

1 **Title Page**

2 **A case report of heterozygous familial hypercholesterolaemia with *LDLR* gene mutation**
3 **complicated by premature coronary artery disease detected in primary care**

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1 Abstract

2 **Background:** Familial Hypercholesterolemia (FH) is an autosomal dominant genetic condition
3 predominantly caused by the low-density lipoprotein receptor (*LDLR*) gene mutation.

4 **Case Summary:** This is the case of a 54-year-old Malay woman with genetically confirmed FH
5 complicated by premature coronary artery disease (PCAD). She was clinically diagnosed in
6 primary care at 52 years old, fulfilling the Simon Broome Criteria (possible FH), Dutch Lipid
7 Clinic Criteria (score of 8: probable FH) and Familial Hypercholesterolemia Case Ascertainment
8 Tool (FAMCAT relative risk score of 9.51). Subsequently, she was confirmed to have a
9 heterozygous *LDLR* c.190+4A>T intron 2 pathogenic variant at the age of 53 years. She was
10 known to have hypercholesterolemia and was treated with statin since the age of 25. However,
11 the lipid-lowering agent was not intensified to achieve the recommended treatment target. The
12 delayed FH diagnosis has caused this patient to have PCAD and percutaneous coronary
13 intervention (PCI) at the age of 29 years and a second PCI at the age of 49 years. She also has a
14 very strong family history of hypercholesterolemia and PCAD, where seven out of eight of her
15 siblings were affected. Despite this, FH was not diagnosed early and cascade screening of family
16 members was not conducted, resulting in a missed opportunity to prevent PCAD.

17 **Discussion:** FH can be clinically diagnosed in primary care to identify those who may require
18 genetic testing. Multidisciplinary care focuses on improving identification, cascade screening
19 and management of FH is vital to improving prognosis and ultimately preventing PCAD.

20 **Keywords:** familial hypercholesterolemia; heterozygous; *LDLR* gene mutation; premature
21 coronary artery disease; case report; multidisciplinary management; primary care

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1 **Data Availability Statement**

2 Data are kept at the Department of Primary Care Medicine, Universiti Teknologi MARA,
3 Selangor, Malaysia. Anonymous data will be shared upon request, and it is subjected to the data
4 protection regulations.

6 **Authors' Contributions**

7 All authors have made substantial contributions to this manuscript. ASR, NQ and HN
8 conceptualized and designed the study. ASR and NQ jointly acquired the funding. MAZAH
9 acquired the clinical history and data. ASR, HAH, NB, MSMY, SSK and NQ interpreted and
10 verified the clinical data. HN led the genetic analysis and interpretation. MAZAH and ASR
11 drafted the manuscript. All authors revised this manuscript critically for important intellectual
12 content and approved the final submitted version.

