# Anemia 2: Fourth year Medical Students/ 17.11.2020

Abdallah Awidi Abbadi.MD.FRCP Feras M Fararjeh, MD Anemia (2): 17.11.2020

Case 2

65 yr old male had gradual onset of "odd" behavior with psychotic symptoms, irritability and parasthesia in hands and feet

He was noticed to have imbalanced gait. Examination showed loss of vibration and proprioception in lower limbs

### **Laboratory tests**

Hb **5** g/dl, MCV **112**,

Retics (corrected)0.009

WBC 3.3k, Platelets 112k

LDH 1900. Serum B12: 30 pg/ml.

IF Ab +PCA+

Achlorhydria+

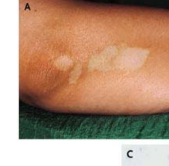
Gastric Bx atrophic gastritis.

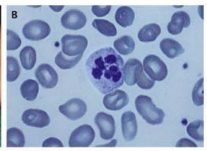
### Physical And Lab

Red Beefy Tongue

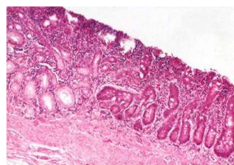
Vitiligo

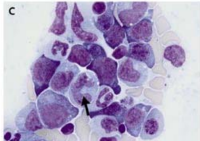






Macroovalocytes.hy perseg



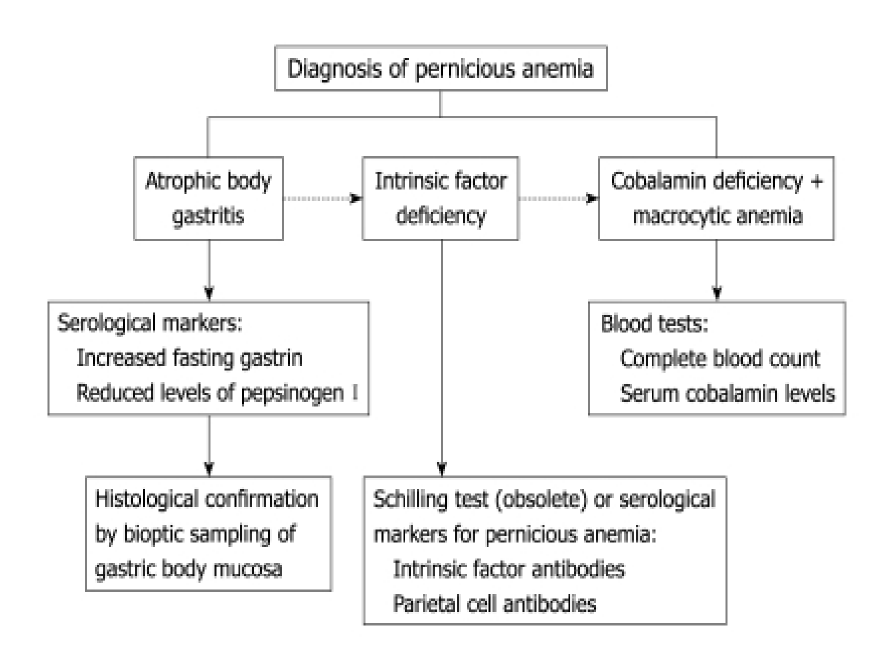


BM: Megaloblasts

Oxyntic G. mucosa atrophy

### Pathogensis of Pernicious Anemia (PA)

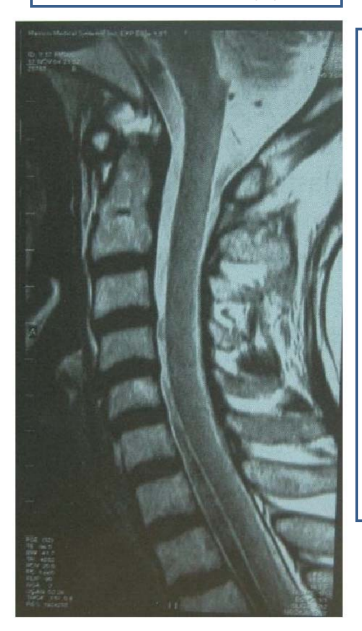
- 1-PA is the end-stage of Atrophic Body Gastritis (ABG) causing oxyntic gastric mucosa damage: achlorhydria.
- 2-It is considered an autoimmune disease (AID).
- 3-AID theory is based on the presence of parietal cell and/or intrinsic factor autoantibodies
  Frequent association with other autoimmune disorders: autoimmune thyroid disease (ATD), type 1 diabetes, and vitiligo



