RESEARCH LETTERS

Pulmonary agenesis: Importance of the diagnosis

To the Editor:

Respiratory diseases are currently the most common disorders seen in paediatric clinical practice. Although infections are the most frequent underlying causes, structural alterations may also be responsible. Congenital malformations of the lung are rare disorders occurring with variable degree of severity. Pulmonary artery agenesis is an infrequent congenital anomaly embryologically resulting from early involution of the proximal portion of the sixth aortic arch during development of the heart. The condition may manifest with total or partial absence of the lung on the same side (estimated incidence of total pulmonary agenesis is 0.0034–0.0097%), with the presence of a hyperplastic and compensating contralateral lung.

Bilateral pulmonary artery agenesis is exceptional and incompatible with life. In such situations the right pulmonary branch generally manifests as an isolated lesion, while the left pulmonary branch is usually associated to cardiovascular malformations, with an increased incidence among males.

Pulmonary artery agenesis is usually well tolerated and manifests with repeated infections, haemoptysis and pleuritic chest pain, among other symptoms. Tolerance becomes complicated if the condition is moreover associated to large vessels towards the affected hemithorax, with a displacement of the mediastinum, heart and Great vessels. Visceral-atrial anomalies, with a diagnosis of pneumomediastinum, pulmonary hypertension and minimal interatrial communication. He was also admitted at 8 months of age due to bronchopneumonia. The chest X-rays revealed bilateral basal infiltration, with an interstitial pattern. Posteriorly, he experienced several repeat pneumonia episodes in the right hemithorax, which were resolved with oral amoxicillin (80 mg/kg/day during 10 days). The clinical examination revealed pale skin, while chest auscultation identified right-side hypoventilation, with basal subcruptilates. There was no intercostal indrawing or signs of breathing difficulty. Weight 40 kg (P97), height 137 cm (P97), temperature 37.9 °C, SatO₂ 98%, heart rate 62 bpm, respiratory frequency 28 rpm, blood pressure 118/89 mmHg, and no other relevant findings. Complementary explorations: Tuberculina test: negative. Pharyngeal smear: negative. Inspiration chest X-ray (Figure 1A): symmetric lung volume with smaller size right hemithorax. The expiration X-ray (Figure 1B) in turn showed persistence of the asymmetry, with no air entrapment. Chest angio-CT without contrast injection (Figure 2): absence of right pulmonary artery with diminished volume of the entire right lung, thickening of the interlobular septae, and presence of enlarged-diameter bronchial, phrenic and intercostal arteries in the entire affected hemithorax. No tracheobronchial tree or mediastinal alterations were noted. Visceral-atrial...
situs solitus. Doppler ultrasound: Normal atrioventricular and ventriculoarterial communications. Intact intracardiac septae. No significant valve disease. Normal global ventricular and segmental function. No ductus or aortic coarctation. Right pulmonary artery agenesis. The spirometry tests revealed an obstructive pattern with a diminished forced vital capacity (1.1) (80%) and reduced FEV₁ (1.52) (70%). Significant reversal (210 ml) was noted after salbutamol inhalation. The findings therefore allowed us to establish a diagnosis of right pulmonary artery agenesis.

Since 1673 there have been reports indistinctly of both pulmonary agenesis and hypoplasia – mostly in newborn and young nursing infants. The diagnosis is particularly difficult to establish when the anatomy is markedly distorted. However, CT and MRI allow us to assess the adjacent airway structures and determine the magnitude of vascular compression, tracheal stenosis and tracheal ring fusion, etc., with the added possibility of facilitating surgical planning if necessary.²

The right pulmonary artery is reportedly more often affected, since the contralateral side of the aortic arch is

Figure 1  Chest X-rays in inspiration (A) and expiration (B), showing asymmetric lung volume with smaller size right hemithorax. No air entrapment is seen.

