Hydatid disease or echinococcosis in humans is caused by infection of the Echinococcus genus cestodes. E. granulosus is the most frequent in Spain and it is associated with cystic echinococcosis [1]. Its frequency has increased in recent years [2]. It usually presents asymptomatic, although an increase in the size of the cyst may lead to the appearance of symptoms. Its manifestation as anaphylactic shock is exceptional, but it should be considered a differential diagnosis that gave the increase in incidence in recent years [3].

A 32-year-old farmer man was admitted as an emergency with pruritus predominantly in the hands, feet, and armpits associated with erythema, facial edema, and generalized raised rash lesions. He referred to respiratory distress and dizziness. The examination revealed hypotension and tachycardia (BP 90/55 mmHg, HR 115 bpm), with a basal oxygen saturation of 90%, without respiratory drainage. Symptoms and hemodynamic status improved after the administration of corticosteroids, epinephrine, and dexchlorpheniramine (BP 110/89 mmHg, HR 52 bpm).

Blood analysis revealed leukocytosis (20,890 / mm3), without other notable findings. Liver enzymes were at normal levels. Chest radiography showed no pathological findings. During hospital care, he presented one episode of fever of 38ºC of unknown origin (negative blood cultures, non-pathological urine analysis, and not associated with respiratory symptoms). After 48 hours of monitoring and close following, the patient remained asymptomatic, and it was discharged home.
The study by the specialty of Allergology showed eosinophilia (15%), high levels of IgE with low C4 (total IgE 1274 kU/ L; complement C4 12.2 mg / dL) and positive serology for E. granulosus (antibody titer: 2560). A thoracic-abdominopelvic computed tomography (CT) was performed and showed a 10.6 cm x 7.3 cm x 8 cm well-defined cystic lesion in the right hepatic lobe with small peripheral calcifications, as well as dense membranes and papillary-looking lesions, corresponding to a CE3 transitional hydatid cyst (WHO classification) (Figure 1).

Following these findings, a surgical intervention was decided and antiparasitic albendazole 500mg every 12 hours was started for a month prior to it. A cystopericystectomy was performed using a right subcostal laparotomy without incident (Figure 2). The patient was discharged after 6 days in hospital without postoperative complications, completing treatment with albendazole 500 mg every 12 hours for an additional month after surgery.

An anaphylactic reaction is a rare presentation of hydatid disease (between 1-7.5%) [7]. It can occur if the content of the cyst is released to a host that has IgE antibodies developed by a previous leak. We can find anaphylaxis without identifying macroscopic ruptures of the cyst [2]. The management of hydatid cysts varies depending on the location, extension in the affected organ, and possible associated complications (rupture or haemorrhage) [8]. Standard treatment includes adjuvant chemotherapy and surgical resection. Cystopericystectomy is considered the curative surgical approach. The recommended chemotherapy treatment (four days before surgery and at least one month after) is 10-15mg/kg of albendazole daily divided into two doses [9, 10].

The most common toxic effect of albendazole is the elevation of liver enzymes (20% of cases). It has also a suppressive effect on the bone marrow. Therefore, it is important to monitor liver enzyme levels and blood count [10]. In conclusion, even though secondary anaphylaxis to hydatid disease is rare (1-7%), it should always be included in the differential diagnosis of an episode of the anaphylactic reaction of unknown cause due to the severity of the disease and the specificity of the required treatment.

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