 (64%) cases had <5%, 30 (23%) between ≥5 and <10%, and 18 (13%) between ≥10 and 19% BM blasts by CM. BM smears/imprints were inadequate (hemodiluted) in three cases, all had an increase of CD34+ blasts by IHC (Suppl Table 1, cases no. 103, 107, and 133) and two had grade 2 marrow fibrosis. Among 14 MDS cases with lower cellularity (<30%) and <5% blasts by CM, IHC detected an increase of CD34+ blast cells in 2 cases (cases no. 85, 91). Seven other cases with <20% blasts by CM were classified as AML-MRC on the basis of histopathological findings.

Comparison of CM blast count and the percentage of CD34-positive blast cells in BM histology

When comparing the two techniques, 95 of 132 (72%) cases were within the same blast range (Table 2). However, the results were not significantly correlated (Suppl. Figure 2, Pearson’s \(r 0.728, p = 0.113\) for groups). BM histology revealed a higher percentage of CD34+ blasts in 23 of 132 (17%) samples (with CD34+ clusters in four patients) as compared to CM, which would allocate the patients to a higher WHO MDS subgroup and a higher IPSS-R prognostic blast score value. Only two of these samples had ≥2 grade 2 marrow fibrosis. Based on the IHC results, 12 patients with <5% BM blasts by CM (all MDS-MLD) would be moved to the MDS-EB category, and 11 patients with 5–9% blasts by CM (MDS-EB1) would be reclassified as MDS-EB2. Higher percentages of CD34+ blast cells in BM histology as compared to the CM blast counts were more often detected in BM trephines as compared to aspirate clots (\(P < 0.05\)), reflecting that uneven distribution of blast cells may be more easily detected in BM trephines than in clots. In this series, the blast frequency and classification were not affected by the presence or absence of marrow fibrosis. In contrast, higher blast percentages by CM as compared to the percentage of CD34 by IHC and/or FCM with respect to classification thresholds were seen in 14 of 132 MDS (10%), which may indicate that not all blasts were CD34 positive. Among those, the presence of aberrant blast populations truly negative for both CD34 and CD117 was confirmed in two of nine cases with available material for FCM. Among samples with <1% CD34-positive blast cells by IHC \((n = 46)\), only four cases had an increase of blasts by CM (between 7 and 11%). Of those four cases, two had ≤1% and one 1% CD34+ myeloid blasts by FCM; the fourth was not diagnostic due to hemodilution.

Comparison of CM blast count and the percentage of CD34+ myeloid blasts by FCM

The blast count in BM smears/imprints and the percentage of CD34+ myeloid blasts by FCM were within the same range in 78 of 88 cases (88%) with respect to the critical WHO blast thresholds. The results of the measurements are summarized in Table 3 and illustrated in Suppl. Figure 3 (Pearson’s \(r 0.782, p < 0.01\) for both groups and absolute values). The percentage of blasts by CM vs FCM was higher in 10 (11%) cases, ranging between 6 and 12% of total BM cells. Direct comparison to FCM results showed <5% CD34+ myeloid blasts in all but one of these cases (with similar results for IHC), of which four had ≤1% CD34+ myeloid blasts, indicating that blast cells were negative for this marker. A higher percentage of CD34+ myeloid blasts by FCM vs CM was seen in one case (9% vs 3%, respectively); the histological material consisted
of a BM clot with 30% cellularity and < 1% CD34+ IHC cells without evidence of fibrosis.

