Portal hypertension as a result of the incomplete surgically treated advanced alveolar echinococcosis: a case description
Ł. Pielok 1*, M. Karczewski 2, W. Cierach 2, P. Zmora 3, E. Lenartowicz 3 and J. Stefaniak 1

Abstract

Background: Infection of Echinococcus multilocularis causes in humans the alveolar echinococcosis. Although the infection has world-wide distribution it is rarely detected. Diagnosis of alveococcosis is difficult because of not typical clinical picture and irregular results of radiological examinations suggesting neoplastic process which begins in the liver tissue or in the biliary tracts. The parasitic growth is slow, so the illness is quite often established in late invasion period. Treatment of long-lasting and late diagnosed infection is difficult and requires cooperation of parasitologists together with surgeons to avoid life-threatening organ dysfunction.

Case presentation: We describe a young male patient, diagnosed, according to the radiological, immunological and histological examination results, infection of Echinococcus multilocularis, who was treated with not radical resection of pathologic mass together with persistent albendazole intake. The right hepatectomy was performed. In addition, visible cysts were removed from the left lobe of the liver in nonanatomical resection and suspicious calcified lesions in hepatoduodenal ligament were also removed. After the operation portal hypertension, with splenomegaly and symptoms of the liver cirrhosis occurred (thrombocytopenia, collateral venous circulation, first degree varices oesophagii). The portal hypertension probably could be a result of incomplete surgery due to extended parasitic infection and liver anathomical changes due to performed procedures, because the portal hypertension and it’s further complications had not been observed before the operation.

Conclusions: Echinococcus multilocularis should be taken under consideration in differential diagnosis of irregular lesions within the liver. Lon-lasting invasion could be responsible for the irreversible secondary liver changes such as cirrhosis and portal hypertension. The surgery treatment (treatment of choice) is difficult and it’s results depends on the invasion period the patient is operated on. After the surgery the patient requires careful follow – up, to detect early complications.

Keywords: Echinococcus multilocularis, Human alveococcosis, Portal hypertension, Liver, Joidance

Background

Alveolar echinococcosis, caused by the metacestode of the fox tapeworm Echinococcus multilocularis, is the most pathogenic zoonosis in temperate and arctic regions of the northern hemisphere [1]. Parasite transmission occurs when eggs of the tapeworm, excreted by the final hosts (usually foxes but also dogs, wolves and cats), are ingested accidentally by humans [2, 3]. In humans E. multilocularis infection is one of the reasons of liver lesions. For many years the illness is not detected because is asymptomatic. Because of increase of foxes population in Poland the risk of parasite transmission to humans is mounting [4]. Diagnosis of alveococcosis is difficult because of not typical clinical picture
and irregular results of radiological examinations (ultrasound of the abdomen cavity - USG, computed tomography-CT, magnetic resonance imaging-MRI) suggesting neoplasmatic process which begins in the liver tissue or in the biliary tracts [5–7]. According to the pathologic lesions localization the PNM (primary liver location, involvement of neighbouring organs and metastatic changes) classification is used to evaluate the disease advanced [8]. Helpful in diagnostics are serology tools performed by screening ELISA- enzyme-linked immunosorbent assay method, which detects non-specific anti-Echinococcus IgG. Western-blot confirms the diagnosis, and EM2-EM18 ELISA detects very specific anty-multilocularis antibodies [9–12]. In controversial cases the diagnosis can also be confirmed after histopathology section of the liver tissue or after performing polymerase chain reaction- PCR, which detects parasite DNA fragments [13, 14]. Long asymptomatic parasite’s development causes that diagnosis is often established in advanced infection period, which delay initiation of specific treatment. All these leads to progressive organ dysfunction with full symptomatic liver cirrhosis [15].

After only several years, when the patient is not treated, cholestasis develops, thrombotic disturbances appears and changes in other distant organs [16]. These all pathologic processes as well as presence of the parasite is responsible for the full symptomatic liver cirrhosis [15].

Echinococcus multilocularis infection should be taking under consideration in differential diagnosis in patients with non –specific liver focuses, specially suspected of neoplasmatic disorders with normal liver function tests-LFTs (GGTP, ALP, ALT, AST) [20]. Early alveococcosis diagnosis and suitable treatment initiation, could protect the patient from life –threatening complications, which correlates with longer survival and better quality of life [21].

