

Type of Article: Review

Title: Clonal Evolution in Myelodysplastic Syndromes: From Molecular Pathogenesis to Targeted Therapeutic Strategies

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Clinical Question Box

How does clonal evolution influence prognosis and therapeutic decision-making in patients with myelodysplastic syndromes?

Clonal evolution underlies the biological heterogeneity, disease progression, and treatment resistance observed in myelodysplastic syndromes. The accumulation and selection of genetic abnormalities shape clonal architecture, influencing prognosis and response to therapy. High-risk mutations and co-mutation patterns identify potential beneficiaries from early aggressive approaches, including targeted therapy or allogeneic stem cell transplantation. Molecular prognostic models such as the IPSS-M improve risk stratification by incorporating genetic data, but their static nature limits the prediction of disease trajectory, highlighting the importance of dynamic, longitudinal molecular monitoring to guide individualized treatment decisions.

Abstract

Myelodysplastic syndromes (MDS) constitute a heterogeneous group of clonal hematopoietic stem cell disorders characterized by ineffective hematopoiesis, peripheral cytopenias, and a significant risk of progression to acute myeloid leukemia. The core of MDS pathogenesis is a dynamic process of clonal evolution: initiating driver mutations in hematopoietic stem cells, under the combined pressure of genomic instability and the bone marrow microenvironment, leads to the continuous acquisition of new genetic abnormalities, forming a complex clonal architecture. This process is the fundamental cause of disease progression, therapeutic resistance, and clinical heterogeneity. This review systematically elaborates on the genetic and biological basis of clonal evolution, including frequent mutations and their interactions in key pathways such as epigenetic regulation, RNA splicing, and signal transduction. It analyzes how clonal competition, therapeutic selection pressure, and immune microenvironment editing drive clonal dynamics. Concurrently, the review details the evolution of prognostic assessment systems from the International Prognostic Scoring System-Revised to the Molecular International Prognostic Scoring System (IPSS-M), marking a revolutionary shift into the molecular era, and discusses both the clinical value and existing limitations of IPSS-M. Building upon this, the article comprehensively depicts the advancement of treatment strategies from immune modulation and epigenetic intervention to the era of precision targeting, integrating the latest evidence and optimized approaches for both lower-risk and higher-risk MDS. Finally, it proposes that future MDS management must integrate dynamic clonal monitoring, biomarker-based strategies, and multidisciplinary collaboration to achieve individualized and precise whole-course MDS management.

Keywords: Myelodysplastic syndromes, Clonal evolution, Gene mutation, IPSS-M,

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1. Introduction

The diagnostic framework and conceptual understanding of myelodysplastic syndromes (MDS) have evolved substantially. Once considered refractory anemia or a preleukemic condition, MDS is now recognized as a heterogeneous group of clonal hematopoietic neoplasms.¹ Both the World Health Organization Classification of Tumours of Haematopoietic and Lymphoid Tissues, 5th Edition (WHO-HAEM5) and the International Consensus Classification (ICC) have adopted the term myelodysplastic neoplasms, underscoring their neoplastic nature.^{2,3} Unlike acute myeloid leukemia (AML), MDS originate not from a single fully transformed leukemic stem cell but from a genetically unstable hematopoietic stem and progenitor cell clone undergoing continuous evolution.⁴ The disease course is therefore defined by clonal evolution, characterized by the stepwise accumulation and competition of somatic mutations and cytogenetic abnormalities, resulting in complex oligoclonal architectures.⁵

Clonal evolution is central to the pathophysiology of MDS, driving disease heterogeneity, influencing progression risk, and shaping therapeutic response.⁶ Under treatment-induced selective pressures, dominant subclones may emerge through immune escape mechanisms or the acquisition of resistance-conferring mutations, ultimately leading to relapse or disease progression.⁷ Consequently, elucidating clonal evolutionary dynamics is essential for predicting disease trajectories, understanding mechanisms of treatment failure, and informing the development of novel therapeutic strategies. Advances in MDS therapy have paralleled increasing biological insight into disease pathogenesis, with prognostic assessment evolving from morphology-based classification systems toward integrated models incorporating cytogenetic and molecular features.⁸ Parallely, treatment approaches have progressed from predominantly supportive care and immunomodulatory therapies to epigenetic agents and targeted therapies

directed against specific genetic alterations or signaling pathways.⁹ Despite these advances, allogeneic hematopoietic stem cell transplantation (allo-HSCT) remains the only potentially curative treatment, with clinical outcomes closely linked to the underlying clonal architecture and patterns of clonal evolution.¹⁰

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2. MDS: Its Clonal Nature and Genetic Basis

2.1 Clonal Origin and Evolutionary Drivers

MDS are clonal disorders arising from hematopoietic stem and progenitor cells, characterized by pervasive genomic instability, including telomere dysfunction and impaired DNA damage–repair mechanisms.¹¹ This instability provides the substrate for clonal evolution by promoting an increased mutational burden. Early in the disease course, clonal abnormalities may be limited to morphological dyshematopoiesis; however, with progression, particularly during transformation to higher-risk MDS or AML, the genetic features of dominant clones become increasingly apparent.⁵ Clonal evolution in MDS is driven by the interplay of intrinsic and extrinsic forces. Intrinsically, the continuous acquisition of genetic lesions fuels clonal diversification, whereas extrinsic selective pressures, most notably therapeutic interventions and the bone marrow microenvironment, shape clonal competition, fitness, and evolutionary trajectories.¹²

