abstract

Myelodysplastic syndrome (MDS) represent very heterogeneous group of clonal stem cell bone marrow disorders with ineffective haematopoiesis leading to cytopenias in peripheral blood and increased risk of blastic transformation and evolution of acute myeloid leukemia. MDS is a disease of older age mostly, in children it seems to be very rare. There are several significant morphological, cytogenetic and prognostic differences of the disease in adults and in children. Adult MDS patients most commonly manifest with symptoms of anaemia, bleeding and infection are uncommon. In childhood, MDS manifests predominantly by neutropenia and thrombocytopenia. In addition, some pediatric MDS patients present also with constitutional disease’s signs and symptoms. Early and correct diagnosis in both age groups is essential for the choice of appropriate therapy and also for next life of patients. However, the diagnosis of MDS is challenging, complex and requiring close correlation of clinical symptoms, laboratory parameters and standardized examination of BM biopsies. The authors present an overview focused on biology of MDS in adults and children, on the differences in the incidence, clinical presentation and treatment. They summarize the possibilities and limits of histopathological diagnosis and differential diagnosis of the disease in different age groups. A major problem in the morphological diagnosis of MDS remains the determination, whether the myelodysplasia is due to clonal disorder. It might result also from some other factors, as significant dysplasia can also occur in reactive conditions, and vice versa, only discrete dysplasia is sometimes observed in MDS patients. Although histomorphological and immunohistochemical analysis of BM biopsy is invasive and time-consuming examination, it has its value in the diagnosis, differential diagnosis and evaluation of therapeutic effect.

Key words: Myelodysplastic syndrome, types of myelodysplastic syndrome, risk stratification, adults, children

Introduction

Myelodysplastic syndromes (MDS) represent group of clonal stem cell bone marrow (BM) disorders with ineffective haematopoiesis leading to cytopenias in peripheral blood (PB) and increased risk of blastic transformation and evolution of acute myeloid leukemia (AML) (1,2). They are very heterogeneous myeloid malignancies, mainly of older age, with slightly predominant incidence in men (excluding type with isolated deletion of 5q) (3). At diagnosis, median age is 65-70 years, the incidence increases with age (less than 1/100 000 before 50 years, more than 20/100 000 after 70) (4,3). MDS is rare haematooncological disease in children, typically occurring in age of 6-8 years. According to WHO classification (2008), pediatric MDS accounts for less than 5% of all haematological malignancies in childhood. The estimated incidence is reported 0.5 – 4 / 1 million yearly (5), although this statement may be distorted due to difficulties with diagnosis of pediatric MDS (6). There are several significant morphological, cytogenetic and prognostic differences of disease in adults and in children. (6, 7). Tab 1

Classifications:
The current WHO classification (2008) distinguishes these MDS subtypes of adults: Refractory cytopenia with unilin ear dysplasia (RCUD) manifests by refractory cytopenia...
Table 1. Comparison of the most common features of adult and pediatric MDS (according to Glaubach T., Robinson L. J., Corey S. J., 2014 and Chatterjee T. Choudhry V. P., 2013)

<table>
<thead>
<tr>
<th>Incidence</th>
<th>Adult MDS</th>
<th>Pediatric MDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median age (years)</td>
<td>3-5 /10^5, &gt;20/10^5 u &gt; 70r</td>
<td>1.8 – 4 /10^6</td>
</tr>
<tr>
<td>Presentation</td>
<td>isolated anemia with / without neutropenia and / or trombocytopenia</td>
<td>↑ bicytopenia – RT &gt; RN and / or anemia</td>
</tr>
<tr>
<td>Etiology</td>
<td>↑ primary (de novo)</td>
<td>↑ secondary and t-MDS</td>
</tr>
<tr>
<td>Morphological subtype</td>
<td>↑ RARS a 5q- syndrome, without RAEB-T, subclassification in RAEB1 and 2 with prognostic significance</td>
<td>↑ RCC, RARS and 5q- syndrome are rare, RAEB-T is used, without data about significance of subclassification in RAEB1 and 2</td>
</tr>
<tr>
<td>BM cellularity</td>
<td>↑ or normal, rare</td>
<td>variable, hypocellular</td>
</tr>
<tr>
<td>Cytogenetics</td>
<td>5/5q- &gt; 7/7q-, 3q a 20q anomalies</td>
<td>-7/7q &gt; + 8</td>
</tr>
<tr>
<td>Genetics</td>
<td>mutations in DNMT3A, ASXL1, TET2, SF3B1, U2AF5, methylating changes</td>
<td>mutations in FANC, SBDS, DKC, TERT, TERC, ELANE, HAX1, WAS, GATA2</td>
</tr>
<tr>
<td>Physical findings</td>
<td>nono</td>
<td>skeletal, cutaneous, genitourinary, cardiovascular and gastrointestinal anomalies (related to IBMFS)</td>
</tr>
<tr>
<td>Therapeutic options</td>
<td>transfusions, LND, HA, HSCT</td>
<td>monitoring, transfusions, HSCT</td>
</tr>
</tbody>
</table>

