Atrial fibrillation as initial presentation of myelodysplastic syndrome (Case report)

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Abstract: We present a case of a patient with myelodysplastic syndrome where in imprecieation persists for case analysis . The cytogenetic study revealed Philadelphia negative with multiple breakpoints and isolated monosomy of chromosome 5 revealing the likely consideration for myelodysplastic myeloproliferative syndrome terminating to acute myelogenous leukaemia (unclassified RAEB category).There is absolute paucity of presentation in the world literature of cases of myelodysplastic syndrome having atrial fibrillation as the initial presentation.

I. Introduction :
Cardiac involvement with leukaemic blast is a well known finding in patients with acute leukaemia 1,2. However involvement of heart with conduction defects in myelodysplasia is reported rarely in the literature 3,4. The symptoms may be ascribed to iron overload & side effects to therapy which are not the reasons in the present case as our patients initial presentation to hospital was with arrhythmias ( Atrial Fibrillation ).

II. History :
Mr KMR , 65 YRS old male was hospitalized with giddiness, palpitations, chest discomfort, weakness, dry cough, shortness of breath, easy fatigability and lethargy which he was experiencing past 1 month prior to the hospitalization . On examination he was afebrile, Heart rate ; 150/min, irregularly irregular, ECG done suggestive of Atrial Fibrillation, Blood Pressure ; 120/80 mm Hg , Respiratory Rate ; 24/min. There was no cyanosis/jaundice. On routine blood count ; Hb-8.5gm% ,TLC-39400/cumm, platelet-1,44,000/cumm. Peripheral smear revealed 5% myelocyte, 5% metamyelocyte, 15% band cells,30% neutrophils, 25% lymphocyte , 2% monocyte , 3% eosinophils , 0% blast cells. A diagnosis of chronic myeloproliferative disorder was considered. Bone marrow biopsy was undertaken along with karyotyping on the 2nd day of hospitalization. His symptoms were attributed to anaemia & rhythm disorder. He was provided with 1 unit of packed red cells & Cardarone & Digoxin for his rhythm problem . 5days later he developed mild febrile illness and breathlessness . The CBC at this time demonstrated Hb- 9.5gm% ,TLC-76,800/cumm, platelet-1,19,000/cumm. Peripheral smear revealed 15% myelocyte, 20% metamyelocyte, 20% band cells,35% neutrophils, 10% lymphocyte , 0% monocyte , 0% eosinophils , 0% blast cells. On 7th day of hospitalization , counts further increased to 99,790/cumm, Hb- 9.9gm% , platelets reduced to 41,000/cumm. Peripheral smear revealed 12% myelocyte, 16% metamyelocyte, 29% Promyelocyte, 17% band cells, 15% neutrophils, 0% lymphocyte , 0% monocyte , 2% eosinophils , 12% blast cells . The clino-hematological diagnosis was consistent with myelodysplastic proliferative disorder terminating to myeloid leukaemia. Bone marrow biopsy was suggestive of MDS & the karyotype revealed Monosomy of chromosome 5 and 12 along with chromosomal breakpoints at 7q11.2,8q22,21q22. Philadelphia was negative. Echocardiographic study identified 55% ejection fraction with mild pericardial effusion and heart rate irregularity. Treatment was instituted with Hydroxyurea & supportive therapy. His condition grew worse on day 10 and developed further cardiac blocks & hypotension with systolic blood pressure of 60 mm of Hg for which pacemaker was inserted & vasopressor therapy was instituted . His vitals remained stable for couple of hours before developing cardiac arrest from which he couldn’t be revived. PostMortem study was not available.

III. Discussion
Myelodysplastic syndrome are all disorders of stem cells in the bone marrow where in haematopoiesis is ineffective of the myeloid class of blood cells . In majority of cases, patient develops cytopenias because of progressive bone marrow failure and 1/3rd cases with Myelodysplastic syndrome get transformed to acute myelogenous leukaemia , some have fatal outcome in few months to years .