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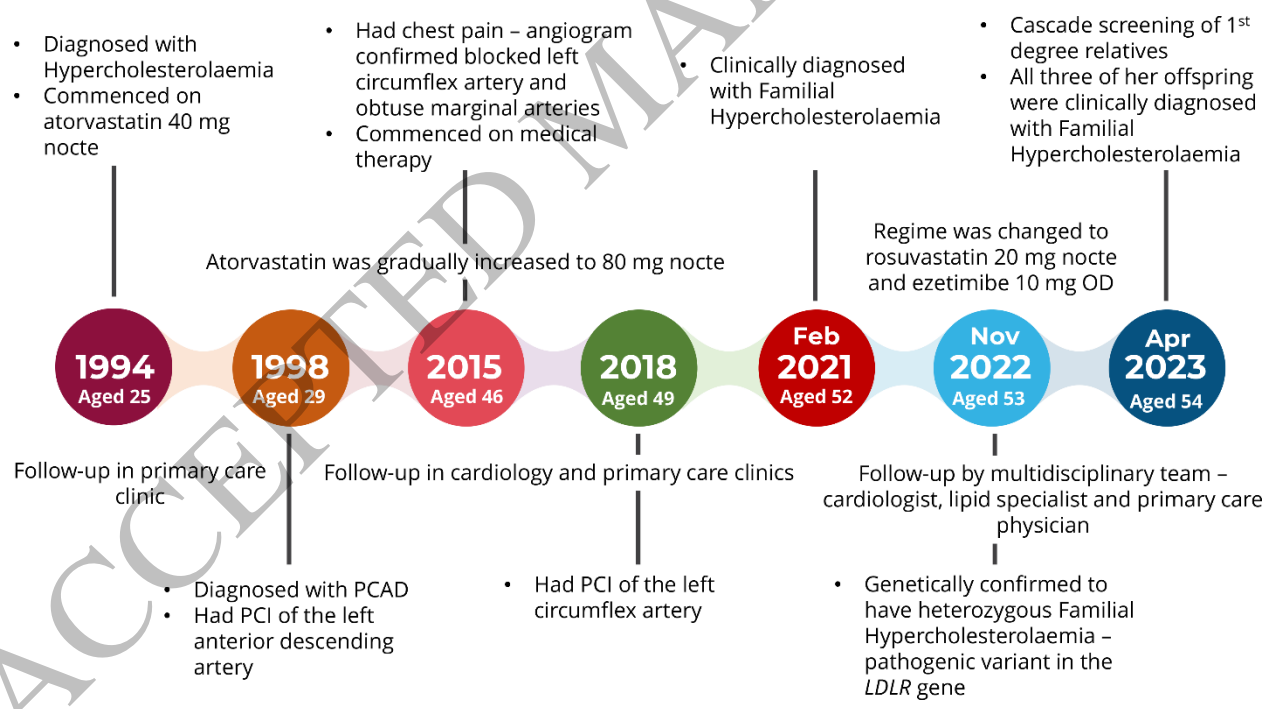


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1 **Learning Points:**

- 2 • This case highlights a missed opportunity to diagnose FH early in a patient with pre-
- 3 existing PCAD. FH should be clinically diagnosed in primary care to identify those who
- 4 may require genetic testing.
- 5 • A multidisciplinary management of FH, including cascade screening of family members
- 6 is vital to prevent premature ASCVD in this extremely high-risk population.
- 7 • This case supports an urgent call for action to improve FH detection and multisectoral
- 8 management in tandem with the global call to action to reduce the clinical and public
- 9 health burden of FH.



Summary figure
328x179 mm (x DPI)

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1 **Article type:** Grand Round Case Report

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3 **Primary specialties involved other than cardiology:** Primary Care Physicians and Lipid
4 Specialists

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6 **Introduction**

7 Familial Hypercholesterolemia (FH) is an autosomal dominant genetic condition
8 predominantly caused by low-density lipoprotein receptor (*LDLR*), apolipoprotein B (*APOB*), or
9 proprotein convertase subtilisin/kexin type 9 (*PCSK9*) gene mutations [1]. An individual with
10 FH has been exposed to a lifelong elevation of low-density lipoprotein cholesterol (LDL-c) since
11 birth, leading to the development of atherosclerotic cardiovascular disease (ASCVD) [1].
12 Heterozygous FH (HeFH) is common, with a global pooled prevalence of 1 in 303 [2].

13 Worldwide, FH is severely underdiagnosed and undertreated [3, 4], especially in primary
14 care, mainly due to a lack of awareness and knowledge of this condition [5, 6]. Clinically, FH
15 can be diagnosed using the Simon Broome Criteria (SBC) or Dutch Lipid Clinic Network
16 (DLCN) criteria based on a weighted combination of LDL-c level, the presence of premature
17 corneal arcus (<45 years old) and/or tendon xanthomas, a personal or family history of
18 hypercholesterolemia and early-onset ASCVD [3, 4]. In primary care, FH can be clinically
19 detected using the Familial Hypercholesterolemia Case Ascertainment Tool (FAMCAT), based
20 on a risk prediction algorithm developed and validated from primary care databases [7]. These
21 tools can identify those who may require genetic testing, especially when resources are limited
22 [3-5, 7]. In this case report, we present a Malay woman with pre-existing premature coronary
23 artery disease (PCAD) since the age of 29, who was clinically diagnosed with FH at the age of

1 52 and was subsequently confirmed to have a heterozygous pathogenic mutation in the *LDLR*
2 gene at the age of 53 years. The timeline of this case is summarized in the **Summary Figure**.