Legend:
- RT – refractory thrombocytopenia
- RARS – refractory anemia with ring sideroblasts
- RAEB / RAEB-T – refractory anemia with excess blasts / in transformation
- -7 – monosomy 7
- +8 – trisomy 8
- LND – lenalidomide
- HSCT – haemopoietic stem cell transplantation
- IBMFS – inherited bone marrow failure syndromes
- RN – refractory neutropenia
- RCC – refractory cytopenia of childhood
- IST – immunosuppressive therapy
- ? - unknown efficacy

and dysplasia limited to one cellular lineage and includes the most common refractory anemia (RA) and less frequent refractory neutropenia (RN) and thrombocytopenia (RT). It accounts for around 10-20% of diagnosed MDS cases (5). Refractory anemia with ring sideroblasts (RARS) is characterised by unexplained anemia, dysplastic changes affecting erythroid lineage and presence of more than 15% of ring sideroblasts. It represents about 3-11% of all cases (5).

Refractory cytopenia with multilineal dysplasia (RCMD) represents MDS with one or more cytopenia and dysplasias in two or more myeloid lineages. This subgroup accounts for around 30% of all MDS (5).

Refractory anemia with excess blasts (RAEB) is characterised by presence of 5-19% of myeloblasts in BM and 2-19% of myeloblasts in PB. According to the blasts count, two clinically different subtypes of RAEB has to be recognized due to different patient’s survival and
incidence of leukemic evolution: RAEB1 (5-9 % blasts in BM and 2-4 % blasts in PB) and RAEB2 (10-19% blasts in BM and 5-19% in PB) (5, 8).

MDS with isolated deletion of 5q (s.c. 5q- syndrome) is characterised by macrocytic anaemia with or without other cytopenia and / or thrombocytosis with the sole cytogenetic abnormality – deletion of 5q, it typically affects elderly women (5).

MDS unclassifiable represents those cases which can not be classified to any of the preceding groups. These patients should be monitored regularly in order to recognize the evolution of the disease in a more specific type (5).

According to the WHO classification (2008), pediatric MDS are classified as follows:

Refractory cytopenia of childhood (RCC) is the most common type with variable clinical course (9), accounting for approximately half of all MDS cases (6). It is characterised by persistent cytopenia with less than 5% BM blasts, less than 2% of PB blasts  and dysplastic changes in two or three lineages or exceeding more than 10% of cells in one lineage (9). In contrast to adult MDS, the WHO classification (2008) does not specify the importance of the number of lineages involved in the dysplasia (which has prognostic significance in adults with MDS). It is recommended, that children who meet the criteria for RCMD, should be considered as RCC, until the prognostic significance of a multilineage presentation is further clarified in children (10).

Advanced pediatric MDS - RAEB (PB blasts 2–19% and/or BM blasts 5–19%), RAEB in transformation (RAEB-T, PB and/or BM blasts 20–29%).

Pediatric RAEB exhibits similar morphological and immunohistochemical features as in adults, however, children have relative stable blood count for weeks to months (11, 12). The progression to secondary AML is variable, from slow period of several months to sudden transformation (7). It occurs in about 30% of cases of pediatric MDS, usually up to 2 years from diagnosis (13). In contrast to adult MDS, significance of subclassification into RAEB1 and RAEB 2 is not clear (14, 15).

