Familial hypercholesterolemia with early coronary atherosclerotic heart disease: A case report

YANHONG LUO1, MIN ZHENG2, YUTIN ZHANG3, SHUANG HE2, LEI ZHANG2, HUICHAO SUN2, XIAOYAN LIU2, TAO TAN3, HUA ZHU3 and JIANFENG HE2

Departments of 1Endocrinology and Genetic Metabolism and 2Cardiology, Children's Hospital Chongqing Medical University, Chongqing 400014, P.R. China; 3Department of Surgery, Davis Heart and Lung Research Institute, The Ohio State University Wexner Medical Center, Columbus, OH 43210, USA

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Correspondence to: Dr Jianfeng He, Department of Cardiology, Children's Hospital Chongqing Medical University, 136 Zhongshan Er Road, Chongqing 400014, P.R. China
E-mail: 1534568197@qq.com

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Abstract. Patients with familial hypercholesterolemia usually present with high levels of serum low-density lipoprotein, xanthomas and early coronary artery disease. A 13 years old female patient was admitted to Children's Hospital of Chongqing Medical University presenting symptoms of heart failure. Laboratory tests showed that her cholesterol and low-density lipid levels were extremely high. Electrocardiogram test revealed that she had sinus tachycardia, QT lengthening and ST-T change. Multiple cardiac function abnormalities were diagnosed by echocardiogram. Multiple coronary artery stenosis was determined by computed tomography angiography. After the combination of lipid lowering, anti-thrombosis, and cardiac remodeling therapies, the patient's symptoms were significantly improved and the patient was discharged.

Introduction

Familial hypercholesterolaemia (FH) is a rare genetic disorder, which affects around 0.2% of the population (1). The disease manifestations are very high levels of serum low-density lipoprotein (LDL), xanthomas and early coronary artery disease (CAD). The genetic mutations of lipid handling genes in patients lead to hyperlipidemia that usually develop into severe atherosclerosis and cause ischemia heart diseases and heart failure. Due to technical challenging and resource limitations, genetic screening is not possible to diagnose patients with FH, especially in developing countries. Thus, establishment of an easy and low cost diagnostic approach for FH is needed.

In the present study, we reported a FH patient admitted in our hospital with symptoms of heart failure. With physical and laboratory tests, the patient showed typical FH symptoms. Cardiac dysfunctions and coronary artery stenosis were further determined by electrocardiogram, echocardiogram and CT angiography. The patient was immediately treated with combination of lipid lowering, anti-thrombosis and cardiac remodel improving therapies. The patient was discharged when the heart failure symptoms were successfully controlled.

Case report

A 13 years old female patient was admitted to Children's Hospital of Chongqing Medical University (Chongqing, China) due to symptoms of heart failure, including lower limb swelling for 9 days, coughing and panic for 6 days. Upon arriving hospital, the patient felt heart palpitations, breath shortness, and fatigue after short walk. The lower extremity of the patient had pitting edema, which had been aggravated with oliguria and sweating for 9 days. In the course of the disease, there was a paroxysmal cough accompanied with the yellow mucus phlegm and bloody phlegm. None of the following symptoms was presented: wheezing and dyspnea, eyelid edema, gross hematuria, foam urine, chest tightness, chest pain, nausea, vomiting, abdominal distension, diarrhea, dizziness, headache, syncope, or fever. Based on her disease history, dermal yellow nodules were found right after her birth, they developed into nodular xanthoma during her growth.

Upon admitted in our hospital, physical examination was performed: Her body temperature was 36.6°C, breath was 30 BPM, heart rate was 142 BPM, and blood pressures were 110/58 mmHg. The skin of double knees, left elbow, ankle, wrist, toe and coccygeal joint had hemispherical yellow eminence subcutaneous nodules (Fig. 1). The size of nodules ranged from 1.0x2.0 to 5.0x6.0 cm. The boundary was clear, texture was tough, and the surface was free from swelling and ulceration. The apex beat was located at the 5th intercostal space about 3.0 cm from left midclavicular line. No presentation of lifting type pulsation and pericardial friction was detected. The heart beat sounded slightly dull and arrhythmia could be detected.
Laboratory tests were performed: blood lipids were extremely high [total cholesterol (TC) 14.73 mmol/l, high-density lipoprotein (HDL) 0.59 mmol/l, LDL 13.44 mmol/l], B type natriuretic peptide was 800 pg/ml, lactic acid was 4.04 mmol/l, blood amine was 61.7 mol/l, CRP was 59 mg/l, PTC was 1.138 ng/ml. In addition, the following tests were performed and showed normal: Blood routine, urine routine, fecal routine, liver and kidney function, electrolytes, myocardial markers, ESR, immunization, autoantibodies, blood glucose and blood gas tandem mass spectrometry.

