CASE REPORT

Late-presenting congenital diaphragmatic hernia in an infant with tuberous sclerosis – case report and a review of the literature

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ABSTRACT

Tuberous sclerosis complex (TSC) is a rare, genetic syndrome, which is characterized by the occurrence of small, benign multilocated hamartomas. The clinical manifestation of the disease is variable, from mild to life threatening. This report presents a 9-month-old male baby suffering from TSC, which was diagnosed prenatally. The child was under constant medical, multidisciplinary monitoring. This boy presented skin lesions, hamartomas in the brain and heart, and observation toward hamartoma of the right retina. The infant was admitted to the hospital because of vomiting, fever and cough. Chest X-ray showed left diaphragmatic hernia with mediastinum shift. He underwent thoracoscopic hernia repair. The postoperative period was complicated by a left pneumothorax, atelectasis and pneumonia, but finally the child recovered and remains under ambulatory monitoring. Every pathological symptom must be imaged and diagnosed, despite good general condition.

KEY WORDS:
infant, congenital diaphragmatic hernia, tuberous sclerosis complex, radiologic symptoms.

INTRODUCTION

Tuberous sclerosis complex (TSC) is a genetic neurocutaneous syndrome that is characterized by multiple, mostly nonmalignant tumours called hamartomas, in many organs, such as the skin, eyes, brain, liver, kidney, heart and lungs [1]. TSC is a chronic, progressive, autosomal dominant disorder [2], and is caused by a mutation in TSC1 (hamartin) or TSC2 (tuberin) suppressor genes. The incidence rate of this syndrome is estimated at 1/6,000–1/10,000 [3]. The clinical manifestation is variable. Most patients have skin lesions and seizures [4], however symptoms can be from mild to life threatening. We want to present a rare case of a neonate suffering from TSC and late-presenting congenital diaphragmatic hernia, which was successfully treated in our Department.

CASE REPORT

A 10-month-old male baby was admitted to the Emergency Room because of the cough lasting for a week, loss of appetite, and persistent vomiting for the last 2 days. The family physician diagnosed a viral infection the day before admission, but he did not perform additional tests. The child suffered from TSC and thus was under constant multispecialty medical care. This syndrome was recognized prenatally, when MRI imaged small tumours in the brain and heart. Manifestation of TSC in this patient.
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was varied: first of all, neurological: cortical tubers, corti-
cal/subcortical tubers, subependymal nodules in the walls
of ventricles and delayed myelination (images in MRI an
CT); eyes: observation toward hamartoma of the right
retina; heart: small multiple rhabdomyoma in both ven-
tricles (in echosonography) caused asymptomatic mul-
tiple episodes of supraventricular extrasystoles (Holter
monitoring). He was treated with vigabatrin because of
infantile spasm. Family history was burdened with TSC.
Four of his family members suffer from TSC: his mother,
both siblings (10-year-old brother and 2-year-old broth-
er) and maternal grandmother. There was no history of
any trauma or accident.

His vital signs were: temperature 36.5°C, respiratory
rate 25 breaths/min, heart rate 100 beats/min., without
cyanosis, oxygen saturation SpO2 96%. On examination,
we found a weight deficiency (< 3 percentile), apathy
and features of dehydration. There were also four small
hypomelanoic macules in the lower limbs and buttocks.
Chest auscultation revealed decreased breath sound on
the left side. Our attention was also drawn to the scaphoid
abdomen. Because pneumonia was suspected, X-ray was
ordered. On the X-ray image, we saw bowel loops into the
left side of the thorax with dislocation of the mediastinum
to the right side and significantly reduced left lung (Fig. 1).
Echosonography showed small multiple hyperechoic
rhabdomyomas: three in the right ventricle (near the
heart apex), with the largest diameters 12 mm × 7 mm,
and two in the left ventricle (Fig. 2). These masses did
not obstruct blood inflow or outflow. He was admitted
to the Pediatric Surgery and Urology Department and
after fluid resuscitation he was taken to the operating
theatre. Before the operation, we started an intravenous
antibiotic (co-amoxiclav). The patient underwent the
thoracoscopic approach under general anaesthesia and
ventilation of both lungs. A nasogastric tube was insert-
ed, and the patient was placed on the right side with ele-
vation of the left upper limb. The first trocar was placed
under the scapula for 5 mm camera placement. Two more
3 mm trocars were inserted in the fifth intercostal space
in the mid-clavicular and posterior axillary line. The in-
sufflation of carbon dioxide was made with low pressures
6 mm Hg. There was a 4 cm posterolateral defect in the
left side of the diaphragmatic dome with the whole of the
small and large intestines inside the chest. During oper-
ation we did not notice any evidence of a ruptured sac.
According to Congenital Diaphragmatic Hernia Study
Group (CDHSG) registry it was a defects are classified
as “B”. It involved < 50% of the chest wall [5]. The left
lung was collapsed. Hernia content was gently reduced
into the abdominal cavity. The diaphragm was closed with
intra-corporeally interrupted 2/0 nonabsorbable sutures.
The patient was ventilated and sedated for four days.
Post-operative time was complicated by a left pneumo-
thonax (treated by chest tube insertion) and with atelec-
tasis and pneumonia of the left lung. He was treated in