PATHOGENESIS

According to pathogenesis, there are two types of MDS: primary or de novo and secondary. Despite extensive studies, particulary in pediatric cases, the causes of origin and progression of primary MDS are not yet fully understood. Pathogenesis of adult and pediatric MDS is multifactorial (16), as the ineffective haematopoiesis may be caused by heterogenous defects of BM stem cells. It is postulated, that genetic defects in the pluripotent progenitor cells lead to genetic instability with consequent numerous molecular and cellular abnormalities (7). Also excessive BM apoptosis plays role in pathogenesis (6). The partial therapeutic response on immunosuppression and autoimmune diseases symptoms suggest participation of deregulation of the immune system. In general, a lot of mechanisms take a part in pathogenesis with deregulation of cellular proliferation, differentiation, maturation and survival (17, 8). Especially genetic instability, epigenetic events, abnormal signal transduction, immune deregulation and finally BM microenvironment seem to have the most important role (18). Secondary MDS occuring after previous chemotherapy and / or radiotherapy is called „therapy-related”. Its incidence is growing due to longer survival of patients. Secondary MDS in childhood can arise in young patients with inherited BM failure disorders (IBMFS) (12). IBMFS represent a small and rare, but important group of diseases with risk of MDS evolution (19) characterised by failure of haematopoietic stem cell to produce blood cells with PB cytopenias (20). It is estimated that about 30% children with MDS share also constitutive disorder (7) Tab 2.

Also some others pediatric genetically determined diseases are associated with higher risk of MDS development, such as MonoMac syndrome, Down’s syndrome, neurofibromatosis, Bloom syndrome or Li-Fraumeni syndrome. In some of these cases, MDS represents the first presenting sing of the disease (15). The risk of clonal evolution and development of secondary MDS in aplastic anemia in children is high, as it occurs in about 25-40% of cases (21).
PROGNOSIS
Searching for prognostic variables for reliable stratifications of MDS patients, predictions of mortality and blastic transformation has started approx. in 1980 (22). Several prognostic scoring systems have been proposed, among them the most important and widely used are IPSS and IPSS-R. Clinical course of the disease is variable, but without therapy and often also despite treatment it is fatal (23). The median survival of patients with de novo MDS is about 2 years. Almost 30% of MDS patients die due to secondary AML, 40% due to BM failure complications such as infections, bleeding or iron overload and last 30% of patient due to cardiovascular complications and non-haematological malignancies. The median survival in therapy - related MDS is only about 3-8 months. The higher age of patients and associated comorbidities contribute to adverse prognosis (24, 4). There were several attempts to create a prognostic score of pediatric MDS, but none of them has been so far significantly applied in clinical practice, also IPSS shows only a limited value (14, 6).

Adult and also pediatric MDS can be divided into chronic low risk disease with slow progression (25) and more aggressive high risk with rapid leukemic evolution and short survival (26, 27).

Table 2. IBMFS, mutated genes and estimated prevalence of myeloid neoplasms (according to Niemeyer Ch. M., Kratz Ch. P., 2008).

<table>
<thead>
<tr>
<th>Disease</th>
<th>Mutated genes</th>
<th>Frequency of myeloid malignancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fanconi anemia</td>
<td>1 from 13 FANC genes</td>
<td>30-40%</td>
</tr>
<tr>
<td>Diamond-Blackfan anemia</td>
<td>RPS19, RPS24, RPS17</td>
<td>5%</td>
</tr>
<tr>
<td>Kostman’s disease</td>
<td>HAX1, ELA2, GFI1, WAS</td>
<td>30%</td>
</tr>
<tr>
<td>Schwachman-Diamond syndrome</td>
<td>SBDS</td>
<td>30-40%</td>
</tr>
<tr>
<td>Dyskeratosis congenita</td>
<td>DKC1, TERC, TERT, NOLA3</td>
<td>5%</td>
</tr>
<tr>
<td>FTD/AML*</td>
<td>RUNX1</td>
<td>?</td>
</tr>
</tbody>
</table>

Legend: *familial thrombocytopenia with propensity to MDS and AML

CLINICAL FEATURES
Most common manifestation of MDS of adult patients includes signs representing consequences of anemia (e.g. such as weakness, fatigue, palpitations etc.), while bleeding and infections are rare. Arthralgia or intermittent fever may be present in some patients. However, in about 50% of cases the early stages of the disease are asymptomatic (28, 3). MDS is suspected in cases with persistent cytopenia lasting more than 6 months, when other possible causes of cytopenia are excluded (29). In childhood, MDS manifests predominantly by neutropenia and thrombocytopenia (6), in a portion of children also constitutional disease's signs and symptoms may be present.

