Alveolar Capillary Dysplasia in a Tertiary Center: A Case Report

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ABSTRACT

Background: Alveolar Capillary Dysplasia (ACD) is an exceedingly rare fatal and lethal developmental lung disorder mainly involving the major pulmonary vasculature, with a dismal prognosis. It usually presents in full term infants who develop respiratory distress on their first days of life as persistent pulmonary hypertension of the newborn (PPHN) that is unresponsive to treatment, and produces respiratory failure early in life. The majority of reported cases were found to be associated with other systemic anomalies, more frequently involving gastrointestinal system, as well as cardiovascular, urogenital, musculoskeletal, and right-left laterality anomalies. Since its first description, significant achievements in research have led to better understanding of the underlying molecular mechanism of ACD, and genetic studies have identified association with genomic alteration in the locus of the transcription factor FOXF1.

Objective: Here we present a case of female newborn who was referred to our tertiary center at the age of 5 months due to chronic hypoxia and failure to gain weight. Eventually, she was diagnosed as ACD.

Conclusion: ACD/MPV is a rare and lethal developmental disorder. Patients suffer from severe hypoxemia that progresses over time, although awareness is growing among physicians it can still be confused with idiopathic pulmonary hypertension as the presentation can be similar. This usually delays the diagnosis and leads to unnecessary suffering of patients and waist of hospitals resources. As soon as the diagnosis is suspected. Genetic testing should be done or histological exam should be performed, ideally before ECMO or even surgeries for CO occurring anomalies.

Keywords: Alveolar capillary dysplasia, Respiratory failure, Chronic hypoxia.

INTRODUCTION

Alveolar capillary dysplasia (ACD) is an exceedingly rare pathology involving the major pulmonary vasculature, with a dismal prognosis.

It usually presents in full term infants who develop respiratory distress on their first days of life. Most case reports are associated with other systemic anomalies, more frequently involving gastrointestinal system, as well as cardiovascular, urogenital, and musculoskeletal system (1,2).

We present a case of female newborn who was referred to our tertiary center at the age of 5 months due to chronic hypoxia and failure to gain weight. Eventually, she was diagnosed as ACD.

CASE REPORT

Our patient is a 5 months old baby girl product of full-term pregnancy by spontaneous vaginal delivery (SVD) at another hospital, she had unremarkable antenatal history and normal pregnancy and maternal history, following her birth she was immediately admitted to neonatal intensive care unit (NICU) due to decreased oxygen saturation where she stayed in for 4 days on supportive therapy and did not require ventilation, she was discharged in stable condition by the end of 4th day.

The patient was stable until her mother noticed cyanosis while crying by the age of 2 months, which was described as bluish discoloration of lips, face and extremities when crying, and it would persist for at least 30 minutes after settling. Eventually, her mother opted to have a medical advice, where she was admitted for investigation of hypoxia and failure to gain weight, as she had gained only 0.5 kg since birth, after failing to reach to a diagnosis she was referred to our tertiary hospital to investigate the reason she is failing to thrive as a case of severe pulmonary hypertension on 1L oxygen and incomplete bowel obstruction.
Fig. (2): Normal lung field, dilated bowel in the left side, no pneumothorax.

So, in our hospital X-rays and barium study was done, that showed mild gastroesophageal reflux and evidence of malrotation with right sided location of the proximal small bowel loops. Markedly dilated bowel loop seen in the right upper quadrant most in keeping with a dilated third/fourth part of the duodenum, could be related to proximal jejunal or distal duodenal stenosis vs possibly annular pancreas. Surgical intervention could not be done as the patient went into pulmonary hypertension crisis and was started on nitrous oxide, diuretics, slindafil and bosentan to control her condition.

During that time high resolution CT chest was done that showed bilateral diffuse ground-glass opacification, and a dilated air filled bowl on the right side. By this time, Whole Exome sequence that was sent earlier came to be Positive for Alveolar capillary dysplasia FOXF1 exon 1 (c.841_862del) heterozygous mutation. Echo reported as sever pulmonary hypertension with aneurysmal atrial septum with fenestrated ASD shunting bidirectional mostly right to left with intact ventricular septum. Swallowing assessment was also done that showed aspiration with thin fluid

Fig. (3): Mild gastroesophageal reflux and evidence of malrotation with right sided location of the proximal small bowel loops.

Declaration of patient consent:
An approval of the study was obtained from King Faisal Specialist Hospital and Research Center, Jeddah (Saudi Arabia) Academic and Ethical Committee. The parents of all children participants were informed that the case would be published as case report and this was accepted. This work has been carried out in accordance with The Code of Ethics of the World Medical Association (Declaration of Helsinki) for studies involving humans.

DISCUSSION
Since the first description of ACD in 1981 by Slot et al. 3 as a rare cause of persistent pulmonary hypertension, around 200 cases have been described so far in literature.

It presents as respiratory distress and cyanosis with pulmonary hypertension that is resistant to usual measures within the first day of life, with a mortality rate reaching 100%. Most important differential diagnosis includes pulmonary hypoplasia, idiopathic persistent pulmonary hypertension of the newborn (PPHN), and surfactant deficiency.
Histological features classically showed reduced number of alveolar capillaries, immature alveolar development, and hypertrophic smooth muscle cells in the pulmonary arterioles. Additionally, in normal histology the peripheral veins in the broncho-vascular bundle were positioned within the interlobular septa, while in majority of ACD cases they were located outside the interlobular septa adjacent to pulmonary arteries, which comprises a phenomena called “misalignment of the pulmonary veins” (3).

Currently, there is no specific treatment for ACD, and management is usually aimed to target the PPHN. As with other cases of persistent pulmonary hypertension, the management of ACD is usually supportive, consisting of clinical respiratory and cardiovascular support with the use of vasodilating medications to control the pulmonary hypertension. In majority of cases, immediate intubation and mechanical ventilation are required at the time of presentation due to the significantly low oxygen saturation levels (8).

Extra-Corporeal Membrane Oxygenation (ECMO) therapy may be used to stabilize patients in critical conditions and when diagnosis is not made yet. However, ECMO may lead to some unwanted outcomes, as most patients of ACD are unable to wean off ECMO once placed on it with the addition of its invasiveness and high cost (9).

FOXF1 and ACD/MPV first described in 1994, FOXF1 is a member of the Forkhead box transcription factors (TFs) and carries an important role in embryologic lung development (10).

A link between ACD and FOXF1 was suggested by Stanckiewicz and Shaw-Smith (11), and in their sample of 141 patients they clearly identified 86 pathogenic variants that were involving FOXF1 gene. In addition, heterozygous genomic variants in the FOXF1 locus of ACD patients have been reported by many other research groups as well.

CONCLUSION

ACD/MPV is a rare and lethal developmental disorder. Patients suffer from severe hypoxemia that progresses over time, although awareness is growing among physicians it can still be confused with idiopathic pulmonary hypertension as the presentation can be similar. This usually delays the diagnosis and leads to unnecessary suffering of patients and waist of hospitals resources. As soon as the diagnosis is suspected. Genetic testing should be done or histological exam should be performed, ideally before ECMO or even surgeries for CO occurring anomalies. Although, survival rates are low there are case reports for patients surviving up to 39 months without transplant, prenatal and postnatal genetic testing could contribute to earlier detection, which may allow adequate consultation about prognosis and decision making.

Fig. (4): Bilateral diffuse ground-glass opacification, and a dilated air-filled bowl on the right side.
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REFERENCES