DISCUSSION

The incidence of congenital diaphragmatic hernia
(CDH) is approximately 1/3,000 live births [6]. The aeti-
ology of CDH is still unclear. Genetic causes of CDH have
been noticed in around 30% of patients [7]. Approximate-
ly 40% of CDH cases are associated with other anomalies
[7] like brain, heart, renal and genitourinary disorders. In
more than 50% of cases it could be diagnosed prenatally
[8]. We can postulate that if our patient underwent pre-
natal MRI imaging it would seem plausible that the CDH
defect would have been diagnosed prenatally. A newborn
with CDH often presents severe respiratory distress.
Whereas, in late-presenting CDH, symptoms could be
either acute or chronic. Gastrointestinal symptoms oc-
cur most often, but none of the complains are specific for
late-presenting CDH. In one study, gastrointestinal prob-

FIGURE 1. Left diaphragmatic hernia with mediastinum shift

FIGURE 2. Hamartomas in the heart
lems were seen more commonly in CDH on the left side [9]. In the case of a late diagnosis, hernia is more common on the right side [10], and this is explained by the fact that the liver is a natural, mechanical barrier to dislocation of the intestines to the chest cavity. Late-presenting congenital diaphragmatic hernia accounts for 10–13% all children with CDH [10]. According to the literature, it is more common in males than females [11] and usually it is left side postero-lateral hernia. In most cases, it is an isolated congenital defect. In our case, chest X-ray was enough to make the right diagnosis, but if the image is ambiguous, chest and abdominal CT can help make the right diagnosis.

In the literature, there was only one case of TSC in a neonate presented as diaphragmatic hernia [12] and we can also find mention about two other cases. One it was a 7.5-year-old girl [13], who presented with a diaphragmatic hernia, multiple cardiac rhabdomyomas and a horseshoe kidney with mutation in the TSC2 gene. The second case of TSC connected with CDH was a preterm neonate, in whom diaphragmatic hernia was found during autopsy, in addition to cardiac rhabdomyomas and intestine malrotation. Ohri et al. who described case of tuberous sclerosis in a newborn who had also multiple rhabdomyomas of heart and diaphragmatic hernia, associate the presence of this congenital malformations with multi-germ layer dysplasia [12]. Defects in morphogenesis associated with CDH are heterogeneous, and seemingly related to numerous pathogenetic mechanisms. Most often abnormalities involve heart, brain, genitourinary system, craniofacial region, or limbs [5].

Most patients with the tuberous sclerosis complex suffer from seizures and differential skin lesions [14]. Signs of TSC in the early infant period are usually mild, and the most common initial disorders in small children are cardiac rhabdomyomas and hypomelanotic macules [15]. As in our patient, if the diagnosis of TSC is made prenatally, careful monitoring must be implemented. According to new diagnostic criteria from 2012 [16] pulmonary manifestation, in the form of lymphangioleiomyomatosis (LAM), is one of the major criteria of TSC. LAM is characterized by cysts and excessive proliferation of the smooth muscles, which replace normal pulmonic alveoli, and it is more commonly seen in women [17]. Patients with LAM are at higher risk of spontaneous pneumothorax [18]. Second pulmonary disease is multifocal micronodular pneumocyte hyperplasia (MMPH) [19], which is a rare, benign hamartomatous subtype of type II pneumocyte proliferation [19]. In contrast to LAM, MMPH occurs both in women and in men. Postoperative complications in our patient could be connected with a baseline disorder and strict future observation should be focused on the lungs. On the other hand, it cannot be ruled out that the postoperative pneumothorax was from residual carbon dioxide from thoracoscopy approach. There is still debate about benefits and risks of chest tube placement.

CONCLUSIONS

Clinical manifestation of TSC is heterogeneous and can create diagnostic problems. Patients with tuberous sclerosis complex should be under strict medical monitoring. What is important, every pathological symptom must be imaged and diagnosed, despite good general condition. Diagnosis of congenital diaphragmatic hernia after the neonate period is rare, but possible. Symptoms are non-specific and mainly relate to gastrointestinal or respiratory systems. The surgical correction of CDH may be performed using open or minimal invasive surgery. The approach could be transthoracic or through the abdomen. In late-presenting CDH patient are usually without pulmonary hypoplasia cardio-pulmonary instability, therefore these patients seem to be good candidates for minimal invasive surgery.

DISCLOSURE

The authors declare no conflict of interest.

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