2.2 Genetic Landscape: Mutational Spectrum and Clinical Relevance

Somatic mutations are detected in over 90% of MDS patients, with more than half harboring multiple concurrent mutations, underscoring the marked genetic complexity of the disease.¹³ These alterations cluster within several key biological pathways (Figure 1):

2.2.1 Ribonucleic Acid (RNA) Splicing Machinery

Mutations affecting components of the RNA splicing machinery, including splicing factor 3B subunit 1 (SF3B1), serine/arginine-rich splicing factor 2 (SRSF2), U2 small nuclear RNA auxiliary factor 1 (U2AF1), and zinc finger RNA-binding motif and serine/arginine-rich protein 2 (ZRSR2), are among the most frequent genetic alterations in MDS. Mutations in SF3B1 are closely associated with the presence of ring sideroblasts and are generally linked to a favorable prognosis; however, this advantage is abrogated by specific co-occurring mutations.¹⁴ For instance, the co-

mutation of SF3B1 with genes such as runt-related transcription factor 1 (RUNX1), BCL6 corepressor (BCOR), BCL6 corepressor-like 1 (BCORL1), NRAS proto-oncogene, GTPase (NRAS), serine/arginine-rich splicing factor 2 (SRSF2), or stromal antigen 2 (STAG2) defines a molecular subset with significantly worse outcomes compared to SF3B1-mutant cases without these alterations.¹⁵ Similarly, emerging evidence suggests that deletion of the long arm of chromosome 5 (del(5q)), combined with an SF3B1 mutation, may also identify a subgroup with an independently poor prognosis, a risk interaction not yet fully captured by current prognostic models like the IPSS-M.¹⁶ Mutations in ZRSR2, which occur more frequently in male patients due to its X-linked location, are often associated with relatively favorable clinical outcomes, although their prognostic significance must be interpreted in the broader mutational context.¹⁷

2.2.2 Epigenetic Regulators

Epigenetic dysregulation represents a central pathogenic mechanism in MDS and involves mutations in genes regulating DNA methylation and chromatin structure.¹⁸ Key regulators of DNA methylation include ten-eleven translocation methylcytosine dioxygenase 2 (TET2) and DNA methyltransferase 3 alpha (DNMT3A), whereas epigenetic regulation of chromatin structure is mediated by histone-modifying genes such as additional sex combs-like transcriptional regulator 1 (ASXL1), enhancer of zeste homolog 2 (EZH2), and lysine methyltransferase 2A (KMT2A).¹⁹ Although mutations in DNMT3A, TET2, and ASXL1 may also be detected in age-related clonal hematopoiesis and, therefore, lack independent diagnostic specificity, they exert substantial biological effects in MDS. Particularly, mutations in DNMT3A and ASXL1 are well-established adverse prognostic markers associated with inferior survival.²⁰

2.2.3 Transcription Factors

Mutations in hematopoietic transcription factors, including RUNX1, GATA binding protein 2

(GATA2), CCAAT/enhancer-binding protein alpha (CEBPA), and ETS variant transcription factor 6 (ETV6), disrupt normal hematopoietic differentiation and contribute to genomic instability.²¹ Notably, germline mutations in GATA2 and ETV6 underlie inherited MDS and AML predisposition syndromes, highlighting the role of transcriptional dysregulation in both sporadic and familial disease.²²

2.2.4 Tumor Suppressor and Signaling Pathway Genes

Alterations in tumor suppressor genes and signaling pathways are strongly associated with disease aggressiveness. Mutations in tumor protein p53 (TP53) represent the most powerful adverse prognostic factor in myelodysplastic syndromes. Biallelic inactivation of TP53 is frequently associated with complex karyotypes, profound resistance to therapy, and extremely poor overall survival. Additionally, mutations affecting signaling pathways, including those involving rat sarcoma viral oncogene homolog (RAS) family members, confer proliferative and survival advantages to malignant clones.²³

Importantly, the clinical impact of individual mutations is highly dependent on their combinatorial patterns. For instance, the favorable prognostic effect associated with SF3B1 mutations is lost when co-occurring with alterations in RUNX1, BCL6 corepressor, BCL6 corepressor-like 1, neuroblastoma RAS viral oncogene homolog, SRSF2, or STAG2. Similarly, in MDS with deletion of the long arm of chromosome 5, coexisting TP53 mutations predict poor responses to lenalidomide and inferior clinical outcomes.²⁴ The biological mechanisms underlying these adverse mutational interactions remain incompletely defined.

