Madelung’s disease – progressive, excessive, and symmetrical deposition of adipose tissue in the subcutaneous layer: case report and literature review

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Abstract: Madelung’s disease is a rare disorder described for the first time in the year 1846 by Brodie. It is characterized by the occurrence of progressive, excessive, and symmetrical deposits of adipose tissue in the subcutaneous layer. Most often, these changes concern the neck, the nape of the neck, arms, and upper back, giving the patients a specific, pseudoathletic appearance. Madelung’s disease is also known as multiple symmetrical lipomatosis, benign symmetrical lipomatosis, and Launois-Bensaude syndrome. The most commonly affected ones are men who drink alcohol in excessive amounts. The fat masses emerging in the course of the disease are painless but can lead to adverse repercussions. Patients may experience dysphagia, dysphonia, difficulty in breathing, and limited mobility of the neck. The reasons for the willingness to take up treatment are also often esthetic reasons. The disease is usually accompanied by numerous metabolic disorders. The etiology of the disease has not been sufficiently explained so far, which creates diagnostic and therapeutic difficulties. Currently used treatment is limited to surgical resection of the resulting lesions or liposuction. Unfortunately, the effectiveness of these activities is limited. Most patients experience recurrence after treatment. This paper discusses the essence of Madelung’s disease, numerous aspects of etiology, the manner of diagnosis, and treatment based on current literature data.

Keywords: Madelung’s disease, lipomatosis, multiple symmetrical

Introduction
Madelung’s disease is also known as multiple symmetric lipomatosis (MSL), benign symmetric lipomatosis (BSL), and Launois-Bensaude syndrome.1–3 It is a rare disorder characterized by progressive, excessive, and symmetrical accumulation of adipose tissue in the subcutaneous layer, thus giving patients a peculiar appearance.1,2,4,5 Enzi6 distinguishes two types of lipomatosis (Figure 1).

Another classification distinguishes three types of disease.7 The first one is characterized by the dominant cervical localization of excess fat; for the second type, the pseudoathletic appearance is considered characteristic, while in the third type – gynecological – changes are located in the area of the abdomen and hips.7,8 Regardless of the type of the disease, there is rarely any accumulation of fat around the larynx, mediastinum, and distal limbs.9,10 The described changes are painless, but they can lead to unfavorable, often-serious repercussions.5,10 Patients may develop dysphagia, dysphonia, and sleep apnea syndrome. There may be dif-
difficulties in breathing and limited mobility of the neck. In patients, polyneuropathy is often observed. It is a generalized motor, sensory, and autonomic neuropathy not directly related to the location of fat mass.\textsuperscript{1,2,5,11} Another indirect consequence of the disease is the occurrence of depressive disorders in patients, resulting from the way they perceive themselves against the background of emerging changes in appearance.\textsuperscript{7}

Madelung’s disease affects men more often – the ratio of men to women varies from 15/1 to 30/1 – and is usually diagnosed between 30 and 70 years of age.\textsuperscript{1,4,12-14} Most cases of the disease are reported in the Mediterranean region – the incidence in Italy has been reported to be as high as 1 in 25,000 men.\textsuperscript{1,4,5,9,15} In contrast, in the Asian population, this disease is extremely rare.\textsuperscript{1,9} In the majority of cases, the disease is diagnosed in men in the 4th and 5th decades of life who consume chronically increased amounts of alcohol, especially in the form of red wine.\textsuperscript{5,10,16} Moreover, 90% of patients also have secondary liver cirrhosis.\textsuperscript{4,7} The most frequently diagnosed phenotype of the disease is of type 1, with a dominant fat deposit around the neck.\textsuperscript{13,14,16}