3 4 **Case Presentation**

5 This 54-year-old Malay woman, an insurance agent, was seen at a primary care clinic in
6 February 2021 when she was 52 years old for a routine follow-up of hypercholesterolemia. She
7 had been on treatment for hypercholesterolemia since the age of 25. Initially, atorvastatin 40 mg
8 *nocte* was commenced, and the dose was gradually increased to 80 mg *nocte* at the time of
9 presentation. There was no history of chronic kidney disease, diabetes or hypothyroidism to
10 suggest a secondary cause for hypercholesterolemia. She was a nonsmoker and did not drink
11 alcohol. She had no history of hypertension or cerebrovascular disease. The Edinburgh
12 Claudication Questionnaire was negative for peripheral vascular disease (PVD) [8]. However,
13 the WHO Rose Angina Questionnaire was positive [9]. She had retrosternal chest pain during
14 exertion, relieved within 5 minutes of taking glyceryl trinitrate 0.5 mg. It occurred once or twice
15 a month. There was no radiation, shortness of breath or any other associated symptoms. Her
16 resting electrocardiogram was normal.

17 This patient was diagnosed with PCAD in 1998 at 29 years of age when she presented to
18 a cardiology clinic complaining of exertional angina and reduced effort tolerance. She
19 subsequently underwent percutaneous coronary intervention (PCI) of the left anterior descending
20 artery. After the intervention, she received follow-up care at a primary care clinic. Unfortunately,
21 she developed another episode of chest pain in 2015 at the age of 46. An angiogram revealed an
22 80% blocked left circumflex artery and obtuse marginal arteries. She was initially treated with
23 medical therapy due to financial constraints but eventually underwent another PCI in 2018 at the

1 age of 49 years. She then continued her follow-up concurrently in the cardiology and primary
2 care clinics.

3 This patient had a strong family history of hypercholesterolemia and PCAD. Both of her
4 parents were treated for hypercholesterolemia. Her father passed away at 63 due to a major
5 adverse cardiovascular event (MACE). Her mother had coronary artery bypass grafting at 72
6 years old. Among her eight siblings, seven were treated for hypercholesterolemia and had either
7 MACE or sudden cardiac death between the ages of 43 and 56 at the time of diagnosis. She has
8 three children who are being investigated for high cholesterol. None of her family members have
9 had genetic testing for suspected FH. Her family pedigree chart is shown in Figure 1.

10 <Please insert Figure 1 here>

11 On examination, she was obese, with a body mass index of 38.4 kg/m². Her blood
12 pressure was 104/74 mmHg. Other vital signs were normal. Bilateral grade 2 corneal arcus were
13 observed (Figure 2), but the patient only noticed them at 52 years old. There was no tendon
14 xanthoma.

15 < Please insert Figure 2 here>

16 The SBC, DLCN score and FAMCAT relative risk score for this patient were deduced
17 based on the clinical history and laboratory investigations that were extracted from her electronic
18 medical record. The highest LDL-c level was 8.0 mmol/L, and the highest total cholesterol (TC)
19 level was 10.7 mmol/L, recorded in October 2020. Therefore, this patient fulfilled the SBC
20 (possible FH), DLCN (score of 8—probable FH), and FAMCAT (relative risk score of 9.51)
21 criteria. She was then offered and counselled for genetic testing, the gold standard for diagnosing
22 FH [10]. Targeted next-generation sequencing of the three FH candidate genes (*LDLR*, *APOB*
23 and *PCSK9*) was conducted [10]. Subsequently, she was confirmed to carry a heterozygous

1 pathogenic variant in the *LDLR* gene (rs769446356) located in intron 2 (noncoding area), in
2 keeping with the American College of Medical Genetics and Genomics (ACMG)
3 recommendation [11]. This patient was then counselled by the primary care physician regarding
4 the genetic diagnosis, the need to intensify her lipid-lowering medication (LLM) and to screen
5 her first-degree relatives. The importance of adherence to lifestyle modification and
6 pharmacotherapy was also emphasized.