Because the patients showed symptoms of heart failure, cardiac related tests were performed. First, electrocardiogram of the patient showed sinus tachycardia, QT lengthening, ST-T change (Fig. 2). Dynamic electrocardiogram test revealed that ventricular premature beat presented 2 times/24 h, atrial premature beat presented onceper 24 h, the average heart rate was 97 beats per minute, and the heart rate variability was poor. Furthermore, cardiac morphology and function were determined by echocardiography (Fig. 3). Multiple pathological changes have been observed, including enlarged left ventricle, enlargement of left and right atrium, and right ventricle, decrease of ventricular septum and left ventricular posterior wall motion, and aortic sinus tube junction stenosis with hyperechoic structure with moderate regurgitation, mild to moderate tricuspid valve regurgitation, decreased left ventricular systolic function, slightly widened inferior vena cava. In addition, abdominal B-ultrasound test revealed a slight swelling of the liver. In order to examine the cause of ischemic heart disease, chest CT angiography was performed (Fig. 4): There were multiple points of stenosis in the ascending aorta, left and right coronary artery and descending aorta, while CT of double lungs showed normal. The bifurcation of the left coronary artery was stenosis, the degree of stenosis was 93%, the left anterior descending coronary artery had stenosis after diagonal branches, the degree of stenosis was 68%, the upper right coronary artery (from the start 27.9 mm) also had stenosis, the degree of stenosis was 42%.

Except for the patients, no skin xanthoma and premature coronary heart disease were found in the family.

Based on the patient's medical history, clinical manifestation, blood lipid and cardiac test results. Diagnosis was made: the patient showed typical FH with coronary atherosclerotic heart disease. She had grad II cardiac insufficiency, nodular xanthoma, atherosclerosis, and vavular calcification. In order to confirm our diagnosis, a genetic test was prescribed to the patient. Unfortunately, the parents of the patient refused to perform the genetic test due to the financial difficulties.

Based on guidance from the 2014 European Atherosclerotic Association and the 2016 British HoFH management, the following treatments were prescribed to the patient: Sodium phosphocreatine 1.0 g IV, qd, 11 days, VitC 3.0 g IV, qd, 11 days, combined coenzyme needle 1 branches IV, qd, 11 days to nourish the myocardium. Milrinone 50 µg/kg, slow intravenous injection for 10 min later, 0.5 µg/kg/min, 6 days, and maintain dobutamine 5 µg/kg/min IV continued 2 h, q8 h, 5 days, digoxin 0.125 mg/time, po, qd for 6 days were used to enhance myocardial contractility. Oral Atorvastatin 20 mg/time, qN for 6 days and diet control were used to lower blood lipid. Ezetimibe Tablets: 10 mg/time, po, bid for 6 days were administrated to stabilize plaque. Benazepril 10 mg/time, po, qd for 5 days and metoprolol 6.25 mg/time, po, bid for 2 days were used to improve myocardial remodeling. Spironolactone 20 mg/time, po, bid for 11 days, furosemide 20 mg/time, po, bid for 10 days and hydrochlorothiazide 25 mg/time, po, bid for 1 day, diuresis. Aspirin 100 mg/time, po, qd for 3 days and clopidogrel 50 mg, po, qd for 3 days were used to prevent thrombosis formation. After treatment for 11 days, the symptoms of the patient were improved, she was discharged.

After returning home, her symptom has been improved. The drugs were subscripted to her after discharge: Oral drug spironolactone 10 mg/time qd for one year, hydrochlorothiazide 12.5 mg/times, qd for one year, metoprolol 6.25 mg/time, bid for one year, hydrogen chloropidogrel 25 mg/time, qd
Figure 2. Electrocardiogram shows sinus tachycardia, QT lengthening, ST-T change.

