A “sweet” hydrothorax in a child on peritoneal dialysis
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Abstract
Peritoneal dialysis (PD) is an established, effective long term renal replacement treatment modality for children with end stage renal disease (ESRD). A rarely reported complication of PD in children is the development of hydrothorax. We report the case of an 8-year-old boy that developed a right-sided pleural effusion during automated PD (APD), in order to raise awareness amongst paediatricians; we also review the diversity of clinical presentation and the available diagnostic tools, discuss theories regarding aetiology and highlight the available treatment options. Hippokratia 2011; 15 (4): 358-360

Key words: peritoneal dialysis, child, hydrothorax

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Case description
The patient presented is an 8-year-old boy with ESRD due to chronic tubular-interstitial nephritis who had been on APD by night for 6 months. His APD prescription was 1000mL (30mL/Kgr) volume exchange by night, without hyperosmolar solutions, with 70 minutes dwell period. Dialysis had been well tolerated apart from a single episode of peritonitis. He had residual renal function consisting of 1500 – 2000mL diuresis per day. On a routine follow – up visit he complained of a dry cough lasting 3 days and dyspnoea. Respiratory rate was 35 breaths/minute, heart rate was 110 beats / minute and his oxyhaemoglobin saturation was 94% in room air. Auscultation revealed attenuated breathing sounds over the lower half of the right chest and dullness was noted on percussion; he was apyrexial and there was no evidence of heart failure or peripheral oedema. Chest radiography confirmed a massive right-sided pleural effusion with a cardiac silhouette of normal size (figure 1). Insertion of a chest drain was uneventful and aspirated 3.6 Litres of straw-coloured pleural fluid over 24 hours. The results of simultaneously drawn serum and pleural fluid are shown in table 1. Bacterial and mycobacterial cultures were negative. To prevent the hydrothorax from enlarging the dialysate dextrose concentration was initially increased to 2.27%, the mean exchange volume was decreased to 500 mL / cycle (15 mL / Kgr), the dwell period was shortened to 40 min and the patient was advised to adopt a semi-sitting position during dialysis. Peritoneal scintigraphy with Technetium-99m macroaggregated human albumin (Tc-99m MAA) revealed diffusely increased isotopic activity in the right hemithorax (figures 2a & 2b), confirming the suspected presence of a pleuroperitoneal communication. Magnetic Resonance Imaging (MRI) failed to demonstrate the communication. APD was temporarily interrupted as pleural effusion continued even with smaller cycle volume, and the patient was transferred temporarily to haemodialysis. APD was resumed 6 weeks later, without clinical or radiographic recurrence of the effusion.

Discussion
Acute hydrothorax is a rare complication of PD. The first case of pleural effusion during APD has been reported by Edward and Unger in 1967², whereas Lorenz published the first paediatric cases in 1979³. The incidence reported varies from 0.9%⁴ to 20%¹; most authorities give an estimate of 2-3%. In our tertiary University Paediatric Department this is the first case of APD-related hydrothorax over 20 years. During this period 63 children (aged 1 month to 18 years) received APD both for acute and chronic renal failure.

It usually affects the right hemithorax and there is no clear sex predominance⁴. Bilateral hydrothoraces in
children on peritoneal dialysis are very rare. The interval between onset of CAPD and development of a pleural effusion can range from a few hours to 8 years. It has been suggested that it is more common in children receiving APD for acute renal failure secondary to haemolytic uraemic syndrome. Krishan et al reported that a major factor in the development of an acute hydrothorax with APD appears to be starting to cycle with 40 mL/kg volumes, whereas increasing to this volume from initial 10 mL/kg cycles over 6 days prevents hydrothorax. However in our case the child was on stable peritoneal dialysis prescription for the last six months with an exchange volume less than 40ml/kg.

Clinical presentation can vary from being a serendipitous discovery in asymptomatic children to respiratory compromise in children in extremis. Our patient was identified during regular follow up. Most children report retention of the dialysate, cough, dyspnoea and costal discomfort. The presence of the effusion can be confirmed by radiography and sonography. The clinician may face difficulties in the differential diagnosis of APD-related hydrothorax when other potential aetiologies of transudative effusions may coexist, such as inadequate ultrafiltration, hypoalbuminaemia, congestive cardiac failure, or tuberculosis pleurisy, especially in chronic cases. Therefore, meticulous attention to history taking and clinical examination is of utmost importance. Failure to recognise this entity may lead to inadvertent prescription of hypertonic dialysis exchanges, with grave consequences.

Hydrothorax in this context is rightfully called “sweet hydrothorax” as hypertonic glucose solution fills the pleural space. Therefore, the hallmark of the diagnosis is demonstrating a positive pleural fluid to serum glucose concentration gradient. In the patient presented herein, glucose concentration in the pleural fluid aspirate was more than three times higher than that in the serum at the same time. The pleural fluid glucose concentration can vary substantially, depending on the size of the pleuropertitoneal communication, the rate of glucose absorption from the pleural space, the concentration gradient between the two compartments and the timing of the thoracocentesis.

The pathogenesis of APD-related hydrothorax remains unclear; the most likely explanation is the unidirectional flow of the peritoneal fluid via congenital or acquired diaphragmatic defects. Contrast studies via the dialysis catheter, peritoneal isotopic studies or even direct inspection at surgery do not always reveal a communication. In our patient Tc-99m MAA scintigraphy was helpful. Alternative mechanisms that have been postulated include passage of fluid to the pleural cavity through lymphatic channels in the diaphragm or lymphatic leakage from the thoracic duct.

Various therapeutic approaches have been proposed. Nomoto et al reported complete resolution of the hydrothorax following brief interruption of the APD – as was the case in the patient presented herein – in 19 out of 50 patients in a large retrospective study. They speculated that this brief interruption allowed the diaphragmatic defect responsible for the pleuropertitoneal communication to heal and made return to APD feasible. Reduced volume dialysis has also been used with success Pleurodesis with tetracycline, autologous blood, or other sclerosing agents instilled into the pleural cavity via thoracic tubes have been proposed in order to allow resumption of full volume APD in patients in whom conservative medical management had failed. Recent advances in video assisted thoracoscopic surgery (VATS) promise great advances both in treating visible lesions such as diaphragmatic

**Table 1: Results of chemical analysis of simultaneously drawn serum and pleural fluid**

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<thead>
<tr>
<th></th>
<th>Serum</th>
<th>Pleural Fluid</th>
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<tbody>
<tr>
<td>Glucose (mg/dl)</td>
<td>91</td>
<td>312</td>
</tr>
<tr>
<td>Total protein (g/dl)</td>
<td>6.8</td>
<td>0.3</td>
</tr>
<tr>
<td>Lactic dehydrogenase (U/L)</td>
<td>218</td>
<td>32</td>
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</tbody>
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**Figure 2a & 2b:** Peritoneal scintigraphy with Technetium-99m macroaggregated human albumin (Tc-99m MAA) revealed diffusely increased isotopic activity in the right hemithorax.
blebs and also mechanical or chemical thoracoscopic pleurodesis.

To conclude, clinicians involved in the treatment of children on APD must have a high index of suspicion for this rare, but potentially life threatening complication.

References