2.3 Germline Predisposition and Clonal Evolution

A subset of MDS arises from a defined germline predisposition and represents a biologically distinct entity. Germline mutations in GATA2 and sterile alpha motif domain-containing 9/ sterile

alpha motif domain-containing 9-like (SAMD9/SAMD9L) markedly increase the risk of MDS and AML, and are frequently associated with syndromic immunodeficiency. GATA2 deficiency and SAMD9/SAMD9L syndromes disrupt the development and function of myeloid and lymphoid immune cells, including monocytes, dendritic cells, and natural killer cells, resulting in profound impairment of both innate and adaptive immunity.²⁵ Affected individuals are highly susceptible to refractory and recurrent infections, such as nontuberculous mycobacterial disease, invasive fungal infections, including aspergillosis, and severe viral infections, including Epstein–Barr virus and human papillomavirus.²⁶ These infections are a primary direct cause of mortality and, critically, promote the establishment of a chronically inflammatory bone marrow microenvironment. This sustained immunoinflammatory state functions as a powerful extrinsic driver that accelerates clonal evolution and perpetuates a self-reinforcing cycle linking immune deficiency, infection, inflammation, and clonal progression.²⁷ These genes are essential for normal immune development, and their disruption results in profound innate and adaptive immune deficiency, predisposing affected individuals to recurrent mycobacterial, invasive fungal, and oncogenic viral infections.²⁸ The resulting chronically inflammatory bone marrow microenvironment imposes strong selective pressure that accelerates clonal evolution.

In SAMD9/SAMD9L-associated disease, clonal evolution often involves somatic genetic rescue, including reversion mutations or uniparental disomy of chromosome 7q, which transiently alleviate the deleterious effects of the germline mutation.²⁹ However, this unstable equilibrium reflects marked genomic instability and predisposes to subsequent acquisition of leukemogenic lesions, particularly monosomy 7 or deletion of chromosome 7q and activating RAS pathway mutations, ultimately accelerating disease progression.³⁰

2.4 Leukemia-Defining Molecular Subtypes

With advances in molecular understanding of MDS, contemporary classification systems have entered a genome-first era, in which certain molecular abnormalities supersede morphologic features, particularly blast percentage, as defining diagnostic criteria.^{2,3} Identification of these leukemia-defining molecular lesions is central to precision diagnosis and directly guides therapeutic strategy, most notably in the nucleophosmin 1 (NPM1)–mutated, MDS1 and EVI1 complex locus (MECOM)–rearranged, also called ecotropic viral integration site 1 (EVI1)–rearranged, and nucleoporin 98 (NUP98)–rearranged subtypes.³¹

2.4.1 NPM1-Mutated Subtype

Both WHO-HAEM5 and ICC define AML by the presence of an NPM1 mutation, irrespective of bone marrow blast percentage (designated AML with mutated NPM1 in the ICC).^{2,3} This approach reflects strong evidence that NPM1 mutations drive a potent leukemogenic program, with clinical behavior and treatment responsiveness comparable to conventional blast-defined AML. Even when presenting with MDS-like morphology, patients typically respond favorably to standard intensive AML induction therapy, consisting of continuous-infusion cytarabine for 7 days combined with an anthracycline administered for 3 days.³² Such patients achieve high complete remission rates and derive clear benefit from subsequent consolidation chemotherapy and/or allo-HSCT. Overall outcomes closely mirror those of typical AML and are clearly distinct from other high-risk MDS subtypes.³³

2.4.2 MECOM–Rearranged Subtype

WHO-HAEM5 recognizes AML with MECOM rearrangement as a distinct entity and mandates an AML diagnosis whenever a MECOM rearrangement is detected, regardless of blast percentage.² In contrast, ICC applies blast-based stratification, classifying cases with <10% blasts as MDS with MECOM rearrangement and those with $\geq 10\%$ blasts as AML with MECOM

rearrangement.³ Despite these differences, both systems uniformly identify MECOM rearrangement as an ultra - high-risk feature, associated with poor therapeutic response and adverse outcomes.

2.4.3 NUP98-Rearranged Subtype

WHO-HAEM5 newly defines AML with NUP98 rearrangement as a distinct entity, allowing an AML diagnosis solely based on a NUP98 rearrangement, even when blasts are <20%.² ICC classifies these cases under AML with other rare recurrent translocations.³ In East Asian adult AML cohorts, NUP98 rearrangements occur in approximately 5% of cases and are strongly associated with chemoresistance and poor clinical outcomes, underscoring their prognostic significance.³⁴

3. Evolution of Prognostic Assessment Systems

3.1 *The Transformative Significance of IPSS-M*

Accurate prognostic stratification is central to individualized treatment in MDS. The IPSS and its revised version (IPSS-R) stratify risk based on bone marrow blast percentage, cytogenetics, and cytopenias, with IPSS-R remaining the most extensively validated model in clinical practice.³⁵ The IPSS-M, introduced in 2022, builds on IPSS-R by incorporating the mutational status of 31 genes, thereby ushering MDS prognostication into the molecular era.²⁴ Real-world studies show that IPSS-M reclassifies approximately 34% of patients, refining the heterogeneous IPSS-R intermediate-risk group into “moderate–low” and “moderate–high” categories and improving risk precision.³⁶ Importantly, IPSS-M identifies occult high-risk disease among patients deemed low risk by IPSS-R due to adverse mutations such as biallelic TP53 inactivation or ASXL1 mutations, supporting earlier or more aggressive intervention.³⁷ Emerging data further indicate that IPSS-M outperforms IPSS-R in predicting outcomes after allogeneic hematopoietic stem cell transplantation, enabling more accurate molecular guidance for transplant timing and candidacy.³⁸