**Case report**

A 53-year-old man with upper body deformity, a strongly limited neck mobility, and weakening of the muscles of the shoulders and upper limbs came to the clinic. The patient had numerous, clearly limited, round, subcutaneous fat masses protruding above the body surface. These changes were arranged symmetrically on the neck, shoulders, upper torso, and the proximal part of the upper limbs. The patient had a pseudoathletic look, and the fatty tissue around the neck formed a typical “Madelung’s collar”. In an interview, the patient admitted that he had been abusing alcohol for 39 years. However, 9 months earlier, he was subjected to tests, on the basis of which he was diagnosed with alcohol-induced decompensated liver cirrhosis. At the time of admission, the calculated patient body mass index (BMI) was 24.7 kg/m\(^2\), with a waist circumference of 81 cm. The patient was not diagnosed with overweight or obesity, and biochemical tests did not show disturbances in either lipid or coagulation parameters. However, it was noticed that gynecomastia, limb muscle abnormalities, and hepatosplenomegaly occurred in the examined man. Laboratory tests showed the following disorders: hypochromic anemia (hemoglobin 11 g/dL, red blood cell count \(3,900 \times 10^6/\)µL), neutropenia (\(1.3 \times 10^3/\)µL), thrombocytopenia (\(100 \times 10^3/\)µL), and slightly elevated hepatic (AST: 130 U/L and ALT: 90 U/L) and pancreatic (lipase: 170 U/L and amylase: 140 U/L) enzymes. The pictures of the patient with the described deformities are shown in Figure 2.
Endocrine workup showed abnormalities. Hyperestrogenism was confirmed with normal testosterone levels and decreased dehydroepiandrosterone sulfate (DHEA-S) levels. The patient was diagnosed with type 2 diabetes based on an oral glucose tolerance test. The results of additionally tested parameters, ie, insulin, C-peptide, and thyroid hormones, were within the reference values. Finally, the patient was diagnosed with a rare disorder of fat proliferation, which is called Madelung’s disease. It was found that the disease phenotype presented by the man corresponds to type 1 according to the Enzi classification. The patient underwent computed tomography to visualize the structure of lesions and determine their location. The study also allowed to determine the degree of penetration of fat masses to deeper structures. Based on the history, clinical status, and examinations, it was decided to qualify the patient for surgical treatment. Surgical resection of excess adipose tissue was performed. In the postoperative period, there were no complications, and the patient’s well-being was assessed as good. The patient remained under observation for 3 years without recurrence.

**Discussion**

Madelung’s disease was first described in the year 1846 by Brodie and then, in 1888, it was also described by Madelung. In 1898, French physicians Launois and Bensaude published the characteristics of this disease based on a group of 65 patients. According to the data for 2012, there are >270 cases of Madelung’s disease in the literature. Chen et al summarized the clinical data of the cases described in the literature in the years 2000–2015 (Table 1).

![Figure 2 Patient with Madelung’s disease.](image-url)
Madelung’s disease remains a disorder of unexplained etiology and unclear pathogenesis. The distribution and type of adipocytes in patients with Madelung’s disease resemble the brown adipose tissue (BAT) of infants. In patients, instead of increasing the size of existing fat cells, new ones appear. The results of a long-term study, in which 7 patients were observed over a period of 12 years, confirm the hypothesis that lipomas formed in the course of the disease are the result of mitochondrial disorders of BAT. Other authors also indicate that disorders of proliferation and differentiation of human BAT cells underlie the described disease.

There are studies that reveal the relationship of Madelung’s disease with diabetes, hyperlipidemia, hypertension, or liver disease. However, the influence of excess body fat on the metabolism of the body is still under discussion. Regardless of this, people with Madelung’s disease often have coexistence of other disorders (Figure 3).

This is confirmed by the described case, in which the patient shows the occurrence of diabetes, hepatosplenomegaly, cirrhosis, and gynecomastia simultaneously with Madelung’s disease. Apart from these disorders, sensory, motor, and autonomic polyneuropathy is additionally observed in approximately 85% of patients with Madelung’s disease.

One of the main factors for the development of this disease is alcohol abuse. Alcohol adversely affects the enzymatic processes in the mitochondria and disrupts adrenergic lipolysis – it can be a cofactor inducing a change in the number and function of β-adrenergic receptors. It is likely that alcohol abuse is the cause of uncontrolled accumulation of adipose tissue in various parts of the body. The described case also presents a long history of consuming excessive amounts of alcohol. The literature describes only a few cases of Madelung’s disease without alcohol in the patient’s diet. Other possible risk factors include nicotine abuse, metabolic disorders, and malignant tumors.

Most likely, there is genetic predisposition based on matrilineal inheritance. It is recognized that the disease may be inherited in an autosomal dominant manner with variable mitochondrial DNA penetration.

