Lomitapide treatment in a female with homozygous familial hypercholesterolaemia: a case report

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Background
Homozygous familial hypercholesterolaemia (FH) is an autosomal-dominant inherited disease presenting with highly elevated low-density lipoprotein cholesterol (LDL-C) levels. Untreated, the patient can develop atherosclerosis and cardiovascular disease already in adolescence. Treatment with statins and ezetimibe is usually not sufficient and LDL apheresis is often required. Lomitapide, an inhibitor of the microsomal triglyceride transfer protein, reduces LDL-C and triglyceride levels and can be used alone or in combination with other therapies in homozygous FH. However, experience with this agent is still limited.

Case summary
We present a young female who was diagnosed with homozygous FH at 6 years of age. She shows a complete lack of normal LDL receptor activity and no cholesterol-lowering effect from statins. The patient was treated with LDL apheresis from 7 years of age. When LDL apheresis treatment extended to twice a week, she began to experience adverse effects, including catheter-related complications, infections, and hospital admissions. When lomitapide treatment was initiated, the frequency of apheresis reduced, the LDL-C levels improved and she has not had any further hospital admissions since. Initially, she suffered from gastrointestinal disturbances. However, after 3 years of treatment with lomitapide 20 mg/day, the patient has not experienced any adverse effects.

Discussion
In this female with homozygous FH adding lomitapide treatment to LDL apheresis has contributed to improved LDL-C levels, a reduction in LDL apheresis sessions and enhanced quality of life. No adverse effects have been reported. These findings suggest that lomitapide can be a drug of choice in patients with homozygous FH.

Keywords
Homozygous familial hypercholesterolaemia • Lomitapide • Case report • LDL apheresis • Hypercholesterolaemia treatment

Learning points
• Homozygous familial hypercholesterolaemia (FH) is a rare autosomal-dominant inherited disease presenting with highly elevated low-density lipoprotein cholesterol (LDL-C) levels.
• Frequent administration of LDL apheresis is often required to lower the LDL-C levels, and if untreated, the patient can develop atherosclerosis and cardiovascular disease already in adolescence.
• Lomitapide is an inhibitor of microsomal triglyceride transfer protein; its mechanism of action is to decrease plasma LDL-C and triglyceride levels. The drug is approved for the treatment of homozygous FH.
Case presentation

The patient is a female born in 1992 who was diagnosed with homozygous FH in 1998 at 6 years of age. At diagnosis, xanthomas were noted on the Achilles tendons and on the skin of one hand (Figure 1), total cholesterol was 20 mmol/L and LDL-C (calculated with Friedewald’s formula) was 18.5 mmol/L. Carotid ultrasound showed increased intima-media thickness, whereas echocardiography and a stress echocardiography were normal. A homozygous mutation (W556R) was identified in the LDL receptor gene, resulting in less than 5% of normal LDL receptor activity in her cultured fibroblasts.12,13

Treatment with atorvastatin and ezetimibe was initiated at diagnosis but terminated because there was no effect on cholesterol levels. At 7 years of age, the patient received weekly LDL apheresis treatment and, in combination with dietary intervention, her LDL-C levels went from 10–12 mmol/L before to 3–4 mmol/L after apheresis therapy. From 13 years of age, the frequency of LDL apheresis treatment had to be periodically increased and from 21 years of age, LDL apheresis was performed regularly twice a week.

During the biweekly LDL apheresis treatment, the patient encountered frequent complications from the dialysis catheter, with bleedings, repeated infections and septicemia, resulting in several hospital admissions and catheter changes. Because of these complications, additional therapy was considered and in the autumn of 2015 treatment with lomitapide 5 mg/day was initiated. The dose was slowly escalated, alternating between 20 and 30 mg/day because of gastrointestinal symptoms (stomach pain, diarrhoea, and nausea). In 2017, the lomitapide dose was uptitrated from 20–30 mg to 40 mg/day and now well-tolerated, which was probably the result of better dietary habits. Because of limited financial support from the city council, the dose was again decreased to 20 mg/day in 2018 (Figure 2).

The initiation of lomitapide treatment rendered a reduction in the frequency of apheresis while still maintaining acceptable LDL-C levels.
Lomitapide treatment in a female with homozygous FH

(Figure 3). Because of fewer apheresis treatments, infections and hospital admissions caused by catheter complications were eliminated. In 2016, the patient was temporarily without lomitapide for 2 months and the frequency of LDL apheresis therapy had to be increased to once a week during that period (Figure 2).