Comparison of the percentage of CD34 blasts by immunohistochemistry and FCM

The two methods showed results within the same blast range in 75 of 89 (84%) cases: 68 of 89 (76%) samples had < 5%, 2 cases between ≥ 5 and < 10%, and 5 cases between ≥ 10 and 19% by both methods (Table 4 and Suppl. Figure 4, Pearson’s r 0.767, p 0.012 for blast groups). Differences in the percentages of CD34-positive blasts (n = 14) were mainly due to higher values in the BM trephines, likely explained by uneven distribution of blast cells and the presence of CD34-positive clusters. In three cases, the CD34+ blast percentage was higher by FCM with similar results for blast percentage in BM smears. Notably, the corresponding biopsy material was BM aspirate clots with high cellularity in two cases (80% and 90%) and low cellularity (30%) in one case. The findings may indicate that BM biopsies, in particular BM trephines, detect higher blast percentages in a fraction of cases that may be more difficult to detect in BM aspirate clots. The percentage of normal early B-cell precursors (by FCM) was < 0.1% in all MDS with ≥ 5% CD34+ blasts by IHC and available material for FCM; an example for this is shown in Suppl. Figure 5.

Higher blast percentages in bone marrow histology in comparison to cytomorphology is associated with shorter overall survival and poor-risk cytogenetics

Data on follow-up time, time to progression and overall survival (OS) were collected for all MDS patients. To study the impact of higher blast counts in BM biopsies on OS, MDS-MLD and MDS-EB patients were assessed separately (Suppl. Figure 6a-b). Patients with MDS-MLD and ≥ 5% CD34+
blasts by IHC (n = 12) had shorter OS as compared to MDS-MLD with <5% by both methods (n = 59); OS at 2 years was 10% and 55% for the two groups, respectively (p < 0.0001, log-rank). However, 9 of 12 (75%) had poor/very poor IPSS cytogenetics as compared to 12 of 59 (20%) MDS-MLD.

**Table 2** Comparison of blast percentage by CM and CD34 IHC in BM biopsies

| Percentage of CD34+ cells by IHC | <5% | ≥5–<10% | ≥10–<20% | Total |
|----------------------------------|-----|---------|----------|------|
| BM blasts %                      |     |         |          |      |
| <5%                              | 72 (55%) | 9 (7%) | 3 (2%) | 84 (64%) |
| ≥5–<10%                          | 9 (7%) | 10 (8%) | 11 (8%) | 30 (23%) |
| ≥10–<20%                         | 3 (2%) | 2 (2%) | 13 (10%) | 18 (14%) |
| Total                            | 84 (64%) | 21 (16%) | 27 (21%) | 132 (100%) |

**Table 3** Comparison of blast percentage by CM and the percentage of CD34+ myeloid blasts by FCM

| Percentage of CD34+ cells by FCM | <5% | ≥5–<10% | ≥10–<20% | Total |
|----------------------------------|-----|---------|----------|------|
| BM blasts %                      |     |         |          |      |
| <5%                              | 64 (73%) | 1 (1%) | 0 (0%) | 65 (74%) |
| ≥5–<10%                          | 7 (8%) | 8 (9%) | 0 (0%) | 15 (17%) |
| ≥10–<20%                         | 1 (1%) | 1 (1%) | 6 (7%) | 8 (9%) |
| Total                            | 72 (82%) | 10 (11%) | 6 (7%) | 88* (100%) |

*89 cases had a bone marrow aspirate for FCM assessment; BM smears were not diagnostic in one case due to hemodilution.
patients \((p = 0.001, \text{ chi-square test})\). MDS progression was documented in 1 of 12 and 10 of 59 MDS-MLD patients; mean last morphological follow-up time was 41 months. For MDS-EB1 patients \((n = 33)\) with blast percentages in the same range \((\geq 5\% - 9\%)\) for both methods \((n = 22)\) or higher by CD34 IHC \((n = 11)\), OS at 2 years was 30\% and 18\%, respectively \((p = 0.78, \text{ log-rank})\); 14 of 22 (63\%) and 6 of 11 (54\%) had poor/very poor IPSS-R cytogenetics \((p = 0.975, \text{ chi-square test})\). Disease progression and/or AML transformation was documented in 10 of 52 (19\%) MDS-EB1/2 patients, and two of these had higher \((\geq 10\%)\) blast counts by BM histology vs CM; mean last morphological follow-up time was 17 months.