### A-Before therapy



### **B-Post-therapy**



A-Hyperinte nse in cervical region

Bcorrected



**Subacute Combined Degeneration of Spinal Cord** 

Other causes of cobalamin deficiency

**Gastric causes of impaired absorption:** 

Gastrectomy/ gastric sleeve operations

Corpus-predominant *H pylori* gastritis

Long-term proton pump inhibitor therapy

Ileal disease or resection

Blind loop syndrome

Fish tapeworm

Severe pancreatic insufficiency

Decreased intake due to vegetarianism

### Other causes of macrocytic anemia

Folate deficiency

Drugs (e.g. metformin, methotrexate, azathioprine, 6-mercaptopurine)

erythropoiesis: hemolysis, response to hemorrhage)

Liver disease (alcoholic, cirrhosis, poor dietary intake)

Hypoplastic anemia, myelodysplastic syndrome

# Case 2: Treatment & Monitoring

No Blood Transfusion

Vit B12 IM injections daily 7-10 days. Then monthly lifelong.

Careful monitoring of response

Careful monitoring for thyroid function & DM

### Response to Treatment

Reticulocytosis in 3-4days, peak 5-10 days Rise in Hgb concentration within 10 days and normalization in 8-10 weeks as well as correction of MCV.

Fall of serum LDH levels within 2 days
Hypersegmented PMN disappear in 10-14 days
Watch closely for severe hypokalemia during early
response.

Megaloblastic changes disappear within 2 days

### Case 2 B

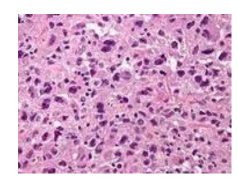
65 yr old male had "anemia syndrome" over the last 6 weeks. He noticed abdominal swelling and weight loss. He had mild fever and night sweats for 2 weeks. No neurological symptoms or signs.

Hb 9, MCV 106, WBC 5.3, Plt 142, Retics (corrected)0.1%. Serum B12 normal. LDH 1100. serum folate was 0.2

**Abdominal Ct** 







### Causes of Folic acid deficiency

#### 1. Inadequate intake

- diet lacking fresh, uncook food; chronic alcoholism, total parenteral nutrition,

### 2. Malabsorption

- small bowel disease (sprue, celiac disease,)
- alcoholism

#### 3. Increased requirements:

- pregnancy and lactation
- infancy
- chronic hemolysis
- malignancy
- hemodialysis

#### 4. Defective utilisation

Drugs:folate antagonists(methotrexate, trimethoprim, triamteren), purine analogs (azathioprine), primidine analogs (zidovudine), RNA reductase inhibitor (hydroxyurea), miscellaneous (phenytoin,  $N_2$ )

# Case 2 B: Treatment and follow-up

Treat the original Cause

Oral administration of folic 5 mg x2daily, for

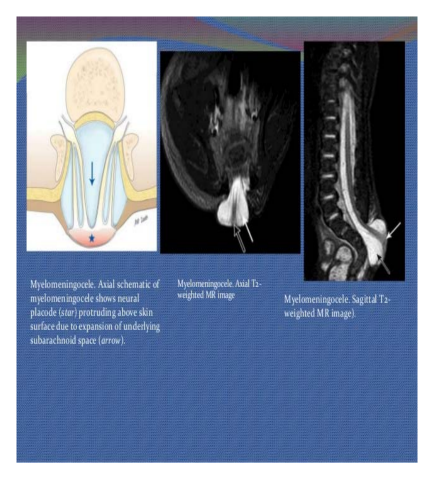
3 months, and maintenance therapy if it is necessary.

Retics after 5-7 days.

Correction of anaemia after 2 months therapy.

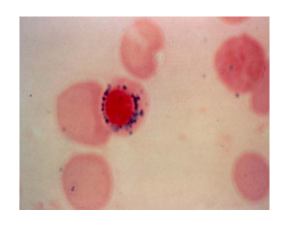
Folic acid has role in neural tube closure in foetus, a pregnant woman should have enough folate to protect her foetus from having neural tube defects

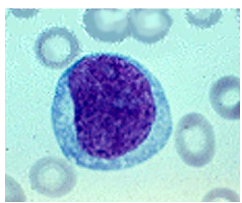




### Case 2 C

48 yr old lady presented with "anemia syndrome" for 3 months. She was found to have splenomegaly. Hb 8g, MCV 107fl, WBC 3.6, plt 95k, retics 0.6%.LDH350 BM: ringed sideroblasts, blasts 8%. Cytogenetics by FISH 11 q del.