The presence of elevated transaminases, induced hepatic steatosis and the development of fibrosis have been reported as adverse effects of lomitapide treatment.3,14 In this patient, alanine transaminase and aspartate transaminase were slightly increased during the dose of 40 mg/day but almost normalized once the dose was decreased to 20 mg/day (Figure 4). A liver biopsy performed in 2009 showed mild fibrosis (Stage 1). The appearance of mild hepatic steatosis on magnetic resonance tomography was noted in 2015, just before lomitapide was initiated and, after 2 years of treatment, a new examination revealed slightly increased hepatic steatosis.

In June 2018, transient elastography, an ultrasound-based method using low-frequency elastic waves to estimate hepatic steatosis and liver fibrosis,15 was performed by an experienced investigator. At that time, the patient was on lomitapide 20 mg/day. The examination indicated a low grade of steatosis (controlled attenuation parameter 247 dB/m) but was suggestive of advanced fibrosis (19.3 kPa, interquartile range 2.5 kPa, success rate 100%) and therefore a second liver biopsy was performed 3 months later. The biopsy showed low-grade microvesicular steatosis and fibrosis Stage 1 without inflammation or signs of steatohepatitis (Figure 5A and 8), providing no evidence to support the notion that lomitapide had induced any serious adverse liver effects.

In 2015, the patient was investigated by a cardiologist because of periods of dyspnoea at exertion and bursts of chest pain. An echocardiogram revealed a tricuspid aortic valve with normal opening, mild scattered increased echogenicity, and mild to moderate central aortic insufficiency. During stress echocardiography, there were no segments with hypokinesia, although the apical segments had a late ventricular contraction pattern (tardokinesia). No obstructive coronary artery disease was present at a coronary angiography performed at the same time.

In 2018, a coronary computed tomography angiography was conducted for clinical evaluation and assessment before a possible pregnancy. Still, no obstructive coronary artery disease was present, but a
calcified plaque ostially in the right coronary artery and the thoracic aorta was noted (Figure 6A and B).

Since her diagnosis, the patient has been on a fat-restricted diet and has had normal growth and development. Her body mass index (BMI) just before lomitapide initiation was 28.2 kg/m²; after 2 years of treatment, her BMI was 26.9 kg/m². She is not participating in any regular physical activities, is not smoking and does not consume alcohol. Serum levels of alpha-tocopherol and retinol have been normal. Vitamin D deficiency occurred periodically and this condition was treated with supplements accordingly.

Discussion
The effect of lomitapide treatment in a young female with homozygous FH and a rare mutation resulting in no functional LDL receptor activity is presented in this case report. Before lomitapide treatment, the patient experienced frequent complications and emotional and social impairment from the biweekly LDL apheresis therapy. The lomitapide treatment reduced the frequency of LDL apheresis while still maintaining acceptable LDL-C levels. The patient had no serious risks or side effects of the new treatment. Moreover, hospital
admissions associated with side effects of frequent LDL apheresis were eliminated and her quality of life improved steadily. Finally, considering the diagnosis of homozygous FH, her cardiovascular status at the age of 26 years is surprisingly normal.

Hepatic adverse effects have been reported in patients treated with lomitapide. In our patient, no progression of low-grade fibrosis occurred during 9 years of follow-up, including 3 years with lomitapide treatment. We therefore conclude that lomitapide did not contribute to harmful liver side effects. However, the slight increase of hepatic steatosis observed in 2018 compared with the earlier examinations could be a result of lomitapide treatment, inappropriate dietary habits, or both. We also observed that the transient elastography falsely indicated an advanced fibrosis stage, which was contradicted by the liver biopsy. Consequently, the use of elastography in patients with homozygous FH must be further evaluated for use in clinical routine.

Lomitapide treatment (20 mg/day) costs approximately 200 000 Euros/year, which is equal to the cost of 1-year biweekly LDL apheresis treatment. When adding lomitapide, the frequency of LDL apheresis decreased to every second week and the total treatment cost increased approximately 25%. Considering the improved quality of life, reduced hospital admissions and complications, we believe there is a positive net effect for all actors: patients, the wider healthcare system and society.

**Patient perspective**

Lomitapide treatment has been successful in this young woman with homozygous FH and a complete lack of functional LDL receptor activity, resulting in a reduction in LDL apheresis sessions, enhanced quality of life with few adverse events, good cardiovascular status and without clinically relevant liver side effects. Taken together, Lomitapide might be considered as a drug of choice for patients with homozygous FH.

**Lead author biography**

Karin Littmann is a MD and PhD student. She works as a specialist in Clinical Chemistry at the Karolinska University Laboratory and part time in the outpatient clinic for patients with dyslipidemias at the Endocrinology Unit, Karolinska University Hospital, Sweden. She is also a PhD student at the Department of Laboratory Medicine, Karolinska Institute, and her PhD project are focused on biomarkers for cardiometabolic diseases.

**Supplementary material**

Supplementary material is available at European Heart Journal - Case Reports online.
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