Case report

Simple schwannomatosis or an incomplete Coffin-Siris? Report of a particular case

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1. Introduction

When we speak about neurofibromatosis (NF), we mean a group of autosomal dominant hereditary diseases linked to the development of tumors of nervous and other organ systems. The three principal entities are NF1, NF2 and schwannomatosis. This last rare condition has been recognized recently and mutation of SMARCB1 gene has been found in 40–50% of familiar forms [1]; nevertheless, mutation of this gene has been found in Coffin-Siris syndrome (CSS), as well. Clinically, schwannomatosis is linked to the appearance of peripheral schwannomas, in absence of other organ involvements. We present the case of a patient with a schwannomatosis and Sjogren syndrome who has developed a spontaneous spleen rupture.

2. Case description

During July of 2013, a 28 years old female patient with an history of Sjogren syndrome, celiac disease and a surgically treated schwannoma, presented to our observation in July 2013 for a pain on the left elbow, where a tumefation was present. After neuroradiological evaluations, a surgical resection was performed and a schwannoma was diagnosed. Genetic exams revealed a puntiform SMARCB1 gene mutation. During 2015, she was subdued to the removal of an another schwannoma located into the cervical medullary canal. Few months later, she was operated in an another hospital for a spontaneous spleen rupture in a possible context of wandering spleen.

Conclusion: We think that the patient could suffer from a partially expressed Coffin-Siris syndrome. No cases of spontaneous rupture in a context of wandering spleen have been ever described as for as schwannomatosis or Coffin-Siris syndrome are concerned. More cases are necessary to establish a direct relationship.
 schwannoma. No neurological deficits appeared after intervention. Postoperative course was regular and patient was discharged one week later. In March 2016 she was admitted to the Emergency of another Hospital for the sudden appearance of an acute abdomen. No trauma history was reported. An urgent CT-scan revealed an hemoperitoneum with a spleen torsion likely. So, an urgent laparotomy was performed and a wandering, congested and ruptured spleen was found and removed. During postoperative period she was subdue to the vaccinations against S. Pneumoniae, N. Meningitidis and H. Influenzae and she was discharged two weeks later. Nowadays, she is in good conditions and no other new lesions suspected for schwannomas have been found.

3. Discussion

Schwannomatosis is the most recent found form of neurofibromatosis, characterized by the development of schwannomas and chronic pain, with two distinct forms of pathology. Schwannomatosis 1 is related to mutation of SWI/SNF-related matrix-associated actin-dependent regulator of chromatin, subfamily B, member 1, so-called SMARCB1, gene; on the contrary schwannomatosis 2 is linked to the germ-line inactivating mutation of leucine zipper-like transcriptional regulator 1, so-called LZTR1, gene [2]. SMARCB1 is considered an oncosuppressor gene because it permits the synthesis of a protein that regulates the transcription of p16INK4a and p21 and represses cyclin D1 [3]. In fact, its downregulation has been described as related to the formation of pediatric chordoma [3], while an epigenetic regulation by some miRNAs has been found in soft tissue sarcomas [4]. Moreover, some missense germ-line mutations of this gene have been related to CSS and to the possibility of malignant tumors [5].

We have described the case of a female patient with an history of multiple peripheral lesions. Two of them were operated and they revealed to be schwannomas. During follow-up radiological evaluation no sign of malignancy was found. So, it seems that her neoplastic history is limited to the presence of peripheral nervous system schwannoma. The particular attitude of her story is that in March 2016 she suffered from an acute abdomen due to an hemoperitoneum in absence of trauma history. During preoperative radiological evaluations, a spleen rupture in a context of hilum torsion was suspected and surgical intervention confirmed those suspects. Moreover, spleen was dislocated lower than its original position. This condition, that predisposes to organ infarction and rupture, is noted as wandering spleen. It is a rare condition, with an incidence of less than 0,2% among all cases of splenectomy [6], due to an absence or underdevelopment of spleen ligaments: gastrosplenic, colicosplenic, phrenolic and phrenosplenic (splenorenal) ligament [7]. Classically, this condition is associated with multiparity and abdominal wall weakness [8] and can cause the torsion of spleen pedicle with a various degree. Possible surgical treatment consist of splenectomy and splenopexy [9].

Our patient did not have any pregnancy and she did not suffer from excessive thinness. Nevertheless, her history was remarkable for a Sjogren syndrome. It is a connective tissue disease, but in literature no case of wandering spleen related to this syndrome were reported. On the other hand, the fact that SMARCB1 gene mutations can be related to CSS cannot be forgotten. This syndrome is characterized by ectodermal anomaly that lead to hypo/aplasia of fifth phalanges and a certain degree of intellectual disability, as hypertrichosis/birusting and development delay. Moreover, organ anomalies, such as cardiac congenital defects and spinal anomalies can be found. Our patient did not have ectodermal anomalies described previously, but she brought us an echocardiography reporting a mild tricuspid, mitral and aortic valvular insufficiency, without clinical signs of congestive heart failure [10]. So, we don’t have sufficient elements to formulate a diagnosis of Coffin-Siris syndrome but we can speculate that a less severe variant of this entity has occurred in our patient. Diets and colleagues, in their recent work, have described four cases of missense mutation in exon 2, resulting in the substitution of arginine with histidine. Clinically, this last finding has been seen related with a more severe form of syndrome, characterized by choroid plexus hyperplasia and hydrocephalus [11]. Our patient reported the mutation c.1120C > T (p.R374W) in exon 9, with arginine replaced by triptophane. Usually, in this exon, mutations related with a full-expressed Coffin-Siris syndrome were reported. Nevertheless, arginine in position 374 was replaced by glutamine [11]. To our knowledge, this type of mutation has never been described. For these reason, we can argue that patient could have a CSS without full phenotype expression, only with weakness of connective tissue. This could explain the presence of valvular insufficiency and wandering spleen, while the birth of peripheral schwannomas could be an acquired condition. In fact, some studied showed how a multiple hit mechanism is necessary to generate schwannomas [12]. A similar situation has been already described by Gossai et al. in 2015 [5] but, on the contrary, their patient had a frank Coffin-Siris syndrome with ectodermal anomalies and intellectual disability.

Fig. 1. MRI studies representing cervical (A) and elbow (B) schwannomas.
In conclusion, we have described the case of a female patient with schwannomatosis and a spontaneous spleen rupture due to a wandering spleen, with a mutation of SMARCB1 gene. To our knowledge, this is the first case described with a similar behavior and we think that it is related to a Coffin-Siris syndrome not fully expressed, probably caused by the particular punctiform mutation of the gene, but more cases are necessary to establish a tight relationship.

4. Conclusion

We described the case of a patient with a schwannomatosis, a spontaneous spleen rupture and a mutation of SMARCB1 gene. In consideration of her clinical history, we think that she could suffer from a partially expressed Coffin-Siris syndrome. No cases of spontaneous rupture in a context of wandering spleen have been ever described as for as schwannomatosis or Coffin-Siris syndrome are concerned. More cases are necessary to establish a direct relationship.

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References