Figure 2  Consecutive angio-CT images with parenchymal window, showing absence of the right pulmonary artery with diminished volume of the entire right lung, thickening of the interlobular septae, and presence of enlarged-diameter bronchial, phrenic and intercostal arteries in the entire affected hemithorax.
normally implicated. A left side location is more often associated with cardiovascular malformations, although this was not confirmed in our case. The literature documents some cases showing irrigation of the parenchyma on the affected side through the persistence of segmental arteries, collateral circulation, intercostal arteries, etc. This is explained by the different embryological origin of these structures, and supports the need for angio-MRI or CT with contrast injection for full evaluation of the patient. The above explains our case, in which right pulmonary artery agenesis was associated to the partial presence of ipsilateral pulmonary tissue. The Spencer classification catalogues the patients as:

- type I (bilateral complete agenesis).
- type II (unilateral pulmonary agenesis): subtype a (associated total absence of bronchi), subtype b (presence of a rudimentary bronchus without lung tissue), and subtype c (poorly differentiated lung parenchyma, the latter corresponding to our own patient).
- type III (lobar agenesis).

Close to 30% of all affected patients may remain asymptomatic for life. However, pulmonary agenesis is associated with chronic and progressive inadequate lung function secondary to the poor development of the tissues, associating skeletal alterations and recurrent respiratory infections. These patients may develop pulmonary hypertension and can present other associated congenital anomalies. Follow-up of the patient and the evaluation of a possible familial or genetic aggregation is important.

Of particular importance for the diagnosis of our patient was the history of repeated respiratory infections. Detailed evaluation of this phenomenon is needed in the context of pulmonary artery agenesis. Respiratory infections are common in childhood, the prevalences being 4%, 2% and 1% per year in children aged 5, 5–9, and over 9 years, respectively. Of these children, a small proportion develop recurrent or persistent respiratory symptoms, as in our patient. The classification of the anatomical distribution of the lungs may be of help in establishing the aetiology of recurrent or chronic pneumonia. The possible causes of disseminated or diffuse infiltrations may be the existence of metabolic, immune or neurological anomalies, while recurrent involvement of a single lung, lobe or segment is suggestive of airway or parenchymal malformations, or the result of atelectasis or hyperinsufflation secondary to obstruction of an airway segment.

Cardiac or large vessel anomalies can give rise to recurrent local pulmonary infections via three mechanisms: 1) diminished blood supply secondary to deficient ventilation (possible explanation in the case of our patient); 2) venous circulatory alterations; or 3) extrinsic large vessel compression of the airway.

The patient prognosis depends on the age at onset of the symptoms, the location of the disorder, the associated malformations, and the presence of lesions on the contralateral side.

Fallot tetralogy is described as the most common cardiac malformation associated to pulmonary artery agenesis — a condition that can be ruled out in our case in view of the age and symptoms of the patient, and the diagnostic exploratory findings. Other related anomalies have also been reported, such as hemivertebrae, hemifacial microsomia, tracheal anomalies, renal alterations or genitourinary malformations, gastrointestinal, hepatic, thyroid or neurological malformations, etc.

At present, the proposed management is based on symptomatic treatment, together with surgery in selected cases. It is estimated that 19–25% of all patients with pulmonary artery agenesis have pulmonary hypertension in adulthood. Knowing that pulmonary hypertension is characterised by a poor production of nitric oxide, possible pharmacological treatment has been evaluated in the form of sildenafil (a selective inhibitor of the enzyme 5-phosphodiesterase, which degrades nitric oxide), with a view to reducing pulmonary hypertension secondary to pulmonary artery agenesis.

Considering that our patient showed no symptoms, no treatment was decided. Two years after the diagnosis the boy continues to lead a normal life.

The present case underlines the need for a high level of suspicion to diagnose congenital lung anomalies, and shows that asthma may appear in such individuals. In this context, appropriate asthma treatment can alleviate the symptoms and reduce morbidity.

References

Cardoon allergy

To the Editor:

The cardoon (Cynara cardunculus), a Mediterranean plant, belongs to the Compositae family closely related to the artichoke. This vegetable is very popular in France, Italy and Spain and the fleshy root and the buds are eaten raw or cooked.

There is no published information regarding cardoon allergy.