7 This patient received long-term follow-up care from a multidisciplinary team of primary
8 care physician, cardiologist and lipid specialist. Despite being on atorvastatin 80 mg *nocte*, her
9 LDL-c level was still high at 8.0 mmol/L, and her TC level was also high at 10.7 mmol/L. The
10 cardiology team changed the LLM regime to a combination therapy of rosuvastatin 20 mg *nocte*
11 and ezetimibe 10 mg daily. The lower rosuvastatin dose of 20 mg was chosen instead of 40 mg
12 to minimize the potential side effects of high-intensity statin in this patient. She responded well
13 to the combination treatment, where her LDL-c level decreased to 5.0 mmol/L, and her TC level
14 decreased to 7.6 mmol/L. However, she still failed to achieve the $\geq 50\%$ reduction in LDL-c or
15 the target LDL-c of <1.8 mmol/L as recommended by the international guidelines [12, 13]. Her
16 LLM will be further intensified by the lipid specialist in the subsequent follow-ups to achieve the
17 recommended LDL-c target of <1.8 mmol/L by maximizing rosuvastatin from 20 mg to 40 mg
18 *nocte* before adding an injectable LLM, such as the PCSK9 inhibitors. The possibility that
19 lipoprotein apheresis may be needed in the future was also discussed with the patient if there is
20 an inadequate response to the maximum tolerated dose of LLM [13, 14]. The cost of treatment
21 with PCSK9 inhibitors and lipoprotein apheresis was also discussed, as these treatments are not
22 currently reimbursed by the government health financing system in Malaysia.

1 The primary care physician conducted a cascade screening of her first-degree relatives.
2 All three of her children were found to have elevated LDL-c levels and were clinically diagnosed
3 with FH. They were started on statin monotherapy by the primary care physician and were
4 referred to the lipid specialist for further management and intensification of LLM [13]. The
5 cardiology team was informed of the FH diagnosis in these children. Table 1 summarizes the
6 important key features of this case, and Table 2 summarizes the clinical histories of her three
7 children.

8 **Discussion**

9 FH is rarely detected in primary care due to suboptimal awareness and knowledge among
10 primary care physicians [5, 6]; therefore, it is often underdiagnosed and undertreated [3, 15]. In
11 this case, the patient was clinically diagnosed in primary care and was subsequently confirmed to
12 have a heterozygous *LDLR* pathogenic variant. *LDLR* gene mutations were responsible for 85–
13 90% of genetically confirmed FH in the Asian population, followed by *APOB* and *PCSK9* [4].
14 Lifelong exposure to elevated LDL-c predisposed this patient to PCAD, as she was diagnosed at
15 the age of 29 years old and subsequently had PCI. Unfortunately, FH was not identified at that
16 stage, and her LLM was not intensified to achieve the recommended treatment target [12, 13],
17 leading to the second PCI at the age of 49 years old. Despite having a very strong family history
18 of PCAD, FH was not diagnosed earlier, and cascade screening of family members was not
19 conducted, resulting in a missed opportunity to prevent premature ASCVD [3]. When left
20 untreated, affected men have a 30% chance of a fatal or nonfatal cardiac event by the age of 50,
21 while affected women have a 50% chance of this event by the age of 60 [1].