Figure 3. Echocardiogram pictures show reduced cardiac functions of the patient. (A) The left ventricle was significantly increased and the left ventricular systolic function decreased (EF 33.6%, FS 16.5%). (B) Left ventricular enlargement, left atrial enlargement and moderate reflux of aortic valve (white arrow). (C) The left ventricle was obviously enlarged, the left atrium increased, and the mitral regurgitation was moderate and severe (the white arrow indicates the mitral regurgitation). (D) Stenosis of the junction of the aortic sinus with hyperechoic structure (white arrow).
for one year, atorvastatin 20 mg/time, qN for one year. After 1 year of treatment, the patient was retested for blood lipids (TC 10.03 mmol/l, HDL 0.48 mmol/l, LDL 852 mmol/l), function of liver and kidney, electrolytes, blood glucose, blood coagulation, myocardial damage markers were normal, ECG was sinus tachycardia, and ST segment was low.

Ethical approval for the present study was obtained from Children's Hospital of Chongqing Medical University and oral informed consent was obtained from both the patient and the patient's parent.

Discussion

FH is an autosomal dominant genetic disease. The most common cause of FH is the mutations of the genes encoding lipid handling proteins, such as LDL receptor (LDLR), apolipoprotein B (ApoB) or proteins convertase subtilisin/Kexin/kexin9 (PCSK 9), resulting in LDL metabolism defects, and LDL cholesterol (LDL-C) levels are abnormally elevated in plasma. The FH patients usually have xanthoma in peripheral tissue, atherosclerosis and other clinical manifestations. Based on a FH study in Jiangsu, China, the Chinese FH diagnostic standard was established (2) (summarized in Table 1). Based on this standard, the patient in current study met multiple criteria of FH, including an early-onset coronary heart disease history and extremely high level of LDL-C (13.44 mmol/l), and xanthoma formation at the tendon sites. FH can further be divided into homozygote type (HoFH) and heterozygote type (HeFH), where the HoFH is a serious rare disease (3). The prevalence of HoFH was 1/(16-100)x10^6, and the patients' serum LDL-C >13 mmol/l. If without any treatment, the HoFH patients usually die from coronary heart disease before the age of 30 (4). The incidence of coronary heart disease in patients with HoFH is 100 times higher than that of the normal population, and most of the patients with HoFH are able to develop in childhood and adolescence (5).

The main clinical manifestations of HoFH are plasma LDL-C level is extremely high (approximately 6-8 times higher than normal), corneal arch, xanthoma at the tendon sites, premature coronary heart disease and aortic valve disease (6). Aortic valve disease is mainly caused by the involvement of the root of the aorta, which leads to the stenosis and calcification of the aorta on the upper part of the aortic valve and the aortic valve, and even the opening of the coronary artery (7-9). Except absent of corneal arch, the patients displayed all other symptoms.

The 2014 European Atherosclerotic Association (European Atherosclerosis Society, EAS) HoFH management guide proposed that the diagnosis of HoFH should be based on the standard of genetic diagnosis or in conformity with the clinical diagnostic criteria. The diagnostic genes include LDLR, ApoB, PCSK9, and LDLRAP1. The clinical diagnostic criteria are: LDL-C>13 mmol/l before treatment or LDL-C>8 mmol/l after treatment, and the presence of xanthoma in the tendon at the age of 10 or the level of parental LDL-C was consistent with heterozygote FH (HeFH) (3). The 2016 British HoFH management guide also recommends the use of genetic diagnostic or clinical diagnostic criteria, in which the genetic diagnostic criteria are basically the same as the EAS HoFH management guidelines. Children and adults diagnostic criteria are: Children LDL C>11 mmol/l and skin tendon xanthoma before 10 years of age; adult LDL-C>13 mmol/l and there were obvious xanthoma of skin tendon, or LDL-C level reached clinical diagnosis standard, while parents were diagnosed as HeFH (8). Therefore, we diagnosed the patient in this study as HoFH.

