

Anaemia Cheat Sheet

by shannenp (shannenprice) via cheatography.com/141934/cs/30489/

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What is A reduction the Hb concentration below normal ranges
Anamia (M: <13g/dL / F: <12g/dL)

, triarria (W. 170g/az / 1. 172g/az)

Aetiology 1. Decreased RCB production 2. Loss of RBCs 3.

Increased RBC destruction

Anaemia Main Categories

Microcyctic / hypochomic Low MCV / Low MCH

Normocyctic / normochomic Normal MCV / Normal MCH

Macrocyctic / hyperchomic High MCV / High MCH

Clinical Presentation

fatigue

dyspnoea

chest pain

dizziness

palpitations

headaches

worsening of other conditions - intermittent claudication

Normocytic Anaemia

Normal MCV, indicating normal sized RBCs

DDx: Anaemia of chronic disease/inflamm-

ation

Bone marrow infilt-

ration

Haemolysis

Acute blood loss

Normocytic Anaemia

Investigations	
FBC	Low Hb, normal MCV
Blood Film	Normocytic, normochromic RBCs
Iron Studies	normal/low serum iron, low TIBC, normal/high serum ferritin
+/- other Ix	Serum erythropoietin (EPO) level is decreased in CKD

Management

Manage underlying consult haem/medical team

cause

EPO replacement

Normocytic Anaemia (cont)

RCC transfusion if severe or symptomatic

Iron supplementation may or may not be needed

Normocytic Anaemia Causes

Anaemia of Chronic Diseases	Chronic renal disease, rheumatic disease, congestive heart failure
Mechanism	depends on underlying pathology
	decrease in release of stored iron
	shortened red cell survival
	impaired marrow response in red cell relacement

Macrocytic Anaemia

Macrocytic	Large RBCs and increased MCV
Anaemia	

Aetiology

Megaloblastic	B12 or folate deficiency
Normoblastic	alcohol excess, reticulocytosis, liver disease

Mechanism

Megaloblastic	Impaired	DNA	synthesis
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Normoblastic Unkowr

Signs and General symptoms of anaemia

symptoms

Pallor +/- glossitis, angular stomatitis B12 deficiency can lead to neurologic

syndrome

Megaloblastic Anaemia

Vit B12 Found in animal sources	
Causes of deficiency	Pernicious anaemia (autoimmune disorder)
	Veganism
	Gastrectomy/gastric absorptive disease
	Chron's disease/coeliac



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Megaloblastic Anaemia (cont)		
Folate	green veg,, organ meat, fortified cereals	
Causes of deficiency	Poor dietary intake	
	Alcohol	
	Anti-epileptic drugs (phenytoin)	
	Methotrexate	
	Coeliac disease	

Managemer	nt Macrocytic Anaemia
Treat underlying cause	consult haem/medical team
B12 deficiency	IM hydroxocobalamin (B12): replenish levels with frequent administration then gradually reduce frequency
Folat deficiency	Oral folate replacement: folic acid 5mg OD

Signs - on CE	
Jaundice	can occur in haemolysis
Koilonychia	spoon shaped nails in IDA
conjunctive pallor	ensure looking at palpebral conjunctiva
sclera icterus	jaundice (haemolysis)
angular stomatitis	B12/folate/iron deficiency
systolic flow murmur	mid-systolic ejection murmur due to increased semi-lunar blood flow

Anaemia Differential Diagnosis			
Microcytic Anaemia	Normocytic Anaemia	Macrocytic Anaemia	
Iron deficiency anaemia (50% of cases)	Anaemia of chronic disease	Vitamin B12	
Thallasaemia	Inflammation:		
Chronic diseases	Chronic infection		

Iron Deficiency Anaemia		
4 main causes	Decreased intake (infant/vegan)	
	Decreased absorption (gastrectomy, IBD, coeliac disease)	
	Increased demand (childhood, pregnancy)	
	Increased loss (chronic slow bleed)	
Potential Symptoms	GI blood loss, heavy menstrual bleeding, Pica	