Cytogenetic data with respect to the whole cohort was available in 129 MDS patients (Suppl. Table 1); three of these cases had inadequate cytomorphological material. For the purpose of statistical analysis, patients were grouped into very good/good \((n = 63)\), intermediate \((n = 13)\), and poor/very poor \((n = 50)\) cytogenetic risk groups according to the revised International Prognostic Scoring System (R-IPSS). Differences in blast percentages affecting critical classification thresholds were highly associated with poor-risk cytogenetic abnormalities; 15 of 22 patients with higher blast counts in BM histology vs CM had poor or very poor cytogenetics (for example case 110, shown in Fig. 1), while 7 patients had good or intermediate cytogenetics \((p = 0.026)\).

### Automated quantification of CD34+ blast cells in bone marrow biopsies

BM slides were scanned using a Pannoramic™ P250 Flash digital slide scanner (3DHISTECH Ltd., Budapest, Hungary) equipped with a CIS color camera (CIS Corporation, Tokyo, Japan) for bright field image acquisition. Automated quantification of CD34+ cells was performed in 25 cases using the Case Viewer 2.3.0 software with QuantCenter 2.2.1 plug-in (3DHISTECH Ltd. Budapest, Hungary) using CellQuant (CQ) application for image analysis. The measurement was performed in predefined, randomly selected annotation areas rather than whole-image analysis, hereby excluding extramedullary tissue, trabecular bone, “empty spaces,” and subcortical areas (Fig. 2). Endothelial cells served as internal control for CD34-positivity. In spite of individual measurement settings for defining characteristics of CD34+ blast cells (including cell size, nuclear shape, staining intensity) in each of the selected BM samples, non-blast cells (e.g., hyperplastic, non-flattened endothelial cells or micromegakaryocytes with strong aberrant CD34 expression) and unspecific structures (e.g. non-cellular background staining) were frequently counted as “CD34-positive.” This error could not be completely eliminated, but calibration of the CQ with the adjustable parameters (color deconvolution for the nuclear counterstain, cell/nucleus size, and size of the cytoplasm) helped to reduce counting of non-blast cells. After these calibrations, the results of the automated count were in some cases comparable to the manual count (Suppl. Table 6). Taken altogether, automated quantification of CD34+ blast cells in BM biopsies is currently still problematic for routine clinical use due to technical pitfalls but also due to the cellular heterogeneity and unspecific or aberrant immunoreactivity in various BM compartments.

### Discussion

Enumeration of blast cells in the BM of MDS patients is a key parameter for correct assignment to MDS subgroups, as well as for the differential diagnosis between MDS and AML. In addition, it is used as a single independent prognostic marker in currently available prognostic risk scores \([4, 35]\). In the updated WHO guidelines \([1]\), despite inaccuracies inherent in manual differential counting, conventional CM is still considered the gold standard to determine blast counts.

Previous studies have shown a generally good correlation between CM and FCM for BM blast enumeration \([17, 19]\), but it has been shown that FCM may underestimate or overestimate blast cell counts in individual cases. The percentage of CD34+ myeloid cells by FCM has been tested as a substitute for a CM count; however, blasts in MDS may not express CD34 \([36, 37]\). Other studies have shown that the CD34^high and/or CD117^HLA-DR^ phenotype of total events showed the highest degree of correlation and agreement with the
morphological assessment of blast counts [19]. However, BM samples taken in a simultaneous setting for cytomorphology and FCM analysis, respectively, may differ in terms of cellularity and blood contamination. FCM has not been accepted as a complementary tool for BM blast count in routine clinical practice but provides valuable information on blast immunophenotype and maturation patterns of various BM cell lineages. In our series, we observed good overall concordance between CM and FCM with respect to critical blast thresholds (88% of cases). In four of 53 MDS with excess blasts (≥ 5% by CM), the percentage of CD34+ myeloid blasts by FCM was ≤ 1% indicating that blast cells lacked this marker.

