1.0 Background

By 4–5 months of age, splenomegaly develops in some infants with sickle cell disease, and by 12 months of age a palpable spleen is noted in nearly half. Although enlarged, the spleen does not properly perform its filtration function. However, its reservoir function is overactive: sequestration of large quantities of blood (often half or more of a child's blood volume) can occur rapidly. This complication, termed acute splenic sequestration, is characterized by pooling of large quantities of sickled RBCs in the splenic red pulp, sudden enlargement of the spleen (within hours), and a precipitous decline in haemoglobin (Hb) and platelets, and an increase in reticulocytes.

Presentation is often (60%) associated with episodes of fever, suggesting an underlying viral etiology. Most commonly occurs in infants and young children between 6 months and 5 years of age with sickle cell anaemia. It may also occur in older patients with any sickle cell phenotype with or without chronic splenomegaly. Often there is no obvious triggering event.

2.0 Clinical/Laboratory Features

A child with an acute splenic sequestration presents with symptoms of:
- acute anaemia (pallor, tachycardia, frank cardiovascular collapse);
- splenomegaly/abdominal pain (pain in the left upper quadrant); and
- evidence of an active bone marrow response (increased reticulocytes) plus or minus thrombocytopenia.

Retrospective reviews have shown a first-episode mortality of as high as 14%. On physical examination, patients show signs of anaemia, hypovolemia, and an enlarged spleen (larger than baseline), sometimes massively so. Mild cases may be asymptomatic.

Haemoglobin concentration is at least 20g/L below the baseline steady state. In severe cases, haemoglobin may decline to life-threatening levels. Reticulocyte counts are usually elevated, which distinguishes this condition from aplastic crisis. The platelet count often declines to <50 X 10^9/L.

The mainstay of management is transfusion to provide circulating erythrocytes and volume. Risk of recurrence is approximately 40–50%, usually within 3 years. Because it is not possible to predict which children will have recurrent attacks, most experts recommend splenectomy after the first major attack (for patients >2 years old), or chronic transfusion to maintain a haemoglobin S level under 50% until the patient can get to surgery once all relevant immunizations have been completed.
3.0 Clinical Practice Guideline

Acute Splenic Sequestration: Guidelines for Management in Children with Sickle Cell Disease

Symptoms of Acute Splenic Sequestration:
- Acute chest pain, tachypnea, and hypoxemia (cardiovascular collapse)
- Tachycardia, hypotension, and syncope
- Evidence of an elevated hematocrit response (increased reticulocytosis) and anemia

Gather history and complete physical exam

ED Management
- Complete tests
- Establish IV access
- Ensure child is on inotropically supportive and O2
- If hemoglobin is <8.0 g/dL, below baseline, transfuse as soon as possible with cross-matched PRBC not exceeding 10% of baseline
- Ensure hematocrit is maintained in consultation with Haematology Team
- Use phenotypically matched blood
- If unstable, give IV fluid bolus followed by PRBC
- Initiate Sickle Cell Fever order set in Epic as indicated

Consult Haematology (notify Sickle Cell Team)

Is the child stable?

NO
- Transfer to CCU
- Consult Haematology (notify Sickle Cell Team)

YES
- Admit to Pediatric Medicine

Inpatient Management
- Child must be on cardiac/circulatory monitor
- Monitor vital signs as per SickleCellPRBs
- Repeat physical assessment:
  - Splenomegaly: q4h (measure with tape and record)
  - O2 saturation
- If hemoglobin is <8.0 g/dL, below baseline, transfuse as soon as possible with cross-matched PRBC not exceeding 10% of baseline, and notify hematologist/immediate transfusion volume (Run history of transfusion volume in consultation with Haematology Team, use phenotypically matched blood. If unstable, give IV fluid bolus followed by PRBC and follow SickleCellPRBs recommendations)
- Continue regularly scheduled medications
- Administer O2 to keep SpO2 >95%

Child discharged home from inpatient unit with appropriate follow-up:
- Evidence of rising hemoglobin and diminishing splenomegaly
- Let child be seen in clinic within 2 weeks of discharge
- Continue regularly scheduled medications
- Follow up within 2 weeks of discharge (including blood culture follow-up)

4.0 References


Attachments:

- Revision History.docx
- SC_Clinic Follow Up Revised 2021_FINAL.pdf
- Splenic Sequestration Care Pathway 2021 Final.pdf