Pneumoperitoneum Secondary to Barotrauma Following a Diving Accident

Neumoperitoneo secundario a barotrauma tras accidente de buceo

We present the case of a 35-year-old male patient who came to our consultation with abdominal distension that had been developing over the previous 10 h, with no other symptoms. The distension had begun immediately after free-diving to a depth of 39 m. His personal history included left inguinal hernioplasty. On physical examination, the patient was in good general condition: no fever, hemodynamically stable and eunepic. The abdomen was soft, with great distension and generalized hyperresonance as well as crepitus upon palpation of the umbilical region, which presented a non-complicated umbilical hernia. The patient had no pain during abdominal palpation or any signs of peritoneal irritation. Air-fluid levels were detected. The remaining physical exploration was unremarkable.

Blood analysis showed 14,490 leukocytes (normal formula), GOT 53 U/l, GPT 74 U/l, CPK 208 U/l, CKMB 40 U/l and myoglobin 46.49 ng/ml. The remaining determinations, including coagulation and venous blood gases, had values within normal ranges.

When we observed the shape of the transverse colon wall on standing chest and abdominal radiographs (Figs. 1 and 2), a normal intraluminal hydro-air pattern was seen with extraluminal free air accumulated in the subdiaphragm region as well as displacement of the lung bases and viscera of the upper abdomen. In addition to the voluminous pneumoperitoneum pushing the abdominal content in the posterior direction and umbilical hernia with fat and gas content, thoracoabdominal CT also revealed pulmonary laminar atelectasis due to compression and minimal pericardial effusion. No intraperitoneal free fluid, intravascular gas or disruption of hollow organs was observed. Electrocardiogram was normal.

Given the good general status of the patient, the lack of significance found in the analyses and the lack of evidence of perforation, we opted for conservative management with no oral intake, parenteral nutrition, normobaric high-flow oxygen therapy (MV at 35%–8 lpm) and low-molecular-weight heparin at a prophylactic dosage. Oral tolerance was started 48 h later, and enteral nutrition was reinstated without complications after 72 h. During hospitalization, the patient passed gas and had normal bowel movements with no pathologic production. He presented isolated fever (38.1 °C) on the third day after hospitalization, which responded to the

* Please cite this article as: Rosado Dawid NZ, Peraza Casajús JM, Bodega Quiroga I. Neumoperitoneo secundario a barotrauma tras accidente de buceo. Cir Esp. 2014;92:368-370.
cases published since 1977.1–6 Perforation of a hollow organ should first be suspected as it is frequently the origin of pneumoperitoneum, since less than half of perforated patients show signs of peritoneal irritation.7 As for the mechanism by which the gas reaches the peritoneum in the absence of organ perforation, many theories have been postulated. It has been accepted that the most probable origin of this air is the lungs. Several authors have demonstrated in animal models that after the rupture of distended alveoli, the gas that is released dissect s pleural bullas and penetrates the pleural space. If the pressure continues, the air enters the retroperitoneum and the subcutaneous cell tissue through dissection of muscle planes and diaphragm defects that lead to the contact between the retroperitoneal and pleural cavities. Anatomical alterations secondary to childbirth or previous abdominal surgery, as in this case, could favor the appearance of pneumoperitoneum. Another possible mechanism is the passage of air under pressure from the pleura through the thoracic duct to the abdominal cavity in a retrograde pathway favored by sudden ascent. It could be possible to consider a mechanism similar to what happens during colonoscopy, where, secondary to the increased intraluminal pressure, microperforations of the colon may occur in areas with thinner walls (for instance, in the diverticula) or in the lymphatic vessels in this area that are small enough to allow for gas to escape without the extravasation of any intraluminal liquid content, then closing up after the distension subsides. Or the air could even pass through walls that are intact but are less thick, with no demonstrated perforation.9

The diagnosis of pneumoperitoneum is based on symptoms and radiology. Abdominal distension and compromised respiratory function are the most frequent symptoms and are directly related to the amount of air leaked. A simple radiograph in the standing position is diagnostic, although tomography is more sensitive and is able to rule out organ perforation. Another useful tool is radiology with hydrosoluble contrast, which locates possible perforations.

If there is clinical compromise or evidence of perforation, the approach is usually surgical with emergency laparotomy, although 10% of cases can be managed conservatively.10 According to the literature, the therapeutic approach in pneumoperitoneum after barotrauma is varied: hyperbaric chamber, puncture-evacuation of the peritoneal cavity, a wait-and-see approach with no oral intake and parenteral nutrition, or surgical intervention. There is no established protocol for the management of these cases. Our criteria are that if there is no evidence of perforation of a hollow organ, there are no other manifestations of barotrauma and the clinical situation allows for it, conservative management can be used, as in the case that we have presented.

REFERENCES

Urgent Surgery in a Kasabach-Merrit Syndrome Associated With a Giant Hepatic Angiosarcoma

Actuación quirúrgica urgente en un síndrome de Kasabach-Merrit asociado a angiosarcoma hepático gigante

The patient is a 70-year-old woman with no prior medical history of interest who came to our Internal Medicine Department complaining of pain and the sensation of a mass in the right hypochondrium. The lab work done at that time showed a normal hemogram and biochemistry. Abdominal-pelvic CT revealed the existence of a liver mass occupying the entire right hepatic lobe, irrigated by portal branches and compatible with a giant hemangioma. The patient was sent to our Hepatobiliary Surgery Unit. Magnetic resonance imaging of the liver was ordered in order to be able to better characterize the tumor, assess its relationship with different vascular structures and plan the surgical technique.

The patient came to our Emergency Department 48 h after the MRI test due to epigastric pain and syncope. She presented with tachycardia (HR 130), hypotension (70/40) and tachypnea (RR 40). The blood work-up showed anemia (Hb 5.8), severe thrombocytopenia (platelets 42,000) and altered coagulation (INR 2.1). The patient was hemodynamically stabilized with fluid therapy and the transfusion of 6 units of packed red blood cells; the coagulation was corrected with the transfusion of 4 pools of platelets and 2000 ml of fresh plasma. At that time, we assessed the magnetic resonance (Fig. 1), which demonstrated the existence of a voluminous liver mass measuring 18 cm × 12 cm in the right liver lobe with intralobial bleeding.

The patient was assessed by the Interventional Radiology Department of our hospital. Abdominal ultrasound showed that there was no intra-abdominal free fluid. At this time, the possibility of performing arteriography and embolization of the right hepatic artery was ruled out due to that fact that the intratumoral bleeding was from the right portal branch. The patient was taken to surgery, where we carried out an extended right subcostal laparotomy and observed a large tumor mass that occupied the entire right liver lobe from the right dome of the diaphragm to the pelvis, which was elastic in consistency and violet in color (Fig. 2). We performed right hepectomy under total vascular exclusion. The patient was discharged on the 8th day post-surgery with no complications. In the pathology study of the surgical specimen, a neoplastic proliferation of epithelioid or fusiform cells was observed with atypia that formed nests or vascular structures interspersed with areas of hemorrhage and necrosis, showing cellular uptake with vascular

Please cite this article as: González Rodríguez FJ, Domínguez Comesaña E, Portela Serra JL, Lede Fernández Á, Piñón Cimadevilla MA. Actuación quirúrgica urgente en un síndrome de Kasabach-Merrit asociado a angiosarcoma hepático gigante. Cir Esp. 2014;92:370–372.