Diagnosis: MDS: RARS/RAEB type I with ring sideroblasts

### What are MDS?

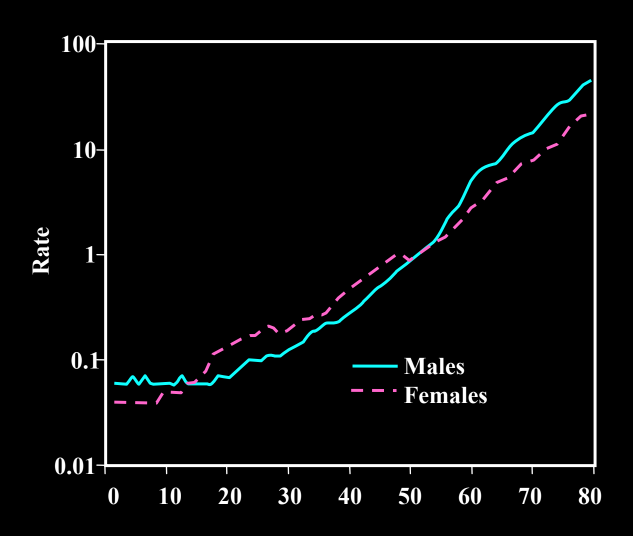
- MDS: a spectrum of heterogeneous malignant hematopoietic stem cell disorders characterized by ineffective and dysplastic changes in BM with
  - ineffective haemopoiesis- dysmorphic cells in blood
  - Variable cytopenia- frequent progression to aml
- MDS may occur
  - a-de novo: primary MDS b-as a result of haemopoietic stem cell injury: secondary or treatment-related MDS
- MDS is associated with significant morbidity and mortality due to
  - cytopenias
  - impaired quality of life
  - risk of transformation to AML

# Epidemiology of MDS

- Epidemiology of MDS
  - common bone marrow disorder
  - the overall incidence is approximately 5 per 100,000 in the general population
  - peak incidence occurs at 60-90 years of age
    - > 20 per 100,000 at 70 years of age
- Typical MDS patient
  - elderly
  - slight male preponderance
  - approximately 50% have a cytogenetic abnormality

# Age-related Incidence of MDS

Leukaemia Research Fund [1984-1993]



## **Pathogenesis**

Poorly understood
Clonal process, thought to arise from single
hematopoietic progenitor cell that acquired multiple
mutations
Global hypomethylation with concomitant
hypermethylation of gene-promoter regions.
Mutation in genes that encode enzymes, such as TET2,
IDH1, IDH2
As role for immunosuppressive agents, suggest

immune system implicated in myelosuppression

and/or marrow hypocellularity

## Clinical features in MDS

- Anaemia
  - > 80% of patients with MDS are anaemic at diagnosis
  - Granulocytopenia
  - 50–70% of patients
  - predisposition for infections
- Thrombocytopenia in 30% of patients
- In MDS
  - chronically low Hb levels associated with cardiac remodelling and increased incidence of heart failure

# Diagnosing MDS

Cytopenia(s) → suspect MDS



Recommended evaluations

- History and physical examination
- Complete blood, platelets, differential, and reticulocyte count
- Examination of peripheral smear
- Bone marrow aspiration with iron stain + biopsy + cytogenetics