3.2 *Clinical Limitations of IPSS-M*

Despite its paradigm-shifting role in molecular risk stratification, the IPSS-M has several important limitations in clinical practice. Although it incorporates the mutational status of 31 genes, gene coverage remains incomplete. Several clinically relevant genes with established prognostic or therapeutic implications in MDS are excluded, including MECOM, ZRSR2, Fanconi anemia complementation group L (FANCL), and phosphatidylinositol glycan anchor biosynthesis class A (PIGA) (Table 2). Additionally, key germline predisposition genes such as DEAD-box helicase 41 (DDX41) and SAMD9/SAMD9L are not included, potentially leading to underestimation of risk

and omission of inherited susceptibility. Such oversights may cause clinical misjudgment and a delay in appropriate therapy. For instance, conventional karyotyping is insufficient to detect MECOM/EVI1 rearrangements due to limitations in resolution and metaphase quality. Thus, the omission of this ultra-high-risk feature may lead to significant underestimation of risk and delay intensive therapy like transplantation.³⁹

A second major limitation is the insufficient modeling of gene–gene interactions and clonal dynamics. IPSS-M assigns weighted risk scores to individual mutations but does not fully capture context-dependent effects of co-mutation patterns, such as the variable prognostic impact of SF3B1 mutations in specific genetic backgrounds.⁴⁰ More fundamentally, IPSS-M represents a static molecular snapshot at diagnosis, whereas MDS is characterized by continuous clonal evolution driven by genomic instability and treatment-related selective pressures, limiting accurate prediction of disease trajectory.⁴¹

Relatedly, IPSS-M does not comprehensively integrate mutational burden or variant allele frequency dynamics, which provide critical insights into clonal size, treatment response, and emerging resistance. Although the allelic state is partially considered for selected genes such as TP53, this level of granularity is not uniformly applied across other prognostically relevant mutations, including STAG2 and RUNX1. Further, IPSS-M does not consider clonal architecture, a defining feature of MDS reflecting hierarchical organization and dynamic subclonal evolution during therapy.⁴¹

Finally, practical and conceptual limitations affect implementation. IPSS-M requires comprehensive next-generation sequencing, increasing technical complexity, cost, and turnaround time compared to IPSS-R.⁴² Moreover, it does not incorporate other prognostically relevant factors such as bone marrow fibrosis, immune dysfunction, inflammatory microenvironmental features,

age, or comorbidities, all of which influence treatment tolerance and real-world clinical decision-making.

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4. Optimizing Treatment Strategies for Lower-Risk MDS

In patients with lower-risk MDS, anemia management remains the central therapeutic priority. Effective correction of anemia is critical not only for improving quality of life and reducing transfusion dependence but also for conferring long-term survival benefits.⁴³ Historically, treatment strategies relied predominantly on supportive care measures and erythropoiesis-stimulating agents (ESAs). However, advances in the understanding of erythroid dysregulation have led to the development of novel agents that more directly target ineffective erythropoiesis and disease biology.

4.1 Erythroid Maturation Agents

Luspatercept, a transforming growth factor- β (TGF- β) superfamily ligand trap, enhances late-stage erythroid maturation by inhibiting aberrant SMAD2/3 signaling. Its clinical efficacy was established in the phase III MEDALIST trial, which demonstrated significant benefit in ESA-refractory, transfusion-dependent lower-risk MDS with ring sideroblasts.⁴⁴ Subsequently, the phase III COMMANDS trial showed superior efficacy of luspatercept compared to ESAs in treatment-naïve, transfusion-dependent lower-risk MDS, supporting its use as a first-line therapeutic option.⁴⁵ Long-term follow-up from MEDALIST confirmed improved overall survival among patients achieving transfusion independence for at least eight weeks, and real-world data published in 2025 reported a 75% reduction in mortality risk among hematologic responders compared to non-responders.^{46,47}

KER-050 (elritercept), a next-generation activin receptor type IIA ligand trap, extends this therapeutic approach by inhibiting activin A and related TGF- β superfamily ligands.⁴⁸ Early-phase clinical studies have demonstrated encouraging efficacy and favorable safety profiles in patients with lower- to intermediate-risk MDS, positioning KER-050 as a promising addition to anemia-

directed treatment strategies.⁴⁹

4.2 Disease-Modifying Approaches

Imetelstat, a first-in-class telomerase inhibitor, has emerged as a potential disease-modifying therapy in lower-risk MDS. In the phase III IMerge trial, imetelstat significantly increased rates of transfusion independence in ESA-refractory patients. Importantly, treatment was associated with reductions in AML transformation risk and declining variant allele frequencies of driver mutations, suggesting biological activity beyond symptomatic anemia control.⁵⁰

4.3 Iron Chelation and Supportive Care

Besides pharmacological therapies, optimal supportive care remains a critical component of management. Treatment of transfusion-related secondary iron overload with iron chelation therapy has been consistently associated with improved survival in lower-risk MDS.⁵¹ Collectively, these data underscore that effective anemia control, through erythroid maturation agents, emerging disease-modifying therapies, and proactive management of iron overload, remains a cornerstone for optimizing long-term outcomes in lower-risk MDS.

