DEFINITION

- Abnormal accumulation of fluid in ≥2 compartments of the fetus (pleural and pericardial effusions, ascites and/or subcutaneous oedema)
- Often associated with polycythaemia and placental thickening
- High but variable mortality rates dependent on underlying cause

TYPES

- Traditionally classified into 2 types:
- non-immune hydrops fetalis occurs in the absence of maternal antibodies; accounts for 90% of fetal hydrops in Western countries
- immune hydrops fetalis occurs when maternal allo-immune antibodies are produced against fetal red cells causing haemolysis; rare since introduction of anti-D immunoglobulins

AETIOLOGY

- Imbalance of fluid movement between fetal intravascular and interstitial spaces
- Multiple causes including cardiac abnormalities (structural or arrhythmias), chromosomal/ genetic, infection, haematological, metabolic and non-cardiac structural anomalies
- No identifiable cause found in 15–31% of babies

ANTENATAL MANAGEMENT

- Hydrops fetalis is diagnosed antenatally via ultrasound
- Refer to fetal medicine team [important as confirmed antenatal diagnosis aids appropriate counselling of families, and further intensive monitoring required throughout pregnancy (discussion of this is beyond the scope of this guideline)]
- Possible antenatal interventions include intra-uterine blood transfusion and in-utero procedures e.g. paracentesis/thoracentesis
- High risk of premature delivery

Refer all antenatally diagnosed hydrops fetalis to a regional fetal medicine centre for further assessment and management

NEONATAL MANAGEMENT

Resuscitation

- Resuscitation and stabilisation can be difficult
- An expert team including a neonatal consultant should be present at delivery
- Manage according to Neonatal Life Support (NLS)

Consider concurrent pleural/ascitic drains to facilitate resuscitation

- In cases of severe anaemia, give urgent Group O RhD negative blood transfusions
- Baby may need further grouped and crossmatched blood transfusions in NNU

Give only CMV negative and irradiated blood

Ventilation

- Ensure adequate oxygenation and ventilation
- May require high frequency oscillatory ventilation [see Ventilation: high frequency oscillatory ventilation (HFOV) guideline] and muscle relaxation
- If pulmonary hypertension present may require nitric oxide (see Nitric oxide guideline)

Cardiovascular system

- Use inotropes to support heart and blood pressure
- If intravascular fluid depletion give colloid
- Strict fluid balance
- If severe compromise may require further pleural and ascitic taps
- Immune hydrops may require exchange transfusion. See **Jaundice** and **Exchange transfusion** guidelines

NEONATAL INVESTIGATIONS

- Due to the extensive list of potential causes, direct investigations according to clinical history and presentation
- Initial investigations to consider include:

	Initial investigations	Further investigations to be considered if underlying cause is not ascertained
Haematology	 FBC (including blood film) Group and direct Coombs test Maternal Kleihauer test 	 Red cell enzyme deficiency (e.g. G6PD deficiency) Red cell membrane defects (e.g. hereditary spherocytosis) Haemoglobinopathies (e.g. thalassaemia)
Biochemistry	 Liver function tests including albumin Urea, creatinine and electrolytes 	 If pleural/ascitic tap done – send for fluid MC+S and biochemistry
Cardiac	 ECG to exclude cardiac dysrhythmias Echocardiography to exclude structural heart defects 	
Placenta	 Send to pathologist 	
Genetic testing	ChromosomesMicroarray	 Investigate for congenital metabolic conditions Non-immune hydrops - discuss whole exome sequencing with genetics team
Infection	 Toxoplasma, rubella, CMV, parvovirus, herpes simplex virus 	
Radiology	Chest X-rayAbdominal X-rayCranial ultrasound scan	 Further investigations to be guided by clinical picture

Even with optimal management, the mortality rate is high. Suggest a post-mortem in the event of a death