Archeological discoveries suggest that there may be an inherited tendency for Madelung’s disease – perpetuated for centuries – among the population of people living in the Mediterranean. This may be evidenced by a statue known as the Capestrano warrior from the 6th century BC and discovered in Italy. It presents an impressive figure with a unique somatic aspect, characterized by large hips and broad shoulders. The silhouette of the Capestrano warrior is very similar to the appearance of a patient with Madelung’s disease. This convergence may be accidental and may also indicate the genetic basis of the occurrence of this disease.

A case report of a familial Madelung disease in siblings, ie, a 41-year-old woman and her 36-year-old brother, has been described in the literature. In molecular studies, the existence of mutations in the lipase-encoding gene has been demonstrated in both patients. In the studies carried out so far, Madelung’s disease is associated with point mutations in mitochondrial DNA, whose genes are important in the regulation of lipolytic pathway processes. A relationship between the described disease and the occurrence of myoclonic epilepsy and shred of red fibers has also been established. In the muscles of patients with Madelung’s disease, the occurrence of radiant red fibers is often found, which indicates mitochondrial dysfunction with reduced cytochrome c oxidase activity.

The local defect of catecholamine-induced lipolysis, adrenergic receptor defects, and the reduction of inducible nitric oxide synthase (iNOS) are potential causes of Madelung’s disease. The implication of these processes is increased adipogenesis. In addition, recently conducted molecular studies using next-generation sequencing have also demonstrated the existence of genetically diverse genes in patients with Madelung’s disease compared to healthy controls.

The diagnosis of Madelung’s disease is based on history, clinical appearance of the patient, and the results of imaging examinations (Figure 4).

The disease should be differentiated from morbid obesity, Cushing’s syndrome, salivary gland disease, lymphoma, Froelich’s syndrome, angiolipomatosis, myxoid liposarcoma, encapsulated lipomas, neurofibromatosis, and lipomatosis in
Madelung's disease

The disease progresses rapidly in the first few years, after which the progression usually slows down or the size of the created adipose tissue mass stabilizes.3,8

The therapeutic approach in Madelung’s disease is reduced to palliative treatment, which includes surgical resection of the adipose tissue, liposuction, or injection lipolysis.1–4 It seems reasonable to correct metabolic disorders and overweight.3 Alcohol withdrawal and weight loss may be helpful in controlling the disease, but they do not guarantee the inhibition of its progression.1,3,16 Surgical procedures are time consuming – often divided into several stages – due to the volume of fat mass, as well as their spread and infiltration into neighboring tissues.3,4 Conventional surgery is recommended, especially
in heavily vascularized and innervated regions, as well as in violation of patency of the airways and esophagus. It is estimated that nearly 63% of patients experience recurrence after surgical resection of adipose tissue, while in the case of liposuction, up to 95% of patients experience recurrence. After injection lipolysis, fibrosis and adhesions form. This impedes surgical treatment or liposuction in the recurrence of lesions. Despite frequent relapses, functional and cosmetic effects obtained in surgical treatment can be satisfactory. Unfortunately, no effective pharmacotherapy has been developed so far in the treatment of Madelung’s disease. The use of salbutamol for adrenergically stimulating lipolysis and thus inhibiting the progression of the disease in larger clinical trials did not show the expected effect.

Conclusion
Madelung’s disease – apart from the essential excessive accumulation of adipose tissue – is characterized by the occurrence of numerous metabolic disorders. In the literature, the authors describe the coexistence, among others, of carbohydrate, lipid, or hormonal disorders along with the disorder in question. Diagnosis of diabetes and abnormal levels of sex hormones in the presented case are the confirmation.

The described case also confirms the role of excessive alcohol consumption in the pathogenesis of the disease. In contrast, impaired liver function – occurring in the majority of patients with Madelung’s disease – may have a greater impact on the metabolic abnormalities than the pathology of adipose tissue alone.

Madelung’s disease is rarely diagnosed, which may not necessarily be due to the low incidence of this disease but may be associated with the diagnostic and therapeutic difficulties, as well as with the availability of few literature reports, often related to individual cases.

Consent for publication
The patient provided written informed consent for publication of the clinical details and images.

Disclosure
The authors report no conflicts of interest in this work.

References