A 12-year-old boy reported asthma and rhinoconjunctivitis in the spring seasons during the last 6 years. He referred, as well, pruritus around mouth after eating nuts, apple, melon, peach, oranges and tomatoes and in the last year, pruritus in mouth and ears and lip swelling after eating raw cardoon. He tolerated cooked cardoon and artichoke.

The patient was tested with a battery of aeroallergens comprising pollens (grass, Olea europaea, Artemisia vulgaris, Parietaria judaica, Platanus acerifolia and Chenopodium album), house-dust mites, fungi, and animal dander. He was also prick tested with commercial food extracts, including nuts (hazelnut, almond, peanut, sunflower seed, pistachio and walnut), fruits (peach, apple, orange and melon), green vegetables (tomato and cardoon), profilin and prick-by-prick with cooked and raw cardoon. Histamine phosphate (10 mg/mL) and sterile 0.9% saline were used as positive and negative controls, respectively.

Specific IgE to extracts from cooked and raw cardoon and pollens from Lolium perenne, O. europaea, A. vulgaris, P. judaica and P. acerifolia was measured by an immune assay (Enzyme AllergoSorbent Test EAST).

SDS-PAGE Immunoblotting was performed according to the method of Laemmli.

Separated protein bands were electrophoretically transferred to polyvinylidene difluoride (PVDF) essentially as described by Towbin et al. and blocked during one hour at room temperature with 5% defatted dry milk in Tris buffered saline. Membranes were incubated overnight at 4 °C with patient’s serum followed by anti-human IgE-horseradish peroxidase conjugate. Detection was carried out by a chemioluminiscient (Amersham ECL Plus Western Blotting Detection System. GE Healthcare). When Western inhibition assay was performed, patient serum was preincubated with cardoon extract (1mg/ml) with the inhibition phases during four hours at room temperature, and afterwards the serum samples were incubated with the PVDF membrane and IgE binding revealed as described herein.

Doubled-blind, placebo-controlled oral challenge tests with raw and cooked cardoon were made.

Skin prick tests with pollen of grass, Olea europaea, Artemisia vulgaris, Parietaria judaica, Platanus acerifolia and nuts (hazelnut, almond, peanut, sunflower seed, pistachio and walnut) were all positive. The prick test with profilin was also positive.

Prick-by-prick with peach, orange, melon, apple and tomato were positive.

Prick-by-prick with raw cardoon was positive, 11×10 mm (with negative controls) and negative with cooked cardoon.

Specific IgE by means of EAST assay was positive to extracts from raw cardoon (0.6 kU/L; class 1), cooked cardoon (0.4 kU/L; class 1) and pollens from Lolium perenne (>100 kU/L; class 5), Olea europaea (>100 kU/L; class 5), Artemisia vulgaris (10.3 kU/L; class 3), Platanus acerifolia (12.5 kU/L; class 3) and Parietaria judaica (1.9 kU/L; class 2).

The protein composition of the raw cardoon extract was studied by the SDS-PAGE and Coomassie staining with protein bands ranging from 66 to 14 kDa being observed. The electrophoresed and electrotransfered raw cardoon extract was incubated with patient’s serum, revealing a broad and intense IgE binding area between 97 and 20 kDa.

Western blotting inhibition of raw cardoon extract with patient serum showed a total IgE-binding inhibition when

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Figure 1 SDS-PAGE immunoblotting inhibition. Lane C: control serum, Lane 1: patient’s serum, Lane 2: patient’s serum preincubated with cardoon extract (1mg/ml) (Positive inhibition control), Lane 3: patient’s serum preincubated with Lolium perenne pollen extract (1mg/ml), Lane 4: patient’s serum preincubated with Olea europaea pollen extract (1mg/ml), Lane 5: patient’s serum preincubated with Platanus acerifolia pollen extract (1mg/ml) Lane 6: patient’s serum preincubated with Artemisia vulgaris pollen extract (1mg/ml) Lane 7: patient’s serum preincubated with lamb extract (1mg/ml).

M: molecular weight marked proteins.