In this work we present a case of a young man with a huge pathological mass within the liver, who was diagnosed alveococcosis and treated with the not-radical operation theater together with albendazole (Zentel, GSK) intake in whom the portal hypertension occurred as a postsurgical complication.

**Case presentation**

A 31-year old male patient admitted to the Tropical and Parasitic Disease Department of Poznañ University of Medical Sciences, Poland, because of the presence of a tumor-like lesion within the liver. The patient had been living in a small village surrounded by forests in which a big foxes population has been detected.

Prior to the admission the patient had suffered from influenza like syndromes, pain in the right subcostal region and suddenly joindance.

He was admitted to the local Surgery Department with suspicion of biliary tract pathology. CT scan gave the evidence of irregular mass with disseminated calcifications. He was diagnosed undifferentiated hepatitis with cholestasis.

Because of atypical radiology results suspicion of Echinococcus infection was done. ELISA serology test was positive (2.9 Units; positive above 1.0). The patient was moved to the Tropical and Parasitic Clinic in Poznañ for further investigations.

On admission day the physical examination was unremarkable. Blood tests showed elevated levels of bilirubine (2 mg%), alkaline phosphatase (172–248 U/l) gamma glutamyl trans peptidase- GGTP (135–262 U/l). USG of the abdominal cavity revealed presence of a huge calcified lesion in the VII-th liver segment with the diameter of 12.3 × 2.8 cm and in the II-nd liver segment a solid hyperechogenic focus with calcifications inside as well as disseminated calcifications in the interhepatic biliary tracts neighborhood. MRI showed the liver enlargement, with irregular tissue. In the VII, VI and V segments polycyclic fluid lesion and disseminated inside the right lobe smaller fluid foci as well as biliary tract widening (Fig. 1).

According to the picturesque data suspicion of alveococcosis was done. ELISA test (Echinococcus IgG) was positive – 50 NTU (positive above 11NTU) and confirmed with positive
Western-blot which revealed presence of specific for Echinococcus multilocularis IgG (7,16,18, 26–28 kDa). ELISA EM2-plus (anti-E.multilocularis) was also high positive (> 3.0ABS). The patient was finally diagnosed the liver alveococcosis with P2M0N0 stage.

When the diagnosis was established the albendazol therapy (2x400mg/day) was initiated together with ursodeokyscholic acid (2x250mg/day) in order to lowered bilirubine level and protect from apoptosis healthy liver tissue. The patient was qualified to the surgery treatment and was moved to the General and Transplant Surgery Department, Poznan University of Medical Sciences. MELD score of the patient was 10 (creatinine 1.13 mg/dl; bilirubin 1,12 mg/dl, INR 1,15, not dialysed).

The right hepatectomy was performed. Access to the liver was achieved by bilateral subcostal incision and then mobilizing the liver from its ligamentous attachments, including the coronary ligament, and left and right triangular ligaments, then anatomic resection of the fifth, sixth, seventh and eighth liver segments was performed. Right portal vein, right hepatic artery and right hepatic duct was ligated and cutted. In addition, visible cysts were removed from the left lobe of the liver in nonanatomical resection and suspicious calcified lesions in hepatoduodenal ligament were also removed (Fig. 2). Postoperative course complicated by lymphorhea, conservative treatment was initiated, obtaining improvement. In ultrasound on the 8th postoperative day, spleen enlargement occurs (141x55mm).

The histopathology examination of the all 3 specimens revealed presence of chronic inflammatory changes with thick wall calcified granulomas and accellular homogeny infiltrates with necrosis cavities.

Additionally, the biological material after surgery was used for molecular genotyping of Echinococcus sp. For this reason, the DNA was extracted from the removed liver tissue with lesions, using the commercially available kit (NucleoSpin Tissue, Macherey Nagel, Dueren, Germany). Next, the isolated DNA was used as a template in the PCR with Echinococcus sp. 12S rRNA-specific primers (EM-H15 and EM-H17), according to Stieger et al. (2007) [22]. Moreover, as a control of DNA quality and presence of PCR inhibitors, we used isolated genetic material to amplify human GAPDH, according to Xiang et al. (2012) [23]. Finally, the PCR product was sequenced and the obtained sequences were aligned using BioEdit software.