22 Early diagnosis and intensive treatment significantly improve the prognosis of individuals
23 with FH [3, 16]. The established clinical criteria, such as SBC and DLCN, and the FAMCAT

1 primary care screening tool can be used as a first step to identify those who may require genetic
2 testing, especially when resources are limited [3, 16, 17]. In contrast to developed nations like
3 the United Kingdom, genetic testing is not frequently available or covered by Malaysia's health
4 financing system [17]. Clinically diagnosed individuals with or without a molecular diagnosis
5 should be treated following the guidelines' recommendations [12, 13].

6 This patient was on a combination treatment of rosuvastatin and ezetimibe. Her LDL-c
7 decreased by 37.5% from the highest recorded reading of 8.0 mmol/L to 5.0 mmol/L. The initial
8 goal is to achieve at least a 50% reduction in LDL-c, followed by further reductions to achieve
9 the recommended target of <1.8 mmol/L [12, 13]. The LLM combinations should be increased to
10 the maximum tolerated dose, e.g., rosuvastatin 40 mg and ezetimibe 10 mg, as the majority of
11 heterozygous FH patients can achieve the guideline-recommended LDL-c target with these
12 combinations [13]. If the target is still not achieved, novel non-statin therapies such as inclisiran
13 injection (a PCSK9-interfering mRNA) or bempedoic acid (an adenosine triphosphate-citrate
14 lyase inhibitor) can be considered [12, 13]. Lipoprotein apheresis, lomitapide and evinacumab
15 are indicated for patients with homozygous FH and those with a severe form of heterozygous FH
16 [12, 13]. However, it is worth noting that bempedoic acid is currently unavailable in Malaysia,
17 and the government health financing system does not currently reimburse other new treatments
18 such as PCSK9-inhibitors and lipoprotein apheresis. Patients have to pay out-of-pocket for these
19 treatments, and many private health insurance companies charge exorbitant fees to cover such
20 treatments. Failure to achieve the recommended LDL-c target has been widely reported due to
21 undertreatment of FH [3, 15], which may be attributable to drug costs and availability issues.

22 Once an index case is identified, cascade screening of close relatives should be performed
23 using a combined phenotypic and genotypic strategy to identify affected individuals [18]. In this

1 case, all three of her children were found to have elevated LDL-c levels and were clinically
2 diagnosed with FH, including her 16-year-old daughter. However, genetic testing could not be
3 conducted due to financial constraints. They were referred to the lipid specialist for further
4 management and intensification of the LLM. In her daughter's case, pre-pregnancy counselling
5 should be conducted if she decides to have children in the future. Fertile women with FH require
6 risk reduction, with particular emphasis on safe therapy during pre-conception, pregnancy,
7 childbirth and lactation [19]. Once conception occurs, early referral to the obstetrician is required
8 for close monitoring to ensure a successful pregnancy outcome [19].

9 This patient and her three children receive multidisciplinary management and long-term
10 follow-up care from a primary care physician, lipid specialist and cardiologist. All three of her
11 children were started on statin monotherapy by the primary care physician because ezetimibe is
12 unavailable in government primary care clinics. They were referred to the lipid specialist for
13 intensification of the LLM, which includes a combination of high-dose potent statins with either
14 ezetimibe or PCSK9 inhibitors [12, 13]. Current evidence showed that a combination of high-
15 dose potent statin with ezetimibe outperformed statin monotherapy in reducing the LDL-c, and
16 patients were more likely to achieve their LDL-c target [13]. Therefore, ezetimibe should be
17 made available in the government primary care clinics so as not to delay the intensification of
18 LLM using combination therapy in patients with high ASCVD risk.

19
20 This case highlights a delay in FH diagnosis in patients with pre-existing PCAD, which
21 may be due to the lack of awareness and knowledge of this condition among doctors. Molecular
22 diagnosis was also delayed as genetic testing is not routinely available or covered by Malaysia's
23 national health financing system. The intensification of LLM using combination therapy was

1 also delayed due to the issues of limited drug availability in primary care. Although this patient
2 and her children are currently receiving multidisciplinary management by a primary care
3 physician, lipid specialist and cardiologist; communication between the multidisciplinary teams
4 can be further improved. In conclusion, an urgent call for action to improve FH detection and
5 management in Malaysia is highly needed. This aligns with the global call to action to reduce the
6 clinical and public health burden of FH by adopting public policy recommendations, including
7 awareness, advocacy, screening, testing, diagnosis, treatment, family-based care, registries,
8 research, cost and value [20]. This multisectoral approach is pivotal to prevent premature
9 ASCVD in this extremely high-risk population [20].