DIAGNOSIS
Early and correct diagnosis is essential for the choice of appropriate therapy and also for next life of patients. However, the diagnosis of MDS is challenging, complex and requiring close correlation of clinical symptoms, laboratory parameters and standardized examination of BM biopsies. It often represents a diagnosis „per exclusionem“ (28, 24).
diagnostic algorithm (as well as classification and prognostication of patients) is based on clinical parameters, the analysis of peripheral blood, BM aspirate and cytogenetic (4), but careful histological and immunohistochemical examination of BM biopsy is often necessary for the final diagnosis and precise classification of the patients (30).

For a correct interpretation of morphological and immunohistochemical findings by pathologist examining the BM biopsy, it is essential to have access to clinical data of the disease, laboratory findings, complete blood counts, results of flow cytometry and of cytogenetics (30).

Typical histomorphological MDS findings in BM biopsies of adults are represented by a distortion of BM microarchitecture, cytological changes and of topographical conditions, all they include (31):

1. Apparently unequal distribution of fat cells and thus different cellularity in various intertrabecular areas. In most cases, the BM is hypercellular, although there are also cases with a relatively normocellular BM. About 10% of cases show so-called hypoplastic MDS with reduced BM cellularity (5). However, it is important to assess the BM cellularity in comparison to the patient’s age.

2. significant differences in the size of hematopoietic islets,

3. increase of BM stromal reticulin fibers of varying grades,

4. a significant increase in the proportion of ring sideroblasts.

The ring sideroblasts are the red cells precursors with the abnormal accumulation of iron in the mitochondrias, which contain at least 5 siderotic granules encircling one third or more of the nucleus (24). They are considered as part of dysplastic changes in MDS, but can be detected also in other disorders and reactive changes as well, thus again they are not specific (24).

5. left shifted maturation of granulopoiesis, creating the „Abnormal Localization of Immature Precursors” (ALIP) phenomenon, characterized by clusters of less mature forms of granulocyte development in the interstitium, including bone’s trabecula and vascular structures.

6. varying degrees of erythropoiesis hyperplasia changing the physiological topographic distribution of its precursors – a typical feature, although somewhat unspecific, is dislocation of red cell precursors to paratrabecular areas. Typical but not exclusively specific features of MDS include anisocytosis and macrocytic and megaloblastic differentiation of erythroid precursors. Dysplastic changes may be nuclear (budding and/or bridging, karyorrhexis, multinuclearity and megaloblastoid changes) or cytoplasmic (presence of ring sideroblasts, vacuolization and periodic acid-Schiff positivity). The increased mitotic activity associated with maturation arrest is characteristic too (32, 28, 5). The assessment of dysplastic changes in both white and red cell lineages should be preferentially done in the BM aspirate as they are hardly detectable in BM biopsies.

7. abnormal cytomorphology and localization of megakaryocytes (mgk)

Number of mgk in BM in the MDS can be normal, increased or even reduced, but significant hypoplasia is rare (31). A typical feature of dysplastic mgk is a tendency to form free and tight clusters, in rare cases even large cohesive infiltrates (31). In contrast dysplasias of white and red cell precursors, the BM biopsy is preferred for evaluations of cytologic atypias im mgk series (33). They are represented by changes of nuclear lobulizations, e.g. hypolobated nuclei with usually dense chromatin, they may also occur as apoptotic forms (24, 32). Another morphological change is the multinucularity of mgk (28, 32). A strong indicator of the MDS is recognition of s.c. micromegakaryocytes (micromgk) (34), but they might be unrecognized in routine histomorphologic slides. To avoid such a mistake it is recommended, to apply immunohistochemical staining identifying typical mgk CD61 and CD41 antigens (32).

8. increased vascularization of BM, and

9. increased expression of CD34 antigen in the immature myeloid precursors.

Generally, a correct BM biopsy diagnosis of a „true” MDS including its differential distinction from reactive conditions, necessarily requires the application of immunohisto-
chemical methods. For the biopsy diagnosis of MDS it is recommended to apply constantly a standardized examination of expression of following antigens: CD34 (myeloblasts detection), myeloperoxidase (precursors of granulopoiesis), CD61 or CD41 (megakaryocytes and micromegakaryocytes), glycophorin A (precursors of erythropoiesis), CD20 (B cells), CD3 (T cells) and of CD68 (monocytes and macrophages).