According to recommendations from the ELN and the revised WHO 2017 diagnostic guidelines, BM trephines should be performed in all patients with suspected MDS for whom BM examination is indicated [15, 16]. Immunohistochemical analysis with CD34 is especially useful for assessing blast percentage in MDS with fibrosis or a hypocellular bone marrow, in which blast percentages are often underestimated. Moreover, it allows the enumeration of CD34+ blast cells and identification of CD34+ cell clusters, which were found to have an independent impact on both overall survival and leukemia-free survival [13]. Previous studies in MDS and AML have shown that CD34 IHC increases diagnostic accuracy [38]. Dunphy et al. found that IHC detected higher CD34+ blast counts as compared to the blast count obtained in smears, which resulted in a change of the initial classification [38]. It has also been illustrated that during follow-up of treated AML patients with CD34+ myeloid blast cells at initial diagnosis, IHC detected higher blast counts (> 5%) as compared to CM (< 5%) in a number of cases [39]. Other studies have proven high overall concordance for positive and negative results for CD34 by IHC in comparison to either CM or FCM [40, 41]. Correlation with FCM may particularly be helpful in MDS cases with micromegakaryocytic hyperplasia and strong aberrant CD34 expression. We demonstrate here a high interobserver correlation for the blinded morphological assessment of CD34+ blasts cells in BM biopsies and good overall concordance between IHC and FCM with respect to critical blast thresholds. However, BM histology detected higher blast percentages by IHC as compared to CM in 17% of MDS cases, which would allocate patients to a higher subcategory and IPSS risk group. Discrepant cases were seen in both MDS-MLD and MDS-EB subtypes and associated with shorter overall survival and poor-risk cytogenetics. In seven of 21 AML patients, final diagnosis was established on the basis of histology and IHC, while cytomorphology of the smears did not fulfill the criteria for leukemia diagnosis. Importantly, discordant cases were more frequently detected in BM trephines as compared to clots from aspirates. Differences in blast percentages between CM and BM histology may partly be explained by uneven distribution of CD34+ blast cells and the presence of CD34+ clusters, which are more easily detected in a trephine biopsy and more frequently seen in high-risk MDS.

The automated assessment of CD34 in BM biopsies was found problematic due to technical issues and positive staining in different cellular and stromal compartments. In addition, the distribution and frequency of CD34-positive blast cells can vary within samples which may require an individual approach for the automated count including the selection of adequate annotation areas and the definition of negative and positive scores (e.g., size of blast cells).

In summary, our findings illustrate that CM does not establish correct blast counts in 100% of cases and should be used with the awareness that BM histology may reveal higher blast percentages, particularly in MDS with high-risk features. Accordingly, as the most reasonable and reliable approach to diagnosing and classifying MDS, we propose the combination of all three methods as gold standard for the assessment of marrow blasts. This should be applied in a simultaneous setting and as part of an integrative diagnostic approach for correct assignment of patients to specific risk categories and MDS subgroups. In cases with increased blast percentages in BM histology, final MDS classification should be based on the higher blast count even if the percentage of blasts by CM is below the 5% threshold. The CM blast count cannot be replaced by FCM, but if FCM finds significant higher counts in the BM aspirate, control sampling may be considered. To our knowledge, there are no larger published studies who have analyzed the percentage of blast cells in BM samples from MDS patients by all three methods (CM, FCM, and BM histology together with IHC) in a simultaneous setting. Additional studies to define standards for the enumeration of CD34 cells in BM samples are currently ongoing by members of the European Bone Marrow Working Group [42]. The findings of our study will be followed up in a larger, population-based cohort of MDS patients at initial diagnosis and under treatment with correlation to clinical data and survival.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.
**Ethical approval** This study was performed in line with the principles of the Helsinki Declaration. Approval was granted by the Swedish Ethical Review Authority (Dnr 2009–00394).

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