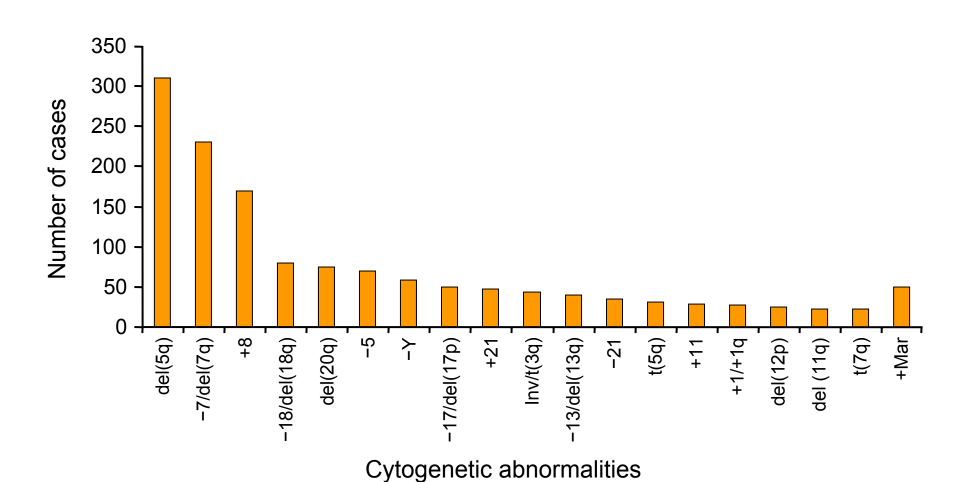
- Serum erythropoietin (prior to RBC transfusion)
- RBC folate and serum vitamin B<sub>12</sub>
- Serum ferritin
- Documentation of transfusion history