IDA Investigations and Management		
Investigations		
FBC	decreased Hb and MCV. Check WCC & platelets (expect normal in IDA)	
Iron Studies	decreased serum iron, serum ferritin, transferrin sat., increased TIBC	
Blood Film	microcytic and hypochromic RBCs, Poikilocytosis / Anisocytosis	
+/- other Ix	Faecal occult blood (FOB), OGD, colonoscopy	
Management		
Manage underlying cause	consult haem/medical team	
Start supple- mental iron	Aim 1-2g raise in Hb every week	
1st line: oral iron replacement eg. Ferrous fumerate		
2nd line: IV iron replacement (Ferrinject)		
3rd line: RCC transfusion (if severe)		
Don't forget to type and screen if giving a blood transfusion		

Microcytic Anaemia	
Low Hb & MCV, indication RBCs	Causes of microcytic anaemia; mnemonic TAILS
Mechanism:	Defect in synthesis of haem
	Thalassaemia - defect in synthesis of globin chain
DDX:	

- T Thalassemia
- A Anaemia of chronic disease
- I Iron deficiency anaemia
- L lead poisoning
- S sideroblastic anaemia



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Macrocytic Anaemia		
Investigations		
FBC	Low Hb, MCV is elevated	
B12/Folate Deficiency	check levels	
Anti-parietal cell anti-body & intrinsic factor antibody	screening for pernicious anaemia	
Anti-tTG & IgA	screening for coeliac disease	
LFTs	GGT may be elevated in alcohol excess	
Peripheral blood smear	anisocytosis, poikiloctyosis, hypersegmented neutrophils	
Don't forget to ask about diet (vegan), alcohol intake, medications		

IDA Iron Studies				
Serum iron levels	LOW	measures amount of iron in transit in blood		
Serrum ferritin	LOW	total iron stored in the body		
Total iron binding capacity	HIGH	TIBC increases in order to try and maximise use of the little iron available		
Transferrin saturation	LOW	level of saturation of transferring with iron: normal is 30%. Reduced in iron deficiency states		

Autoimmune Haemolytic Anaemia (Haemolysis)		
Warm AIHA	Antibody active at body temp	
	Aetiologies include: rheumatic disease and lymphoproliferative disorders	
	IgG antibodies +/- complement	
Cold AIHA	Antibody active only at lower temps	
	Aetiologies include: infections (eg. mono) and lymphoma	
	IgM antibodies	

Haemolytic Anaemia		
Investigations		
FBC	Low Hb, normal MCV	
Reticulocytes	elevated	
LDH	elevated	
Haptoglonbin	low	
LFT's	unconjugated Bilirubin - elevated	
Direct Antiog- lobin (Coombs) Test	if + then autoimmune haemolysis likely	
Blood Film	look for specific abnormalities	
Management		
Treat underlying cause	consult haem/medical team	
Stabilize pt	consult haem re. need for transfusion	
Warm AIHA	1st line: corticosteroids, 2nd line: Rituximab, Azathioprine, Cyclosporin, 3rd line: splectomy	
Cold AIHA	Avoid cold temps & treat underlying cause +/- immunosuppressant (rituximab)	

Haemolytic Anaemia	
Haenolytic Anaemia	Haemolysis: destruction of red blood cells
Aetiologies	
Autoimmune	Warm, cold, transfusion reaction, drug induces
Haemoglobinopathies	sickle cell, hereditary spherocytosis, thalassaemia
Infections	malaria
Enzyme defects	G6PD
Microangiopathic haemolytic anaemia (MAHA)	haemolytic uremic syndrome, TTP, DIC, eclampsia/HELLP
Mechanical haemolysis	heart valve prosthesis
Rare	Paroxysmal noctural, haemoglobinuria (PNH)



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