Based on the PCR reactions with Echinococcus-specific primers, we confirmed that the patient was infected with Echinococcus sp. (Fig. 3a). Further sequence analysis revealed the complete similarity with Echinococcus multilocularis 12S rRNA (Fig. 3b), suggesting that the clinical outcome and after surgery complications were not correlated with parasite genotype and potential more pathogenic Echinococcus multilocularis strain.

After the operation albendazol treatmet was continued. With no other major or minor complication patient with normal level of bilirubin, alanine transaminase and aspartate transaminase was discharge on 17th postoperative day.

The patient’s follow –up, performed 10 months after the operation gave the evidence of thrombocytopenia (80G/l), leucopenia (3.1G/l) and syderopenic anemia. MR of the abdomen cavity showed presence of numerous hypodensic partially calcified lesions within the remaining liver segments. Moreover critical portal vein constriction (the diameter 3-4 mm), collateral venous circulation in the liver hil were detected. The examination revealed also splenomegaly as a result of the portal hypertension (Fig. 4). Performed endoscopy of upper gastrointestinal tract revealed presence of first degree varices oesophagi as well as gastritis and duodenitis. Recently, patient had control MRI (06.03.2020) and there are no active outbreaks of alveococosis in the liver parenchyma, compared to previous studies, the image is stable. Collateral vessels are visible in the liver cavity. The spleen is enlarged, by a maximum length of about 16 cm (before operation spleen length was 14 cm). Patient is clinically asymptomatic. But he still requires regular, every 6 months follow-up in the Tropical and Parasitic Clinic.

Discussion and Conclusions
Infection caused by Echinococcus multilocularis is the most dangerous parasitic zoonosis in Europe. Left untreated, it has a very high mortality rate after located in vital organs such as liver, lungs, brain [24]. The alveolar cyst causes a maliging tumor-like lesions with infiltrative, proliferative and destructive character, which locates in the liver primarily, then metastasizes to the other organs.
Echinococcus multilocularis is a parasite with capabilities of mass-forming affect neural and vascular invasion. Within the liver, the parasite’s unrestricted growth is responsible for focal calcifications and finally the whole necrosis. Due to vascular and neural invasion Buddi-Chiari syndrome may be the clinical presentation of this condition [26].

Disseminated forms of alveococcosis appears to be associated with invasive growth of the parasite and the related necessity to perform vast resection of the liver and adjacent organs, as well as resection and prosthetic repair of the major vessels [17].

Long lasting and late diagnosed alveococcosis leads gradually to the liver dysfunction with its internal structure rebuilding. Consequences of this are changes in the portal vein circulation and liver cirrhosis [16]. Hepatofibrosis as a result of the parasite-host interaction is a pathological feature of Echinococcus multilocularis infection that destroys normal liver tissue leading to the portal hypertension and the formation of peripheral fiberboards around the metacestode is a major reason of clinical symptoms appearance [27, 28].

To avoid serious and often irreversible life-threatening complications for the patient the diagnosis of Echinococcus multilocularis infection should be early obtained and surgical treatment should be immediately initiated, because delayed surgery gives inadequate results. Surgical treatment can be radical or sparing. Radical procedures are associated with a lower risk of relapse but they carry a greater risk of postsurgical complications. Moreover, in advanced alveococcosis only palliative operation can be performed to improve quality of the patient’s life. Postsurgical frequency of recurrences is connected to incomplete parasite’s excision [29]. Lately, in literature you can find descriptions of new surgical techniques such as associating liver partition and portal vein ligation, which can be safely in young individuals [30].

Described, minimally invasive surgery, the laparoscopic and robotic approach which require high expertise in liver surgery [31] could be suitable in treatment of small foci of Echinococcus multilocularis. But in advanced cases with parasites dissemination laparatomy used to be the technique of choice.

In the described above case the portal hypertension probably could be a result of incomplete surgery due to extended parasitic infection and liver anatomical changes due to performed procedures, because the portal hypertension and it’s further complications (thrombocytopenia, splenomegaly) had not been observed before the operation. Moreover, the results of the molecular investigations suggest that the clinical outcome and after surgery complications were not correlated with parasite genotype and potential more pathogenic Echinococcus multilocularis strain.
showed in this manuscript proved that *Echinococcus multilocularis* should be taken under consideration in differential diagnosis of irregular lesions within the liver. Long-lasting invasion could be responsible for the irreversible secondary liver changes such as cirrhosis and portal hypertension. The surgery treatment (treatment of choice) is difficult and it’s results depend on the invasion period the patient is operated on. After the surgery the patient requires long life albendazole intake and careful follow – up, to detect early or late complications.