Diagnosis of MDS based on morphologic and clinical criteria

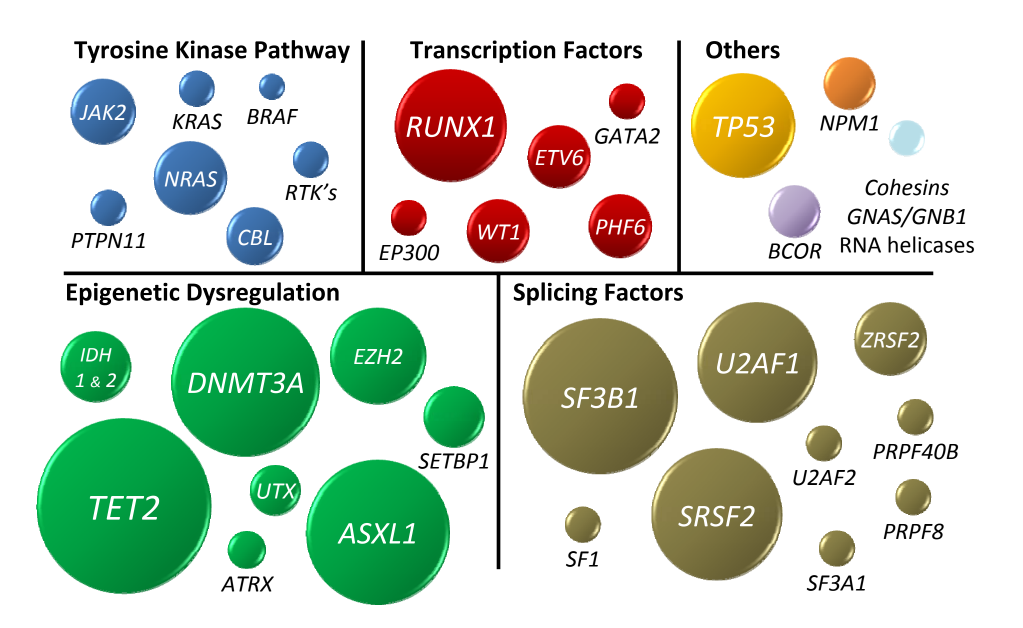
## Subtypes of MDS: WHO classification

Disease	Blood findings	Bone marrow findings
	Anaemia	Erythroid dysplasia only
Refractory anaemia (RA)	No or rare blasts	< 10% grans or megas dysplastic
, ,	< 1 × 10 <sup>9</sup> /L monocytes	< 5% blasts, < 15% ringed sideroblasts
Refractory anaemia with ringed	Anaemia	Erythroid dysplasia only
sideroblasts (RARS)	No blasts	< 10% grans or megas dysplastic
Sidelobiasis (NANO)	NO DIASES	≥ 15% ringed sideroblasts, < 5% blasts
	Cytopenias (bicytopenia or	Dysplasia in ≥ 10% of cells in two or more
Refractory cytopenia with	pancytopenia)	myeloid cell lines
multilineage dysplasia (RCMD)	No or rare blasts	< 5% blasts in marrow, no Auer rods, < 15%
	No Auer rods, < 1 × 109/L monocytes	ringed sideroblasts
Refractory cytopenia with multilineage dysplasia and ringed sideroblasts (RCMD-RS)	Cytopenias (bicytopenia or	Dysplasia in ≥ 10% of cells in two or more
	pancytopenia)	myeloid cell lines
	No or rare blasts	≥ 15% ringed sideroblasts, < 5% blasts, no Aue
	No Auer rods, < 1 × 109/L monocytes	rods
Refractory anaemia with	Cytopenias	Unilineage or multilineage dysplasia
excess blasts-1 (RAEB-1)	< 5% blasts	5–9% blasts, no Auer rods
excess biasis-1 (NAED-1)	No Auer rods, < 1 × 109/L monocytes	5-570 blasts, no Adel Tods
Refractory anaemia with	Cytopenias	Unilineage or multilineage dysplasia
excess blasts-2 (RAEB-2)	5–19% blasts	10–19% blasts, Auer rods ±
excess biasis-2 (NALD-2)	Auer rods ±, < 1 × 10 <sup>9</sup> /L monocytes	10 10 % blasts, Audi 10us ±
Myelodysplastic syndrome,	Cytopenias	Unilineage gran or mega dysplasia
unclassified (MDS-U)	No or rare blasts, no Auer rods	< 5% blasts, no Auer rods
MDS associated with isolated	Anaemia	Normal to increased megakaryocytes with
del(5q)	< 5% blasts	hypolobulated nuclei
ao.(04)	Platelets normal or increased	< 5% blasts, no Auer rods, isolated del(5q)

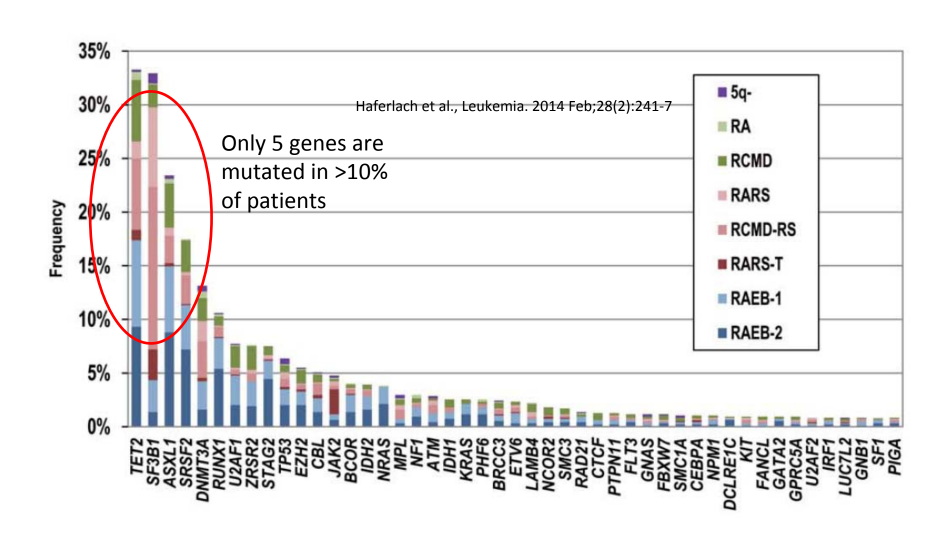
# Frequencies of the most common cytogenetic anomalies in patients with MDS



### **Point Mutations in MDS**



# Many mutations are very rare



# WHO classification-based Prognostic Scoring System (WPSS)

Variable	0	1	2	3
WHO category	RA, RARS, isolation 5q-	RCMD, RCMD-RS	RAEB-1	RAEB-2
Karyotype <sup>*</sup>	Good	Intermediate	Poor	_
Transfusion requirement	No	Regular	_	_

\*Karyotype: **good**: normal, -Y, del(5q), del(20q); **poor**: complex (≥ 3 abnormalities), chr 7 anomalies; and intermediate: other abnormalities.

Score	WPSS subgroup	Median survival (months) Italian cohort	Median survival (months) German cohort
0	Very low	103	141
1	Low	72	66
2	Intermediate	40	48
3–4	High	21	26
5–6	Very high	12	9

Malcovati L, et al. J Clin Oncol. 2007;25:3503-10.