4.4 Exploration of Spliceosome Modulation

Direct targeting of the spliceosome, a core machinery frequently mutated in MDS, has been explored. H3B-8800, an oral small molecule modulator of the SF3b complex, demonstrated the potential to induce transfusion independence in a subset of patients with spliceosome-mutant, lower-risk MDS in a phase I trial.⁵² However, overall efficacy was modest, with no standard complete or partial remissions observed, and clinical development has been discontinued.⁵³ This experience validates the spliceosome as a therapeutic target but highlights the challenges in achieving robust clinical efficacy with single-agent modulation.

In the management of lower-risk myelodysplastic syndromes, adequate doses of ESAs remain

the classic first-line therapy for patients with serum erythropoietin (EPO) levels <500 IU/L.⁵⁴ Luspatercept has also emerged as an established first-line option, particularly for patients with SF3B1 mutations or ring sideroblast (RS)-associated MDS, in whom it demonstrates favorable efficacy in those with EPO levels between 200–500 IU/L and even <200 IU/L.⁵⁵ In addition, the MEDALIST trial showed activity in non-RS, non-SF3B1 lower-risk MDS.⁴⁴ The phase 3 COMMANDS trial further demonstrated superior erythroid responses with luspatercept compared with darbepoetin alfa in ESA-naïve, transfusion-dependent lower-risk MDS, establishing luspatercept as a frontline option for a broader patient population irrespective of RS status.⁴⁵ In real-world practice, the choice between luspatercept and ESAs should be individualized according to local reimbursement policies and patient financial considerations.

Imetelstat is indicated for transfusion-dependent lower-risk MDS patients who have failed, lost response to, or are ineligible for ESAs.⁵⁰ It is particularly valuable in patients with baseline EPO >500 IU/L, a population with an anticipated poor response to ESAs and reduced benefit from luspatercept. The U.S. Food and Drug Administration approval was based on the IMerge phase 3 trial, and imetelstat is now included as a National Comprehensive Cancer Network Category 1 preferred option for this indication.⁵⁶

For patients with del(5q) MDS, lenalidomide remains the standard of care.⁵⁷ However, its efficacy is attenuated with increasing TP53 mutation burden. In del(5q) patients harboring TP53 mutations, a sequential strategy using frontline hypomethylating agents (HMAs) to reduce the TP53 clonal burden followed by lenalidomide has shown potential efficacy in a recently reported case, suggesting a novel therapeutic approach for this challenging subset.⁵⁸

For patients with EPO >500 IU/L who exhibit clinical features predictive of response to immunosuppressive therapy, including age ≤ 60 years, bone marrow blasts $\leq 5\%$, hypocellular

marrow, presence of a paroxysmal nocturnal hemoglobinuria (PNH) clone, or STAT3-mutant cytotoxic T-cell clones, immunosuppressive therapy combined with thrombopoietin receptor agonists may achieve favorable outcomes.⁵⁹ When these strategies are ineffective, treatment with hypomethylating agents or enrollment in clinical trials should be considered. For patients with significantly compromised quality of life, early evaluation for HSCT is recommended.⁶⁰

5. Optimizing Treatment Strategies for Higher-Risk MDS

The primary therapeutic goals in higher-risk MDS(encompassing moderate-high, high, very high risk categories) are to delay disease progression, eradicate malignant clones, and prevent transformation to AML. While HMAs such as azacitidine and decitabine(including the oral fixed-dose combination of decitabine with cedazuridine) have long served as the therapeutic backbone, treatment paradigms are rapidly evolving toward mechanism-driven combination regimens and precision-based strategies designed to overcome resistance and address high-risk disease biology.

5.1 Refinement and Expansion of Combination Therapies

5.1.1 HMA Plus BCL-2 Inhibition

Combination therapy with azacitidine or decitabine plus the BCL-2 inhibitor venetoclax, adapted from its success in AML, has demonstrated rapid and deep cytoreductive responses in high-risk MDS. Across real-world and early clinical studies, overall response rates of 62–80% have been reported, and this regimen is widely used as an intensified strategy or bridge to allo-HSCT.^{61,62} However, in the randomized phase III VERONA trial, the addition of venetoclax to HMA therapy failed to confer a statistically significant improvement in overall survival compared to HMA monotherapy (hazard ratio, 0.908).⁶³ Although higher response rates were observed in molecularly adverse subsets such as TP53-mutated disease, survival benefits remained modest. Consequently, HMA–venetoclax is best regarded as an effective disease-modifying and bridging regimen, rather than a new standard first-line therapy.

5.1.2 HMA Plus Isocitrate Dehydrogenase (IDH) Inhibitors

Targeting mutant IDH constitutes a precision-medicine strategy applicable to approximately 3–13% of patients with MDS.^{64,65} In IDH1-mutant disease, the IDH1 inhibitor ivosidenib is approved for relapsed or refractory MDS following failure of HMA therapy.⁶⁶ Its combination with

azacitidine is currently being evaluated in the frontline setting in the randomized phase III PyramIDH trial.⁶⁷

Similarly, the IDH2 inhibitor enasidenib is under investigation as monotherapy and in combination with hypomethylating agents across multiple disease settings.^{68,69} Early-phase clinical studies have demonstrated feasibility and biological activity, although definitive efficacy data and survival outcomes require further validation in larger controlled trials.⁷⁰ These approaches exemplify the ongoing shift toward genomically informed treatment selection in higher-risk MDS.

