Complicated Meconium Ileus: Case Report with Review of Literature

Dr. Anand¹, Dr. Sreeramulu .P.N², Dr. Rahul Singh R³

¹Associate Professor, Department of Pediatric Surgery
²Professor, Department of General Surgery
³Post Graduate in Surgery, SDUMC. Kolar

Abstract: Meconium ileus (MI) is a unique form of neonatal obstruction almost always associated with Cystic Fibrosis. Uncomplicated cases respond well to nonsurgical management. However, few patients develop complications in the postnatal period, requiring emergency operation. A two day old newborn presented with clinical features of low intestinal obstruction suspected to be due to meconium ileus. After resuscitation, a trial of conservative treatment, with Mucomist enemas, was attempted without success. Exploratory laparotomy revealed typical MI with midileal volvulus. Resection of the gangrenous, volvulated ileum with manual evacuation of meconium, both proximally and distally, followed by end to end anastomosis was done. Patient made uneventful recovery. Followup tests did not reveal cystic fibrosis, the growth and development of the baby has been uneventful.

Keywords: Meconium ileus, Volvulus, Gangrene, Cystic Fibrosis

1. Introduction

Meconium ileus (MI) is defined as an obstruction caused by inspissated meconium, at the level of the terminal ileum.¹ It is a unique form of neonatal small bowel obstruction with or without association of cystic fibrosis (CF). Nearly 80% cases of Meconium Ileus is associated with Cystic Fibrosis. Only around 10-20% of cases have been reported to have isolated meconium ileus without any features of CF. However MI complicated by perforation, Atresia, volvulus, gangrene etc., of the affected ileum, requires emergency surgical intervention, with majority undergoing a two staged procedure.² The Prognosis of those babies presenting with uncomplicated MI is generally good and a trial of conservative management can be given.

2. Case Report

A two day old, term male baby, weighing 2.6kg, presented with clinical features of gross abdominal distension, bilious vomiting and non-passage of meconium, of 1 day duration. General physical examination, revealed a lethargic and ill looking baby with gross abdominal distension & moderate respiratory distress. There was no erythema or edema of the abdominal wall. A vague, ill-defined mass was felt in mid abdomen. Bowel sounds were present. He was febrile with temperature of 99 °F; respiratory rate 48/pm, and pulse 143bpm.

Figure 1: Clinical Photograph Showing Distended Abdomen

Figure 2: Contrast Enema showing Micro Colon showing gaseous distension
Meconium ileus is a condition where in extremely viscid, protein rich, inspissated meconium, causing intraluminal obturator type of obstruction of distal ileum by thickly passed mucus plugs that resemble rabbit pellets. Nearly 80% cases of Meconium Ileus is associated with Cystic Fibrosis. More common in the Caucasians with equal incidence in both sexes. Only around 10-20% of cases have been reported to have Isolated meconium ileus without any features of Cystic Fibrosis. Our case was among those 20% which is unusual with no signs of prematurity, no family history. Other causes of Meconium Ileus have also been studied like- Pancreatic aplasia, colon aganglionosis, Pancreatic ductal stenosis, ileocaecal atresia, familial conditions, prematurity.

Two types of Meconium Ileus are described Simple (uncomplicated)/Complicated.

Simple (uncomplicated) MI: baby at birth is normal, no abdominal distension but as time goes by develops abdominal distension, bilious vomiting, careful palpation of abdomen reveals a “doughy” abdomen with indentable masses (meconium clogged bowel).

Complicated Meconium Ileus: Baby is born with significant abdominal distension, respiratory distress with poor general condition. Examination reveals signs of meconium peritonitis/obstruction, ascites. Rectal wash with saline will expel a small amount of thick, viscid meconium. A mass palpable suggesting pseudo cyst, probably due to inueter bowel perforation. The common complications that are encountered are ileal atresia or stenosis, ileal perforation resulting in meconium peritonitis, volvulus with or without pseudo cyst formation.

On a plain radiograph, simple meconium ileus shows unevenly dilated bowel loops with or without air fluid levels due to increased viscosity of the meconium not allowing air interface with the fluid. Occasionally, has a mottled appearance on radiographs during the first 2 days of life, “bubbly” appearance of the distended intestinal loops (Neuhaussser’s Sign) particularly in the right lower quadrant. Contrast enema will show a micro colon (micro colon of meconium clogged bowel). Undue precautions should be taken to resuscitate the sick babies prior to subjecting them to any contrast studies. In a complicated meconium ileus radiographs will reveal signs of obstruction/ giant cysts, ascites and/or calcifications.

Prenatal ultrasound can help in the earlier diagnosis so that suitable precautions can be taken postnatally, findings include echogenic bowel, which can be dilated and thick.
walled, polyhydramnios, fetal ascites, peritoneal wall calcifications, intra-abdominal cysts. Postnatal findings include distended bowels with meconium (in simple MI), ascites, and giant intra-abdominal cysts (in complicated MI).

Meconium ileus is often the first sign of Cystic Fibrosis. The basic genetic abnormality in Cystic Fibrosis is mutation in CFTR (cystic fibrosis transmembrane regulator) gene located on 7q31 chromosome, F508 mutation, causing 3 base pair deletion where phenylalanine residue is removed at 508 position. Abnormal CFTR gene produces abnormal electrolyte content in the environment external to intestinal apical cells, which leads to reduced clearance of secretions from tubular structures lined by the affected epithelium. Chronic intestinal obstruction, abnormal respiratory tract, exocrine pancreas insufficiency, elevated sweat chloride levels are the features.

Meconium ileus (simple) should be differentiated from meconium plug syndrome, Hirschprung’s Disease and bowel atresia.

Majority of these patients are usually sick and dehydrated with electrolyte disturbances. They need intensive resuscitation with IV fluids and antibiotics before subjecting them to invasive procedures or surgery. In simple MI nearly 68% of cases can be managed conservatively with enemas using Omnipaque, gastrograffin, under fluoroscopic control warm saline enemas containing 1% N-acetyl Cysteine. Two failed attempts and operative intervention is needed. Surgery involves Enterotomy and evacuation of all the abnormal meconium assisted by local irrigation with Mucomist (1% N Acetyl Cysteine) or Gastrografin (in a fit patient). Other cases are managed by staged procedures like Mckulikz, Bishop Koop, and Santulli Blanc. Complicated MI are managed by laparotomy, resection followed by ileostomy. Intestinal continuity is established 6-8 weeks post operatively.

Prognosis is the best in isolated, uncomplicated cases without Cystic fibrosis. Patients with CF have chronic problems like repeated chest infections, pancreatic insufficiency with malnourishment, recurrent attacks of constipation and abdominal distension (meconium ileus equivalent), fertility disorders etc.

We present this case because of its rarity and to create an awareness that meconium ileus could be a reason for distal neonatal intestinal obstruction. Instead of a staged procedure this complicated MI was managed by a single staged resection and anastomosis due to reasonably good general condition and absence of overt peritonitis.

4. Conclusion

The presentation of simple meconium ileus with volvulus can be missed if not taken into consideration. Early surgical intervention is required to manage such a situation to avoid complications. Meconium ileus can be a reason for distal neonatal intestinal obstruction.

References