Review Article

Exploring the acute myeloid leukaemias

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ABSTRACT

Abstract: The acute myeloid leukaemias are genetically a diverse group of neoplasm with varied clinical behavior and response to treatment. Advances in immunophenotyping, cytogenetics and molecular genetics have resulted in better understanding of their genesis. Risk stratification of different variants is now emerging. Therapy strategies are now increasingly being developed considering the inherent biological behavior of the different subtypes. It is anticipated that in the future, deeper secrets of these once fatal diseases will be unraveled by advances in newer genomic techniques. It is hoped that future use of gene specific tailored therapy and strategies will result in longer survival in cases showing poorer prognosis at present.

History and classifications

The classification and genetics of acute myeloid leukaemias (AML) have undergone major changes since the 1970’s as rapid advances in immunology and genetics have made significant progress in the understanding of this varied group of disorders (fig. 1). The current 2008 WHO classifies precursor myeloid neoplasm into 7 major types and 24 subtypes’ entities (Table 1).

The FAB; 1976 was a landmark classification in its days. It was essentially a morphological classification. Initially it was very useful as it standardized nomenclature used by different authors all over the world. Immunophenotyping was introduced in revised FAB Classification in 1982 to differentiate AML Mo from ALL and to diagnose M7. Cytogenetics and molecular genetics were not used. It did not consider MDS associated AML or prognosis (risk groups categorization).

The WHO (2001) classification was the second landmark classification which incorporated morphology, cytogenetics, molecular genetics and immunological markers of AML. It was an acceptable classification having prognostic value. It separated favorable leukaemias with balanced translocations which respond better to treatment from the unfavorable ones. The unfavorable group included AML with:

- Complex karyotypes
- Partial or complete loss or deletions of chromosomes 5 and/or 7
- Multilineage dysplasia and
- Multi drug resistance.

Keywords: Acute myeloid leukaemia; Immunophenotyping; Cytogenetics; Molecular genetics; Gene profiling

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The WHO (2008) classification basically continues to use the same approach used as in WHO 2001 classification, but added other parameters. It starts with analyzing basic morphology and also incorporates clinical, morphology, immunophenotyping, cytogenetics and molecular genetics for diagnosis and classification. As a result the initial morphological diagnosis may sometimes have to be amended after further investigations parameter reports become available (fig.2).

Immunophenotyping and karyotyping:

Immunophenotyping is rapidly replacing cytochemistry to differentiate acute leukaemias. They are useful to confirm myeloid or mixed phenotypic acute leukaemia lineage (MPAL; fig.3). Immune markers are also useful to separate minimally differentiated acute myeloid leukaemia from acute lymphoblastic leukaemia and recognition of acute megakaryoblastic leukaemia. Strict criteria for immunophenotyping diagnosis need to be followed when investigating suspected MPAL as over use of immunophenotyping can lead to an over diagnosis of MPAL. About 50-60 % of de novo AML show karyotype abnormalities. By karyotyping AML are classified into low, intermediate and high risk groups (Table 2). Many AML cases have however normal karyotypes can still have genetic mutations. MDS related AML are now better understood due to karyotyping.

AML with recurrent genetic abnormalities

They have been separated as a distinct entity as they are associated with characteristic clinical, morphological and immunophenotypic features that form a clinicopathologic-genetic entity having prognostic significance. The most frequent are t(8;21), inv(16) or t(16;16), t(15;17) and t(9;11). Distinct morphological changes may be seen with some of the translocations. Translocation t (8:21) (q22;q12) AML cases may show blasts with perinuclear hofs and chunky pink cytoplasmic granules which can be misinterpreted as granulocytes (fig.5). Inv (16) (p13:1q22) and t (16;16) (p13.1:q22) cases may show abnormal eosinophils with large basophilic granules.

Mutation analysis in AML

There are two types of genetic mutations in AML. Class I mutations provide a proliferative and/or survival advantage and do not affect differentiation. They occur late in the disease or in relapse. They include: KIT, FLT3-ITD and FLT3-TKD. Class II mutations impair haematopoietic differentiation and apoptosis. They occur early in the disease. They include: NPM1 and CEBPA (fig.5).

Correlation of some mutations in AML with prognosis

NPM1 (nucleophosmin) mutations are the most common mutations in AML. They occur in 30% of de novo AML and 50-60% of AML with normal cytogenetics. An NPM1 mutation without any FLT3-ITD mutation is associated with good induction therapy and better prognosis. CEBPA (CCAAT binding protein enhancer alpha) mutations are seen in about 5-9% of AML. Lower relapse rate and improved survival are seen in CEBPA gene mutation. FLT3-ITD (FMS related tyrosine kinase 3 with internal tandem duplications) mutations are associated with shorter survival and unfavourable prognosis. KIT mutations associated with t (8; 21), inv (16), t (16; 16) are associated with poorer prognosis. Some of the other mutations and their present clinical utility are listed in table 4.
Acute myeloid leukemia

AML with myelodysplasia related changes

These myeloid neoplasms include AML showing dysplasia: in more than 50% of cells of 2 or more myeloid lines in a pretreatment specimen. They can be seen de novo or following MDS or MDS/MPD. The cases are seen mainly in elderly and rare in children. There is often severe pancytopenia. They can show dysgranulopoiesis, dyserythropoiesis, and dysmegakaryopoiesis. The blasts are positive for CD34 and pan-myeloid markers (CD13 and CD33). Frequently aberrant expression of CD56 and CD7 is seen. There is increased incidence of multidrug resistance glycoprotein: MDR-1. Multilineage dysplasia has adverse effect on remission.16

Therapy related myeloid neoplasm

They are seen after cytotoxic chemotherapy and or radiotherapy. They are of two types: alkylating agent and radiation therapy related and Topoisomerase II inhibitor related.