5.1.3 Next-Generation BCL-2 Inhibitors

To address the resistance and toxicity associated with first-generation BCL-2 inhibition, next-generation BCL-2 inhibitors such as lisaftoclax are being developed. Lisaftoclax exhibits optimized selectivity and tolerability, and early phase I/II studies combining lisaftoclax with azacitidine in treatment-naïve higher-risk MDS and AML have demonstrated manageable safety profiles and encouraging antileukemic activity.⁷¹ This approach represents a promising evolution of BCL-2–based therapy.

5.2 *TP53-Directed Therapy*

TP53-mutated MDS, frequently associated with complex karyotypes, represent one of the most adverse prognostic subtypes, largely resistant to conventional therapies and constituting a major unmet clinical need.⁷²

Efforts to directly restore p53 function have thus far produced mixed results. Notably, the phase III trial evaluating the p53 reactivator APR-246 (eprenetapopt) in combination with azacitidine failed to achieve its primary endpoints of complete remission and overall survival improvement.⁷³ Consequently, therapeutic strategies are increasingly shifting toward intensified

combination approaches, including triplet regimens incorporating APR-246, azacitidine, and venetoclax. In contrast, MDM2 inhibitors, which are restricted to TP53–wild-type disease, have demonstrated limited clinical activity in MDS, prompting growing interest in next-generation approaches such as PROTAC-based targeted protein degraders.⁷⁴

5.3 Modulation of the Immune Microenvironment

Targeting the immunosuppressive bone marrow microenvironment has emerged as an important complementary therapeutic strategy in higher-risk MDS. Immune checkpoint modulation through TIM-3 inhibition has revealed encouraging early activity; notably, the anti–TIM-3 antibody sabatolimab, when combined with HMAs, demonstrated favorable tolerability and an overall response rate of 62.9% in a phase Ib study, and the randomized phase III STIMULUS-MDS2 trial has completed accrual with results awaited.^{75,76} Parallely, macrophage-directed approaches are gaining traction, exemplified by bexmarilimab, an antibody targeting Clec4e-1 on immunosuppressive macrophages, which, in combination with azacitidine, achieved a preliminary overall response rate of 45% in HMA–refractory higher-risk MDS, highlighting the therapeutic potential of macrophage reprogramming within the bone marrow niche.⁷⁷ However, first-generation CD47 blockade strategies (e.g., magrolimab in combination with HMAs) have been constrained by dose-limiting toxicities and inconsistent clinical efficacy.⁷⁸

6. Allo-HSCT

Allo-HSCT remains the only potentially curative therapy for MDS.⁷⁹ Long-term outcomes indicate a 5-year overall survival of approximately 33.9-52.8% following transplantation.⁸⁰ Optimal transplant outcomes depend on careful patient selection, effective pre-transplant disease debulking, and post-transplant surveillance, including minimal residual disease (MRD) monitoring using high-throughput sequencing to enable timely preemptive intervention.⁸¹

6.1 Risk-Based Transplant Selection

The IPSS-M scoring system has improved identification of patients most likely to benefit from allo-HSCT; however, accurate diagnostic classification remains the cornerstone of transplant decision-making. Molecular and cytogenetic features increasingly guide both the indication and timing of transplantation, emphasizing the need for integrated clinicogenomic assessment.²⁴ Under the WHO-HAEM5 classification, cases harboring NPM1 mutations, MECOM rearrangements, or NUP98 rearrangements are now classified as AML or distinct AML subtypes rather than MDS. For these patients, frontline management should follow AML treatment principles, including intensive induction chemotherapy and early incorporation of allo-HSCT, with urgency comparable to that applied in high-risk AML.

In patients meeting diagnostic criteria for MDS, decisions regarding allo-HSCT should be individualized according to molecular subtype. Indolent subtypes, including isolated SF3B1 mutation and biallelic TET2 mutations, generally do not warrant allo-HSCT.⁸² Subtypes requiring individualized consideration, such as del(5q) with high-risk co-mutations, DDX41-mutated disease, and U2AF1 or SRSF2 mutations, require integration of molecular features, blast burden, and clinical factors to guide transplant timing.⁸³ Conversely, high-risk molecular subtypes, including TP53-mutated disease with complex karyotype, -7/del(7q) with SETBP1 or RAS

mutations, and EZH2–ASXL1 co-mutations, strongly favor early allo-HSCT and consideration of clinical trial enrollment when feasible.⁸⁴

Germline predisposition syndromes substantially complicate allo-HSCT decision-making. Patients with germline susceptibility (e.g., GATA2 or SAMD9/SAMD9L mutations) frequently exhibit baseline immune dysfunction, recurrent infections, and organ impairment, necessitating highly individualized risk–benefit assessment.⁸⁵ Donor selection is particularly challenging, as germline genetic screening is mandatory to avoid related donors carrying the same pathogenic variants.