In about 75% of children with RCC the BM cellularity is decreased with peculiar patchy distribution of haematopoesis in an otherwise fatty marrow (so-called „patchy pattern”) (11, 35). In these cases the essential marker of histopathological RCC diagnosis in BM biopsy is the presence of at least one island with at least 20 precursors with affected maturation and increased mitotic index. Granulopoiesis is reduced, with a possible shift to the left in maturation, but without increased proportion of myeloblasts. Elements of megakaryocytic series can be reduced or completely absent, while the finding of micromegakaryocytes is pathognomonic also for MDS in children (11, 35). In some cases, however, morphology is different and corresponds with RCMD of adult age (36), what is sometimes described as „diffuse type” of RCC (37).

DIFFERENTIAL DIAGNOSIS

A major problem in the morphological diagnosis of MDS is the determination, whether the presence of myelodysplasia is due to clonal disorder or results from some other factors (5). It is accepted that finding of manifesting dysplasia in more than 10% of cell of at least one cell lineage qualifies the MDS diagnosis. However, significant dysplasia may occur also in non-MDS cases and even in healthy persons too. Therefore some studies suggest that the assessment criteria might be tightened (32). Dysplastic changes alone are therefore not unique to MDS, as we know the number of diseases or factors that may cause secondary dysplasia and in adulthood. In particular, they might be summarized as follows:

1) megaloblastic anemia (MA) due to folate or vitamin B12 deficiency is one of the most common causes of secondary dysplastic changes in BM. These changes primarily affect the red lineage with megaloblastic differentiation and changes in topography of varying degrees (38). The total number of white series precursors can be easily reduced resulting in decreased physiological ratio of white and red series. Severe anemia may be associated with mild thrombocytopenia and the BM biopsy may show bizarre forms of megakaryocytes (39). Differentiation of MDS and MA based on the morphological (and immunohistochemical) BM changes is virtually impossible. Thus, clinical informations about a) levels of folic acid and vitamin B12, b) disorders of the stomach affecting the parietal cells (producing intrinsic factor needed for the absorption of vitamin B12 from food (39, 40), c) excessive abuse of alcohol (39) and finally, d) all chronic disease (infection, inflammation and malignancies) which lead to anemia of chronic disease (8) are very important and helpful.

2) chronic renal diseases can cause secondary dysplastic changes due to decreased production of erythropoietin, the primary regulator of erythropoiesis.

3) drugs can also cause BM changes that mimic MDS and induce cytopenias in PB. The most common are co-trimoxazole, mycophenolate mofetil, and other immunosuppressive agents, corticosteroids, most chemotherapeutic agents, growth factors and many other (8).

4) other - hypothyroidism, hepatopathy, haematopoietic and non-haematopoietic malignancies, autoimmune diseases, idiopathic inflammatory bowel disease, intoxication with heavy metals, virus diseases, etc. (8)

As in adults, the MDS diagnosis in children is usually morphological and therefore subjective (15). The diagnosis of pediatric MDS is challenging, especially in children with low blasts count without clonal marker, or in cases with discrete dysplasia only (14). In contrast, a significant dysplasia can also occur in reactive conditions of childhood. In pediatric disease, it is important to distinguish in particular:

1) MDS with low blasts count from aplastic anemia and other nonclonal diseases:
   a) aplastic anemia - defined by aplasia of all three hematopoietic cell lineages.
   Similar to most hypocellular RCC there is a marked decrease in BM cellularity below 30% for age with reduction of hematopoietic cells and substituting fatty marrow, but erythroid...
islands and micromegakaryocytes are not present in BM biopsy (Tab. 3.-11, 41, 35). BM architecture is essentially preserved with persisting non-hematopoietic cells, sometimes with the impression of a chronic inflammatory infiltrate (21, 35). However, if the BM cellularity is significantly reduced, to assess the status of haematopoietic cells can be extremely difficult (15). Overexpression of p53 in cases of MDS detected by immunohistochemistry appears to be useful (14). For the differential diagnosis of RCC and aplastic anemia it might be essential to repeat the BM biopsy (37) to receive and consider representative marrow spaces (12).

b) IBMFS with pancytopenia – shows in BM biopsies similar morphology to MDS and essentially indistinguishable changes from RCC. It is therefore necessary to exclude these syndromes mainly clinically using an extensive physical examination, assessment of family history and relevant laboratory and molecular measurements (11, 35). Histological examination can detect the progression of the diseases to secondary MDS, especially in the cases of increasing the blasts count or rising BM cellularity with progressive pancytopenia in the PB (11, 14).

c) secondary myelodysplasia due to viral infections (EBV, CMV, parvovirus), rheumatic diseases (juvenile idiopathic arthritis) or nutritional deficiencies (15).