World health organization has evolved new classification scheme (2008) based on genetic findings 5,6 However haematocytological screening tests are still the effective tools for deciding the nature of pathology.
Cytogenetic analysis help to evaluate impact on disease outcome of Myelodysplastic syndrome. The variables included age, gender, besides cytogenetic analysis. Cytogenetically, poor outcomes were complex & are described as less than or equal to three abnormalities, &/or presence or chromosome 7 anomalies. Good outcomes noted with deletion 5q alone, Y alone, deletion 20q alone. The present case had revealed 7a11.2, 8q22, 2q22 including chromosome 7 anomaly, thus suggesting the poor outcome as also noted in our case. The patient succumbed to death within a short period of 2 weeks. The variables described by their statistical power separated such patients into distinctive subgroup of risk. High risk (8%) describes for acute myeloid leukaemia developing in 0.2 years reported by international MDS risk analysis workshop.6 The dyslastic syndrome under new WHO system is dealt as under.5

1) RARS- Refractory anemia with ring sideroblast with refractory cytopenia, neutropenia, anemia, thrombocytopenia
2) RABSS- Refractory anemia with ring sideroblast with thrombocytosis which is essence of myelodysplastic myeloproliferative disorder.
3) RAEB- Refractory anemia with excess blast.
RAEB-1 5 to 9% blast
RAEB-2 10 to 19% blast
Auer bodies may be seen in RAEB 2 which has a poor prognosis.
4) The category of RAEB-T. Refractory anemia with excess blast. In transformation is eliminated and considered to have acute leukaemia with deletion of long arm of chromosome 5.
5) Myelodysplastic myeloproliferative overlap syndrome
   A) 5q syndrome (typically seen in older women with normal or high platelet count)
   B) Refractory cytopenias of childhood.
   C) Myelodysplastic unclassifiable; Where in patients with RA or RAEB occasionally present with leucocytosis or thrombocytosis instead of usual thrombocytopenia as noted in the cases.

Cardiac involvement in patients with myelodysplasia is often attributed to the presence anaemia, iron load and chemotherapy.7,8. The case under discussion had presented with arrhythmia and anaemia of short duration where no chemotherapy agent & or over iron dose were received by the patient. Such myelodysplastic patient usually present with symptoms attributed to cytopenias & the clinical course progressing to bone marrow failures or transformation to leukaemias. We believe that the in conduction deformities in the patient developed due to infiltration of malignant haematopoietic precursors with myelodysplasia. Similar type of case was reported by Mateen et al as the first case of affection of heart with myelodysplastic syndrome. Leukemic/myelodysplastic cardiac involvement in life is usually not suspected because of subclinical nature of the symptoms & signs with this pathology.9 However cardiac involvement in leukaemia extramedullary (heart) spread is relatively common ranging 37 to 44 %.2

In our reported case, extramedullary cardiac involvement by dysplastic haematopoietic precursors by itself is considered most reasonable hypothesis as the patient presented initially with atrial fibrillation. Long before the patient heralded transformation to leukaemia, yet another postulation can be entertained as explained by Bin cohette et al in 1998.10 The granulocyte macrophage colony stimulating factor overproduction may leads to autonomous colony formation in the bone marrow of patients with myelodysplastic syndrome which may also partially explain the proliferation of malignant haematopoietic cells elevation in the heart. This extramedullary cardiac spread could have heralded the conducting system leading to the devastating cardiac irregularities in the present case.

References:

5. Wikipedia.Myelodysplastic Syndrome
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Figures:

**Fig: 1** Peripheral smear (H & E stain) x100 showing immature cells - Myeloblast, Promyelocyte, Metamyelocyte & band forms consistent with the myeloblastic disorder.

**Fig 2**: Karyotyping analysis showing 45,XY,-5,t(7;8;21;12)(q11.2;q22;q22;q13),-12,-mar. Analysis revealing a total of 42 autosomes, 2 sex chromosomes & a marker chromosome of unknown origin. There is monosomy of chromosome 5 & 12 with presence of 4 break rearrangements between 7q11.2, 8q22, 21q22 & 12q13 region.