**Acknowledgements**

Not applicable.

**Authors’ contributions**

LP – write the manuscript, WC- write the surgery section, performed the operation theater, MK- performed the operation, collected the data, PZ write the molecular fragment, EL – performed molecular examination, JS – made conception of the work, analyzed the results and created a part of the discussion. All authors have read and approved the revised manuscript and all confirmed that it is the case report.

**Funding**

Not applicable.

**Availability of data and materials**

Not applicable.

**Ethics approval and consent to participate**

The patient, described in the manuscript, during admission to the University Hospital, had written the permission for diagnostics, treatment and data collections performed during the hospitalization. No other declarations were required from ethical point of view in our Poznań University of Medical Sciences, Poland.

**Consent for publication**

The written consent for the publication of the patient and identifying images was obtained from the patient. The completed written version is available in the Patient’s Medical History (Tropical and Parasitic Department of Poznań, University of Medical Sciences).

**Competing interests**

The authors declare that they have no competing interest.

**Author details**

1. Department and Clinic of Tropical and Parasitic Diseases, Poznan University of Medical Sciences, Przybyszewskiego Street 49, 60-355 Poznań, Poland.

2. Department and Clinic of General and Transplant Surgery, Poznan University of Medical Sciences, Przybyszewskiego Street 49, 60-355 Poznań, Poland.

3. Institute of Bioorganic Chemistry Polish Academy of Sciences, Z. Noskowskiego Street 12/14, 61-704 Poznań, Poland.

**Received: 2 March 2020 Accepted: 27 May 2020**

**Published online: 05 June 2020**

**References**

1. Kotva J, Isaksson M, Jardine CM, et al. *Echinococcus multilocularis* infection, southern Ontario, Canada. Emerg Infect Dis. 2019;25(2):265–72.

2. Vuitton DA, Zhou H, Bresson-Hadni S, Wang Q, Pierroux M, Raoul R, Giraudoux P. Epidemiology of alveolar echinococcosis with particular reference to China and Europa. Parasitology. 2003;127(Suppl):S87–107.

3. Nahorski W, Kracz J, Pawłowski Z, et al. Human alveolar echinococcosis in Poland: 1990–2011. PLoS Negl Trop Dis. 2011;7(11):e1986. https://doi.org/10.1371/journal.pntd.0001986.

4. Tylikowska A, Pilarczyk B, Pilarczyk R, Zysko M, Tomaż-Marciniak A. Presence of tapeworms (Cestoda) in Red Fox (Vulpes vulpes) in North-Western Poland with particular emphasis on *Echinococcus Multilocularis*. J Vet Res. 2019;63(1):71–8. https://doi.org/10.2478/jvetres-2019-0005 eCollection.2019 Mar.

5. Stefaniak J, Paul M, Kaczprzak E, Polański J, Chomicz L, Myjak P. Case definitions and clinical evaluation of patients infected with *Echinococcus multilocularis* treated in the Poznań Center. Ann Parasitol. 2004;50(3):447–54.

6. Aoki T, Hagiwara M, Yabuki H, Ito A. Unique MRI findings for differentiation of an early stage of hepatic alveolar echinococcosis. BMJ Case Rep. 2015;19:2015. https://doi.org/10.1136/bcr-2014-208123.

7. Abdulrauf AG, Milonova VB, Tserenko IA. Computed tomography diagnosis of hepatic space–occupying lesion of different nature. Khirurgia (Mosk). 2005;6(3):61–5.

8. Kern P, Wen H, Sato N, et al. WHO classification of alveolar echinococcosis: principles and application. Parasitol Int. 2006;55(Suppl):S283–7.

9. Poloñeva, et al. Use of an enzyme immunoeast test system with cystic *Echinococcus* antigen to diagnose *echinococcosi alveolaris*. Med Parasitol. 2011;2(4):44–5.

10. Liange M, Janin V, Bresson-Hadni S, Vuitton D, Houin R, Pierroux R. Immunodiagnosis of Echinococcus infections: confirmatory testing and species differentiation by a new commercial western blot. J Clin Microbiol. 2000;38:3718–21.

11. Sułma M, Szostakowska B, Nahorski W, Sikorski K, Wójc尼斯cowicz, Wąg P. The usefulness of commercially available serological tests in the diagnosis and monitoring of treatment in patients with alveolar echinococcosis. Clin Exp Hepatol. 2019;5(4):327–33.