**Table 3.** Morphological differences between refractory cytopenia of childhood and aplastic anemia (according to Baumann I., Führer M., Behrendt S., et al., 2012 and Chatterjee T. Choudhry V. P., 2013)

<table>
<thead>
<tr>
<th></th>
<th>Refractory cytopenia of childhood</th>
<th>Aplastic anemia</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Erythropoesis</strong></td>
<td>Patchy distribution</td>
<td>Lacking foci or a single small focus with &lt;10 cells with maturation</td>
</tr>
<tr>
<td></td>
<td>Left shifted maturation</td>
<td>Lacking or marked decrease, very few small foci with maturation</td>
</tr>
<tr>
<td></td>
<td>Increase mitoses</td>
<td></td>
</tr>
<tr>
<td><strong>Granulopoiesis</strong></td>
<td>Marked decrease</td>
<td>Lacking or very few, no dysplastic megakaryocytes</td>
</tr>
<tr>
<td></td>
<td>Left shift</td>
<td></td>
</tr>
<tr>
<td><strong>Megakaryopoiesis</strong></td>
<td>Marked decrease</td>
<td>Lacking or very few, no dysplastic megakaryocytes</td>
</tr>
<tr>
<td></td>
<td>Dysplastic changes</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Micromegakaryocytes</td>
<td></td>
</tr>
<tr>
<td><strong>Lymphocytes</strong></td>
<td>May be increased focally or dispersed</td>
<td>May be increased focally or dispersed</td>
</tr>
<tr>
<td><strong>CD34+ cells</strong></td>
<td>No increase</td>
<td>No increase</td>
</tr>
</tbody>
</table>

2) **MDS with excess blasts from the primary, de novo AML.**

There are significant differences in clinical features, cytogenetics and in response to therapy between MDS and de novo AML reflecting fundamental biological differences (7). Primary AML characterized by a specific recurrent translocations is chemosensitive, on the contrary, MDS and secondary AML are chemoresistant and defined by numerous chromosomal abnormalities (14). In contrast to adults, it is not yet well-defined boundary threshold of blast percentage to separate RAEB from AML. That was the reason to maintain in the classification of pediatric MDS the subgroup RAEB in transformation ("RAEB-T"), with the proportion of blasts in BM between 20-29%. In such borderline cases with dysplasia and in the absence of cytogenetic markers, it is recommended to repeat the BM examination two weeks later. If the blast count has increased above 30%, or organomegaly or a significant leucocyte increase in the PB has appeared, it is most likely the condition represents a primary AML (7). If the blast count is stable, then an arbitrary period of 4 weeks is suggested before establishing a diagnosis of RAEB-T. However, most children with primary AML present from the beginning as a frank AML (14).
In the presence of specific translocations, e.g. t (8; 21) (q22; q22) and t (15; 17) (q22; q12), or inversion, for example inv16 (p13; q22), the disease should be considered as AML, irrespective of the blast proportion in BM (7, 14).

TREATMENT

Differences between adult and pediatric MDS are also reflected in therapy options. There are numbers of therapeutic procedures (supportive, stimulating, immunomodulatory with lenalidomide) in adult patients with low and intermediate risk disease, which the main aim to extend the life of hematopoietic cells (42). In patients at high risk, the dominating effort is to delay leukemic evolution by hypomethylating agents, intensive chemotherapy and transplantation of hematopoietic stem cells (33). Compared to the adult population, a major therapeutic goal in childhood is to cure pediatric patients (15) and the only potentially curative option remains transplantation of hematopoietic stem cells. In some pediatric patients, immunosuppressive therapy may be effective (10). Hypomethylating agents have the potential to modify the course of the disease and have become a standard in the treatment of disease in the adult patient population, but the effects in children still remains unclear (43, 15).

CONCLUSIONS

MDS both in adults and in children is a severe disease, regardless of subtype. Diagnosis, classification and selection of optimal treatments are currently based on a comprehensive evaluation of the patient, including assessment of morphological, immunophenotypic and cytogenetic findings. Although histomorphological and immunohistochemical analysis of BM biopsy is invasive and time-consuming examination, it has its value in the diagnosis, differential diagnosis and evaluation of therapeutic effect. However, clinicopathological mutual cooperation is necessary for correct interpretation of observed changes.

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