Finally, after literature search, we found one previous study that reported a 17 years old female patient. She had hypercholesterolemia on 2 year old, xanthoma at age of 9, and the clinical manifestations of coronary heart disease at age of 12. In addition, her TC and LDL-C was significantly increased, the electrocardiogram has ST segment depression, cardiac color Doppler showed stenosis and mitral regurgitation. There was sign of heart failure. The aorta stenosis and coronary artery stenosis were detected by angiography. The genetic examination revealed that she has exon 541T>C homozygous mutation of LDLR gene (9). Based on the clinical manifestation, we believe our results are in agreement with those reported by Zhao et al (9).

Thus, we showed the feasibility of a clinical diagnosis of HoFH without a genetic diagnosis. We believe this is extremely importantly in developing countries, where genetic diagnosis might be challenging and not available. In this case, we had successfully used lipid-lowering therapy, anti-thrombosis...
therapy and cardiac remodeling therapy to improve cardiac function of the patient.

According to the clinical diagnostic criteria, only about 1/4 of FH cases can be predicted. Most patients were diagnosed until middle age, thus missing the chance of early treatments. It has been estimated that there are about 26 million potential FH patients in China (10), but the rate of diagnosis and treatment in clinics is still very low (11). According to the international FH foundation, population screening should be based on age, sex and specific LDL-C level. Systematic screening strategy is very important for identifying FH proband, which is the key to discover new cases for family members (12). Therefore, we should perform blood lipids, electrocardiogram, echocardiography and carotid ultrasound examination on parents, siblings and other family members, and complete family genetic testing, in order to identify FH patients and perform early treatment.

The current treatment options for FH include lifestyle changes, drug therapy and liver transplantation. Lifestyle changes are the most important aspect of lipid-lowering therapy, including reducing uptake of saturated fatty acids and cholesterol, eating cellulose rich foods, stopping smoking and controlling salt consumption, proper physical activity, and weight loss. Conventional lipid-lowering drugs include statins and ezetimibe. The National Lipid Association of the United States suggested that the level of LDL-C in patients with FH decreased to below 2.5 mmol/l or decreased by >50% compared with that before treatment (12). But HoFH is a serious hereditary disease. Even when multiple drugs combined with lifestyle intervention, it is still difficult to achieve LDL-C compliance (13). The EAS HoFH management guide recommends that HoFH patients take high-intensity tolerable dose statins as initial treatment, gradually combine with other types of lipid-lowering drugs such as ezembo, and PCSK9 inhibitor, or plasma lipoprotein replacement, in addition, some patients may choose to undergo liver transplantation (14). In recent years, with the in-depth study of the mechanism of cholesterol metabolism, new target drugs are emerging. The most representative new drugs are mainly two oligonucleotide inhibitors ( mipomersen and lomitapide) (15) and two PCSK9 monoclonal antibody inhibitors ( alirocumab and evolocumab) (16). Mipomersen and lomitapide can significantly reduce blood lipids in patients with HoFH, but the cost is high and the incidence of adverse reactions is high. In contrast, PCSK9 inhibitors are better choices at present because of their therapeutic effects and high potency ratio.

Clearly, our case report has limitations. One of them was there was no genetic test for the patient to finally confirm the case was a HoFH at the gene level. Luckily, some clinical evidences strongly suggested it was a HoFH case. First, the immediate family members are all normal without known skin xanthoma, indicating they might be heterozygous carriers. Second, the symptoms of the patients are consistent to the diagnostic criteria from both the 2014 European Atherosclerotic Association and the 2016 British HoFH management guidance. Most importantly, after following treatment of HoFH, the patient has recovered and was finally discharged. Thus, we feel our case is deserved to be achieved and valuable for the physicians from developing countries, where the genetic tests might not be feasible. The other limitations include that we do not have information for the family history of coronary heart disease neither did we check the patient's second-degree family members for the symptom. We failed to obtain this important information due to quick improvement of the patient's symptom and discharge of the patient.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

YL, TT, HZ and JH participated in the manuscript writing; YL, TT, MZ, YZ, SH and LZ participated in data collection; XL participated in manuscript editing and data analysis; HZ and JH participated in the data analysis and manuscript revision. All authors have read and approved the manuscript.

Ethics approval and consent to participate

The study has been approved by IRB of Children's Hospital of Chongqing Medical University.
Patient consent for publication

The patient provided oral consent for publication of this study.

Competing interests

The authors declare that they have no competing interests.

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