12. Demirkazik M, Koltas IS, Inceboz T, Korkmaz M, Gumurdukl. Use of the ELISA (Em2-Em18) and western-blotting methods on diagnosis of alveolar echinococcosis. Türkiye Parazitol Derg. 2019;43(Suppl-1):13–7.

13. Stefaniak J. Guidelines for diagnosis and treatment of liver alveococcosis caused by *Echinococcus multilocularis*. Wiad Parazitol. 2007;53(1):189–94.

14. Atanasov G, et al. Alveolar echinococcosis – spreading disease challenging clinicians: a case report and literature review. World J Gastroenterol. 2013; 19(26):4527–61.

15. Dybczic M, Borkowski PK, Padzik M, Balaza W, Chomicz L. Molecular determination of suspected alveolar echinococcosis requiring surgical treatment in human cases from Poland. Ann Parasitol. 2018;64(4):339–42.

16. Nowak SP, Piekołk J, Stefaniak J. Thrombosis of inferior vena cava in the course of advanced alveolar echinococcosis. Pol Arch Int Med. 2019;129(3):204–5.

17. McNamara DP, Li Z, Yang S, Grey DJ, Yang YK. Case study emphasizing he liver transplantation as a strategy of surgical treatment with special focus on liver transplantation. Transpl Infect Dis. 2016;18(5):661–6.

18. Chomicz L, Szubert P, Fiedor P, et al. Human cystic and alveolar echinococcosis as indicator to liver transplantation. Transplant Proc. 2003;35(6):2260–1.

19. Vanista J. Clinical picture and treatment of alveolar echinococcosis. Epidemiol Mikrobiol Immunol. 1997;46(4):166–7.

20. Sieger C, Hegglin D, Schwarzenbach G, et al. Spatial and temporal aspects of urban transmission of *Echinococcus multilocularis*. Parasitology. 2002;124: 631–40.

21. Vanista J. Clinical picture and treatment of alveolar echinococcosis. Epidemiol Mikrobiol Immunol. 1997;46(4):166–7.

22. Sieger C, Hegglin D, Schwarzenbach G, et al. Spatial and temporal aspects of urban transmission of *Echinococcus multilocularis*. Parasitology. 2002;124: 631–40.

23. Vanista J. Clinical picture and treatment of alveolar echinococcosis. Epidemiol Mikrobiol Immunol. 1997;46(4):166–7.

24. Engler A, Ski R, Beer M, et al. Simple liver cysts and cystoid lesions in hepatic alveolar echinococcosis: a retrospective cohort study with Hounsfield analysis. Parasite. 2019;26:82. https://doi.org/10.1051/parasite/2019057.

25. Dirler AT, Bolukbas CS, Acici M, Umir S. Overview of Echinococcus multilocularis in Turkey and in the world. Türkiye Parazitol Derg. 2018; 43(Suppl 1):118–35.

26. Bansal N, Vij V, Restogi M, Wadkowan M, Kumar A. A report on three patient with *Echinococcus multilocularis*: lessons learned. Indian Gastroenterol. 2018; 37(4):385–8.

27. Niu F, Chong S, Qin M, Li S, Wei P, Zhao J. Mechanism of fibrosis induced by *Echinococcus* spp. Diseases. 2019;7(3):E51. https://doi.org/10.3390/ diseases/030051.
interaction. PLoS Negl Trop Dis. 2019;13(3):e0006959. https://doi.org/10.1371/journal.pntd.0006959.

29. Kowalczyk M, Kurpiewski W, Zieliński E, et al. A rare case of the simultaneous location of Echinococcus multilocularis in the liver and the head of the pancreas case report and review of literature. BMC Infect Dis. 2019;19(1):66. https://doi.org/10.1186/s12879-019-4274-y.

30. Akbulut S, Cicek E, Udu M, Sahin TT, Yilmaz S. Associating liver partition an portal vein ligation for staged hepatectomy for extensive alveolar echinococcosis first case report in the literature. World J Gastrointestinal Surg. 2018;10(1):1–5.

31. Ruzzente A, Alamo L, Conci S, Bagante F, Compagnoro T, Ciangherotti A, Guglierri A. Laparoscopic treatment of Caroli disease. Laparosc Surg. 2020. https://doi.org/10.21037/js.2019.11.01.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.