6.2 Infection Control

Pre-transplant infection control represents a major challenge in patients with inherited predisposition syndromes. Active or refractory infections, particularly nontuberculous mycobacterial infections and invasive fungal diseases, may constitute contraindications to transplantation or carry an extremely high risk of transplant-related mortality.⁸⁶ Achieving adequate infection control often requires prolonged, multidisciplinary antimicrobial therapy, which can delay allo-HSCT and increase the risk of disease progression; nevertheless, optimization of infectious status before conditioning remains essential, albeit frequently difficult.⁸⁷

Post-transplant outcomes are further influenced by delayed immune reconstitution due to underlying immunodeficiency, resulting in high rates of severe infections, including bacteremia (22.3%) and pulmonary fungal infections (12.4%).⁸⁸ Reduced tolerance to conditioning regimens, related to poor baseline performance status and prior infections, necessitates individualized conditioning intensity and tailored graft-versus-host disease prophylaxis to balance disease control with transplant-related toxicity.

7. Further Direction

Future therapeutic strategies for MDS must be dynamic, precision-driven, and grounded in a comprehensive understanding of clonal evolution. Longitudinal monitoring of variant allele frequencies and clonal architecture using high-throughput sequencing enables response-adaptive treatment strategies, with post-treatment MRD status emerging as a powerful predictor of relapse and a critical guide for timely preemptive intervention.⁸⁹ Moving beyond static risk models, integration of artificial intelligence–driven, multiparametric frameworks that incorporate clinical variables, molecular profiles, and imaging data holds promise for predicting clonal evolutionary trajectories and optimizing treatment sequencing, thereby enabling individualized, dynamic prognostic systems that surpass the limitations of current tools such as IPSS-M.⁹⁰

Ultimately, effective control of clonal evolution will require rationally designed therapeutic combinations, administered concurrently or sequentially, to target both dominant and emerging subclones. Future drug development should prioritize novel and historically challenging targets, including transcription factors, downstream effectors of spliceosome dysregulation, and components of the bone marrow microenvironment, as well as synthetic lethality–based strategies to overcome adaptive resistance mechanisms.⁹¹

8. Conclusion

MDS represent a biologically heterogeneous group of disorders in which clinical outcomes are increasingly shaped by underlying molecular architecture and clonal dynamics. Though advances in genomic profiling, risk stratification, and targeted therapeutics have refined diagnosis and treatment selection, durable disease control remains limited for many high-risk subgroups. Future progress will depend on integrating precision medicine, dynamic MRD-guided strategies, and rational combination therapies to overcome clonal evolution and resistance, with allogeneic transplantation and clinical trial enrollment remaining essential components of curative and disease-modifying approaches.

Acknowledgments: None

Funding statement: None

Ethics approval statement: Not applicable

Patient consent statement: Not applicable

Conflict of interest disclosure: The authors declare that they have no competing interests.

Data Sharing Statement: The raw data are available upon reasonable request to the corresponding author via email.

Generative AI Declaration: During the preparation of this manuscript, the authors used ChatGPT to assist with proofreading. All content was subsequently reviewed and edited by the authors, who assume full responsibility for the accuracy and integrity of the published work.

Authors' contributions: Nai-Bo Hu was responsible for study conceptualization, investigation, preparation of the original draft, project administration, and approved the final manuscript.

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Table 1. Comparative Classification of Selected Molecular Abnormalities in WHO-HAEM5 and ICC Frameworks

Molecular Abnormality	WHO-HAEM5 (2022): Classification and Principles	ICC (2022): Classification and Principles	Shared Clinical Implications
NPM1 mutation	AML with NPM1 mutation; classified as AML regardless of blast percentage.	AML with NPM1 mutation; classified as AML regardless of blast percentage.	Strong leukemogenic driver; mandates AML-directed therapy and is generally sensitive to intensive chemotherapy.
MECOM rearrangement	AML with MECOM rearrangement; classified as AML regardless of blast percentage.	Classification based on blast percentage: <10% blasts: MDS with MECOM rearrangement; ≥10% blasts: AML with MECOM rearrangement	Ultra-high-risk feature; conventional therapies are largely ineffective, necessitating intensive induction and early consideration of allo-HSCT.
NUP98 rearrangement	AML with NUP98 rearrangement; recognized as a distinct entity even with <20% blasts.	Classified as AML with other rare recurrent translocations, including AML with NUP98::NSD1.	Adverse prognostic marker; associated with chemoresistance and poor outcomes in both adult and pediatric AML.

