INTRALUMINAL STONE IN A PD CATHETER: THE THIRD WORLD CASE

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Résumé

Jusqu’à présent, seulement 2 cas de lithiases intraluminales du cathéter de DP ont été décrits dans la littérature. Une femme âgée de 68 ans, en DPCA a été admise à l’hôpital pour une difficulté intermittente de drainage par son cathéter péritonéal. Un calcul a été découvert dans la lumière du cathéter. Sa composition était identique à celle déjà décrite dans les deux cas précédents: une composition d’hydroxyphosphate de calcium carbonaté (carbapatite 83 %) avec une couche de protéine (17 %). Le calcul ne pouvait pas migrer à travers la partie intrapéritonéale du cathéter en raison de sa taille (3 x 4 mm), plus importante que la lumière du cathéter, malgré les épisodes journaliers de rinçage par injections et drainages du dialysat. Dans les 3 cas le dialysat contenait du lactate et 1,75 mmol/L de calcium.

Abstract

Only 2 cases of intraluminal peritoneal catheter lithiasis have been reported so far. A woman (68 years of age) on CAPD was admitted to hospital because of intermittent outflow obstruction of her peritoneal catheter. A stone was found in the catheter lumen. Its biochemical composition was the same as that of the 2 other cases previously described: calcium-hydroxycarbonate-phosphate with an admixture of protein. The stone could not migrate through the intraperitoneal tip hole of the catheter because its size (3 x 4 mm) was greater than the catheter lumen (2.7 mm). That suggests in situ development, directly in the catheter lumen despite inflow-outflow dialysate daily regular rinsings. In the 3 cases, dialysate contained lactate and 1.75 mmol/L of calcium.

Mots clés : dialyse péritonéale, catheter, calcul

Keywords : peritoneal dialysis, catheter obstruction, stone
So far, only 2 cases of intraluminal lithiasis of the PD catheter have been described in the literature: in a 62-year-old man after 6 years of PD [1], and in a 65-year-old woman after 2 years of PD [2].

A 68-year-old woman, who has been on CAPD for 1 year, reported to the hospital for “one-way obstruction” type of drainage difficulties. The X-ray demonstrated that the catheter was in place. Thrombolysis was attempted by instillation of Alteplase according to the protocol of the service, followed by 2 weekly injections of heparin. In the aftermath, however, irregular drainage and normal infusion were consistent with continued peritoneal dialysis. One month later, the patient returned to the hospital for an impossible drainage. On examination, we discovered a lithiasis completely obstructing the catheter lumen, at the connector level of the extension line. Lithiasis was extracted after section and repair of the catheter. The dialysis then resumed without any problem.

Lithiasis was oval, irregular, 4 x 3 x 2 mm size with a light yellow-brown color. The analysis revealed it was composed of carbonated calcium hydroxyphosphate (83% carbapatite) with a layer of protein (17%). The patient had started PD one year earlier for renal failure secondary to chronic tubulointerstitial nephropathy. She had no history of nephritic colic or lithiasis. Six months before the cholelithiasis, she presented a peritonitis without identified organism, of good evolution under antibiotic therapy. She was treated with CAPD with 3 continuous daily exchanges, including a long nocturnal exchange of an icodextrin-based solution. The calcium concentration of the dialysate was 1.75 mmol / L, the magnesium concentration was 0.50 mmol / L (0.25 mmol / L for icodextrin). On admission, the patient had poorly controlled secondary hyperparathyroidism with 2.15 mmol / L serum calcium and 2.48 mmol / L phosphoremia. The intact PTH was 215 ng / L.

Secondary hyperparathyroidism has been suggested as a leading cause of peritoneal lithiasis [1, 2]. As in our observation, Antoniou described a significant hyperphosphoremia due to poor adherence to chelation therapy [1]. However, in the case described by Skaro [2] hyper-
parathyroidism was perfectly controlled and the patient had no hyperphosphoremia.

In our case as in that of Antoniou [1], there is a history of peritonitis. However the case reported by Skaro [2] had no peritoneal or emergent infection.

The physicochemical composition of the lithiasis is similar to that of the two other cases reported in the literature: carbonated calcium hydroxyphosphate with a protein layer. Its shape and color are also similar. In the two cases previously described, the authors evoke the formation of a lithiasis in the peritoneal cavity, and its secondary migration in the PD catheter. However, in our observation, the size of the calculation suggests lithogenesis in situ, directly in the catheter lumen. Indeed, the internal diameter of the catheter is about 2.7 mm and our lithiasis was 4 x 3mm. With this size, the lithiasis would not have been able to cross the intraperitoneal distal orifice of the catheter from the peritoneal cavity. In addition it was necessary to cut the catheter to extract the lithiasis. In the case reported by Antoniou [1], the lithiasis measured 3.9 x 3 mm and the catheter had to be deposited. The lithiasis described by Skaro [2] was smaller: 2.5 x 1.5 mm, and could be aspirated with the syringe.

This observation suggests a local mechanism at the origin of the development of lithiasis in the catheter lumen, despite the multiple rinsing of the catheter by the peritoneal dialysis solution. In the three cases described, the lithiasis was of the same physico-chemical composition; the dialysate contained lactate, and had a calcium concentration of 1.75 mmol / L.

In spite of the rarity of this complication, the presence of lithiasis of the peritoneal dialysis catheter must be evoked in case of drainage problems.

**DISCLOSURE**

Authors declare no conflict of interest

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