Renal Failure due to Bardet-Biedl Syndrome

A Case Report

Sukru Ulusoy Kubra Kaynar Semih Gul Kubilay Ukinc

Department of Internal Medicine, Faculty of Medicine, Karadeniz Technical University, Trabzon, Turkey

Key Words
Bardet-Biedl syndrome - Renal failure - Hemodialysis

Abstract
Objective: To describe a case of Bardet-Biedl syndrome involving renal failure and retinal dystrophy. Case Presentation and Intervention: A 50-year-old female patient presented to the emergency service with uremic symptoms and metabolic acidosis. Polydactyly, retinitis pigmentosa, obesity, strabismus, nistagmus and renal failure were found. Because she had end-stage renal failure, hemodialysis therapy was started. She has been well for 18 months, without any complication on hemodialysis. Conclusion: Bardet-Biedl syndrome should be considered in patients with polydactyly, retinitis pigmentosa and renal failure.

Introduction

Bardet-Biedl syndrome, a form of Laurence-Moon-Biedl syndrome, has five recognized features: retinal dystrophy, obesity, dysmorphic extremities, renal abnormalities and hypogonadism (in male patients). Polydactyly, syndactyly or brachydactyly may be seen. The degree of polydactyly, a prominent feature of this syndrome, varies from patient to patient, ranging from a wide fifth metatarsal or metacarpal to a complete sixth digit [1]. Scores on tests of intelligence are usually low. Mental retardation, polydactyly and hypogonadism are not necessarily present in female patients [2]. The autosomal recessive disorder Bardet-Biedl syndrome is heterogeneous with at least four gene loci (BBS1–4): 11q13 (BBS1), 16q21 (BBS2), 3p12 (BBS3) and 15q22 (BBS4) [3]. A fifth locus [4] and a sixth locus [5] map to chromosome 2q31 and chromosome 20, respectively. These alleles may act in conjunction with mutations at other BBS loci to cause or modify the BBS phenotype. The prevalence is 1:160,000. It usually presents between 10 and 20 years of age. The ocular defect in Laurence-Moon-Biedl syndrome involves the photoreceptor cells [6]. In addition to retinitis pigmentosa, strabismus may be observed [7]. Early otolaryngologic, audiologic, speech pathology, and dental evaluation of these individuals is recommended [8]. Bifid epiglottis [8] and disturbance of tooth formation have been reported. Hypertrophy of the interventricular septum and dilated cardiomyopathy, insulin-resistant diabetes mellitus, empty sella, clindactyly, and congenital hepatic fibrosis can also be found. Although hypogonadism is rare in female patients, vaginal atresia, hypoplasia of uterus...
and ovaries, and ectopic urethra should be sought. Uremia is an important cause of morbidity and early mortality in these patients. Structural or functional abnormalities of the kidneys can be observed. The spectrum of renal involvement can range from calyceal clubbing or cysts to diffuse renal cortical loss [9]. Light microscopy may reveal varying degrees of increase in mesangial cellularity and matrix or sclerosis of the glomerular tuft [10]. Renal disease occurs in 70% of the patients [10].

We describe a patient with renal failure, who had signs and symptoms of Bardet-Biedl syndrome.

Case Report

A 50-year-old single, nulliparous woman with normal secondary sex characteristics presented to our emergency room in September 2000 with a history of nausea, vomiting and dyspnea. She had a history of frequent urinary system infections. On examination, she was febrile at 38.4°C, and had an erythematous lesion with yellowish discharge located near the vulva. Kussmaul’s respiration was observed. She had been blind since aged 40. Ophthalmoscopic examination revealed bilateral retinal pigmentation and strabismus. There were bilateral scars due to amputation of a sixth toe on both feet. There had been postmenopausal for 2 years. Her thyroid and sex hormone levels were normal. After 3 weeks of hemodialysis she was well. The patient had diffuse cortical loss. This has also been reported [16].

Discussion

In this condition death generally occurs at a considerably younger age than in the general population with renal disease frequently noted as the primary cause of or the contributing cause of death. Renal disease seems to reduce life expectancy considerably [11]. Renal transplantation appears to have a good outcome in these patients. However, after renal transplantation, morbid obesity can be observed [12]. Peritoneal dialysis has been attempted in these patients but due to infections and the hepatic involvement of the disease, frequent conversion to hemodialysis is required [13] as in this patient.

Genetic consulting should be provided for those who want to get married and be pregnant. Prenatal diagnosis may be made by ultrasound when large echogenic kidneys (which can mimic infantile polycystic kidneys) and polydactyly may be found [14, 15]. In a prospective cohort study; 38 patients with this syndrome were identified and studied. Of these, 86% were blind, 26% being blind by the age of 13 years [16]. Our patient was blind by the age of 10. Whereas fetal lobulation and calyceal cysts/diverticula/clubbing are characteristic in this syndrome, our patient had diffuse cortical loss. This has also been reported in this syndrome [16].

Conclusion

This report illustrates the importance of the clinician considering the diagnosis of Bardet-Biedl syndrome in adults with retinal dystrophy, polydactyly and renal failure.

References