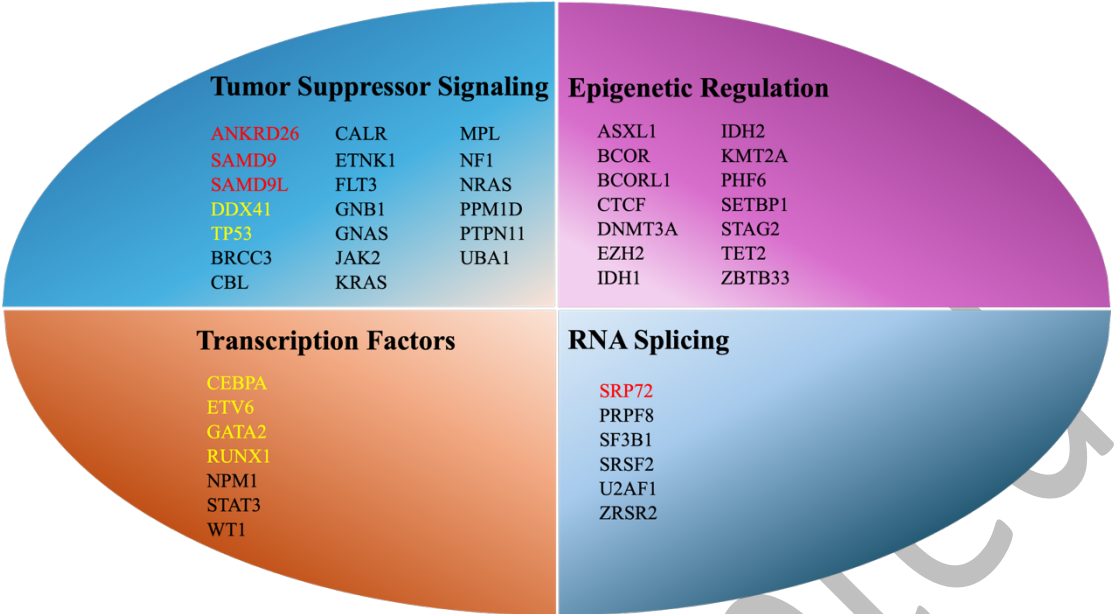
Table 2. Key Molecular Determinants Included or Omitted in IPSS-M Risk Stratification

Gene	IPSS-M Status	Key Function	Clinical Significance in MDS	Potential Influence
TP53	Included	Tumor suppression; DNA damage response	Strongest adverse prognostic marker; biallelic inactivation linked to complex karyotype, treatment resistance, and poor survival	Adequately incorporated
ASXL1	Included	Epigenetic regulation	Consistently adverse; associated with advanced disease and leukemic transformation	Adequately captured
SF3B1	Included	RNA splicing	Often favorable, but effect is highly co-mutation-dependent	Contextual risk not fully integrated
SRSF2, U2AF1	Included	RNA splicing	Associated with poor prognosis and AML progression	Appropriately included
RUNX1	Included	Transcriptional regulation	Adverse; linked to differentiation arrest and genomic instability	Adequately incorporated
DNMT3A, TET2, EZH2	Included	Epigenetic regulation	Heterogeneous effects; DNMT3A/EZH2 adverse, biallelic TET2 potentially indolent	Partially captures complexity
IDH1, IDH2	Included	Metabolic–epigenetic regulation	Actionable targets; inform use of IDH inhibitors	Correctly included
MECOM (EVII)	Not included	Transcriptional regulation	Ultra–high-risk marker; chemoresistance and poor survival	May delay intensive therapy
ZRSR2	Not included	RNA splicing	Often favorable; male predominance	Impairs the accurate diagnosis of lower-risk MDS
DDX41	Not included	RNA helicase; immune signaling	Common germline predisposition gene; implications for donor selection	Inherited risk not captured
SAMD9/SAMD9L	Not included	Innate immunity; growth regulation	Germline predisposition; linked to monosomy 7 and clonal evolution	Underestimates inherited susceptibility
PIGA	Not included	GPI-anchor biosynthesis	Associated with PNH-like clones; affects supportive care	Supportive implications overlooked
BCR::ABL1	Not included	Tyrosine kinase signaling	Suggests alternative diagnoses; mandates TKI therapy	Diagnostic pathway may be missed

Table 3. Selected Emerging and Investigational Therapies in MDS

Regimen	Mechanism (with HMA)	Population	Key Data	Current Status
HMA + Venetoclax	BCL-2 inhibition	Higher-risk MDS	VERONA: no OS benefit; real-world ORR ~62–80%	Cytoreductive/bridging therapy
Azacitidine + Ivosidenib	IDH1 inhibition	IDH1-mutant MDS	Approved post-HMA; phase III frontline trial ongoing	Targeted therapy under evaluation
Azacitidine + Enasidenib	IDH2 inhibition	IDH2-mutant MDS	Early-phase activity	Investigational
Lisaftoclax + Azacitidine	Next-generation of BCL-2 inhibition	Int/high-risk MDS/AML	Phase I/II: safe, active	Promising investigational regimen
APR-246 + Azacitidine	p53 reactivation	TP53-mutant MDS	Phase III negative	Strategy under revision
Sabatolimab + HMA	TIM-3 blockade	Higher-risk MDS	Phase Ib ORR 62.9%; phase III pending	Immune-based approach
Bexmarilimab + Azacitidine	Clever-1 macrophage targeting	HMA-refractory MDS	Phase I/II ORR ~45%	Early-phase immunotherapy
H3B-8800	Spliceosome modulation	Spliceosome-mutant MDS	Limited efficacy	Discontinued
Magrolimab + Azacitidine	CD47 blockade	Higher-risk MDS/AML	Toxicity/limited efficacy	First-gen CD47 agent

Figure 1. Key Mutational Pathways in MDS



In mutation annotation, black usually indicates somatic mutations , while red clearly denotes germline mutations(requiring further confirmation with VAF); yellow may represent either somatic or germline mutations and requires comprehensive evaluation based on the specific locus and VAF.