# Case 2 C

### **WPSS**

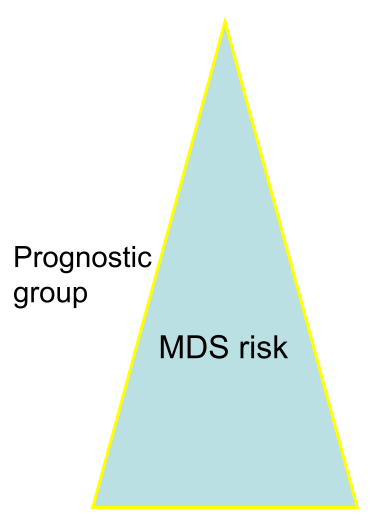
WHO category =2

Cytogenetics intermed. = 1

Bld Trx = 0

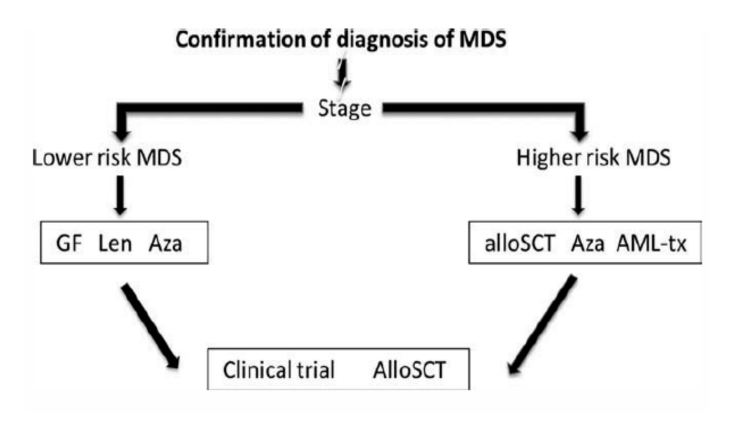
Total score 3. ms 21-26 months

# MDS: therapeutic options



- "Best supportive care", including iron chelation
- Haemopoietic growth factors
- Immunosuppressive treatment
- Differentation induction
- Immunomodulatory drugs
- Arsenic trioxide
- Low-dose chemotherapy
- Epigenetic treatment
- Intensive chemotherapy
- Allogeneic SCT

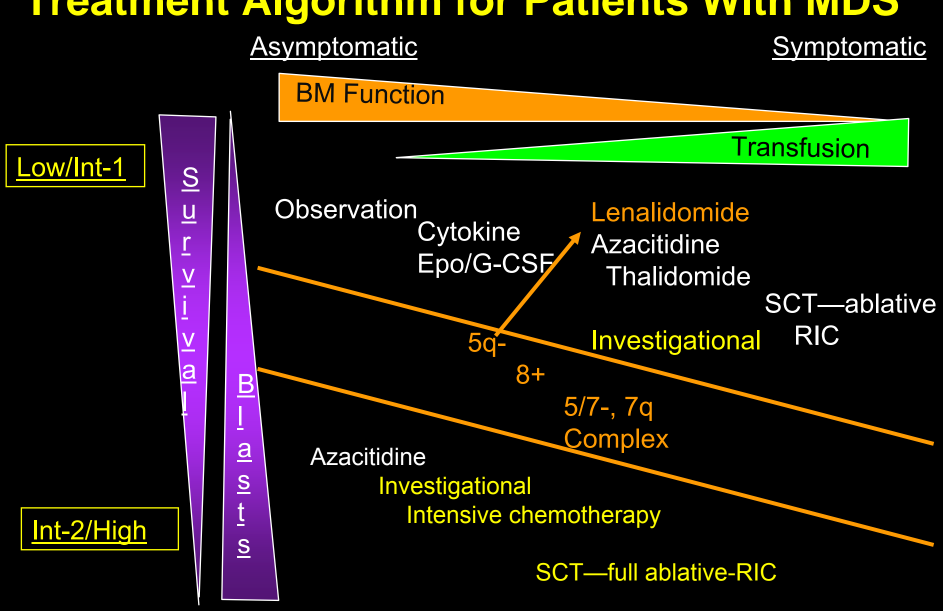
### proposed general treatment algorithm



Treatments can be complicated by advanced age, comorbidities, chronicity of the disease.

Garcia-Manero, G. Am J Hematol. 2012.

## **Treatment Algorithm for Patients With MDS**



RIC = reduced intensity conditioning. From Silverman. In: Holland et al, eds. Cancer Medicine. 7th ed. BC Decker; 2006, .