Alkylating Agent/Radiation Therapy Related AML/ MDS

They occur 5-6 years after initiation of treatment with alkylating agents or radiotherapy. The risks are related to age and also to cumulative dosage. Two-thirds of cases present as MDS. One-third of cases present as overt AML with MDS features. All myeloid cell lines affected.

Dyserythropoiesis, ring sideroblasts are seen in 60% of cases (one-third in excess of 15% of erythroid cells). They also show hypo granulation and nuclear hypolobation of granulocytes. Dysplastic megakaryocytes are seen increased in 25% of cases. Basophils are increased in 25% of cases. Auer rods are seen in a minority of cases. Bone marrow biopsy is hypercellular in 50% of cases, normocellular in 25%, hypopcellular in 25%, and shows fibrosis in 15%. The types of AML commonly developed include M2, M4, M5, M6, and M7.

Table 4: Some gene mutations in AML and their significance

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<th>Established</th>
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<th>Potential</th>
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<tr>
<td>Favourable mutations</td>
<td>NPM1, CEPBA</td>
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<tr>
<td>Unfavourable mutations</td>
<td>FLT3-ITD, KIT</td>
<td>MLL-PTD, WT1, FLT3-TDK</td>
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<tr>
<td>Unfavourable over expression</td>
<td>BAACL, EVL, ERG, HOXA9, MN1, MEIS1</td>
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<td>Clinical testing</td>
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Immunophenotype in these lesions include blasts that often express CD34+, CD33+, CD13+, frequent aberrant expression of CD56 and CD7+. There is an increase incidence of MDR-1 expression in blasts. These lesions show increased cytogenetic abnormalities similar to de novo MDS, RCMD, RAEB. These include unbalanced translocations, deletions of chromosomes 5 and 7 (long arms), chromosomes 1, 4, 12, 14, 18, complex chromosomal abnormalities. They show poor response to therapy, poor survival is seen.17

Topoisomerase II Inhibitor Related AML/MDS

They are associated with epipodophyllotoxins and related compounds that target DNA-Topoisomerase II, examples etoposide and teniposide, also anthracyclines, such as doxorubicin and 4-epi-doxorubicin. They affect all age groups. They show shorter latency: 12-130 months (median: 33-34 months). Latency can be less than 6 months. They usually presents as overt AML without a previous MDS.
phase. There may be a significant monocytic component. Most are acute monoblastic or myelomonocytic, occasionally acute promyelocytic leukemia or acute megakaryoblastic leukemia. The bone marrow is usually hypercellular. Usually balanced translocations involving 11q23 (MLL gene) and primarily t (9;11), t (11;19), and t (6;11) are seen. Other translocations include t (8; 21), t (3; 21), inv (16), t (8;16). They usually show good initial response to therapy, but relapses frequent and survival variable (especially poor with 11q23).

**AML, not otherwise specified (AML NOS):**

They include AML without recurrent genetic abnormalities and those with no myelodysplasia, post therapy related or Down's syndrome association (fig.6 and table6). Promonocytes in AML monoblastic variant are considered equivalent to blasts. Two new entities acute basophilic leukaemia and acute panmyelosis with myelofibrosis have been included with AML NOS.

**Myeloid proliferations associated with Down’s syndrome**

Transient myeloid abnormalities often occur with DS, though AML related with DS is also seen. About 10% of Down’s syndrome show transient myeloproliferative disorder (TPD). The blasts are CD13, CD33 (myeloid) and CD41, CD61 (megakaryocytic) positive. The cases can be asymptomatic or show severe disease. Most regress spontaneously without any treatment. About 20-30% progress to AML.

Down’s syndrome (DS) is associated with acute leukaemia (fig.7). AML M7 is 500 times more common in DS than normal children. Half of leukaemias in DS are AML. ALL is 40 times more common in DS. AMKL is more common and confers a better prognosis in Down’s syndrome. GATA-1 mutation occurs early in DS and is associated with increased risk of future leukaemia. GATA-1 mutations in DS with AML show better response to therapy and better overall prognosis compared to non-DS children with AML.18

**Microarray analysis and specific targeted therapy for AML**

Microarray analysis or gene profiling involve using large numbers of up to 13,000 oligonucleotide gene probes. The aim is an attempt to try to study the pathogenesis, predict prognosis and sub classify the different AML variants into meaningful groups. Genes with poor prognosis have been identified like HOXB2, HOXB5, PBX3, HOXA4, and HOXA10. However microarray methods are still yet to replace standard genetic methods.19

Use of specific agents targeted against gain of function mutations like FLT3, JAK2 and cKIT are being developed using tyrosine kinase inhibitors The drugs include sorafenib, quizartinib, testauctinib, midostaurin, pacritinib, PLX3397,
and CCT137690. Aberrant proteins that are produced in AML like PLK1 can also be a target for specific therapy.

CONCLUSION

The last decade has seen a large upsurge of interest and has resulted in many discoveries in the fields of acute myeloid leukaemias, cytogenetics and molecular genetics. A molecular classification of leukaemias is a distinct reality appearing in the distant horizon. Gene expression profiling, micro-RNA analysis, methylation analysis and whole genome sequencing analysis are expected to deliver more accurate descriptions of intricate molecular changes in AML. The data will help better understand the clinical behavior in distinct genetic groups resulting in better response to therapy and improved prognosis. The fields are exciting, it is hoped that all these events will lead to better understanding and management of these neoplasm, many of which still show dismal prognosis at present.

REFERENCES