10

11 **Statement of informed consent**

12 This patient participated in a study titled 'Reducing Premature Coronary Artery Disease by Early
13 Identification of Familial Hypercholesterolaemia'. Written informed consent was obtained from
14 this patient upon recruitment into the study. The authors also confirmed that written informed
15 consent for submission and publication of this case report including images and associated text
16 has been obtained from the patient in line with the COPE guidance.

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18 **Conflicts of interest**

19 None declared.

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7
8 **Figure legends**

- 9 1. **Figure 1.** Family pedigree chart.
10 2. **Figure 2.** Grade 2 corneal arcus in both eyes.

11
12 **Table legend**

- 13 1. **Table 1.** Clinical summary of the indexed case.
14 2. **Table 2.** Clinical summary for the three offspring of the indexed case.

15

1 **Table 1.** Clinical summary of the indexed case.

| Details | |
|---|--|
| Age (Year) | 54 (born in April 1969) |
| Gender | Female |
| Personal History | |
| Premature coronary artery disease | Yes CAD Had PCI at the age of 29 and 49 years old |
| ROSE Angina Questionnaire | Positive |
| Premature cerebrovascular disease | No |
| Edinburgh Claudication Questionnaire | Negative |
| Chronic kidney disease | No |
| Diabetes | No |
| Hypothyroidism | No |
| Family History | |
| Premature coronary artery disease (Male <55y; Female <60y) | Yes Seven out of eight siblings had either an adverse cardiovascular event or sudden cardiac death at the age of 43 to 56 years old |
| Premature cerebrovascular or peripheral vascular disease (Male <55y; Female <60y) | No |

| | | | |
|--|--|--|---------------------------------|
| Hypercholesterolemia | Yes Both parents and 7 out of 8 siblings | | |
| 1 st -degree relatives with corneal arcus | No | | |
| 1 st -degree relatives with tendon xanthoma | No | | |
| Physical Examinations | | | |
| Blood pressure (mmHg) | 104 / 74 | | |
| Body mass index (kg/m ²) | 38.4 | | |
| Waist circumference (cm) | 115 | | |
| Xanthomas | No | | |
| Premature corneal arcus (<45 years old) | No The patient noticed at the age of 52 years old | | |
| Fasting Serum Lipid | Normal Range | 14.10.2020 - the highest TC and LDL-c ever recorded | 14.11.2022 |
| TC (mmol/L) | < 5.2 | 10.7 | 7.6 |
| LDL-c (mmol/L) | < 1.8 | 8.0 | 5.0 |
| HDL-c (mmol/L) | > 1.0 | 1.8 | 1.7 |
| TG (mmol/L) | < 1.7 | 1.9 | 1.8 |
| Lipid Lowering Medications | Atorvastatin 80 mg <i>nocte</i> | | Rosuvastatin 20 mg <i>nocte</i> |

| | |
|--|---|
| | Ezetimibe 10 mg once daily |
| Other Medications | Valsartan 40mg once daily Acetylsalicylic acid 100mg + glycine 45mg once daily Bisoprolol 7.5mg once daily Isosorbide mononitrate 90mg once daily Glyceryl trinitrate 0.5mg as needed |
| Clinical Diagnostic Criteria | |
| SB Criteria | Possible FH |
| DLCN Score | 8 (Probable FH) |
| FAMCAT Relative Risk Score | 9.51 |
| Mutation | |
| Gene | <i>LDLR</i> (NM_000527.4) |
| Intron | 2 |
| Nucleotide change | c.190+4A>T |
| Chromosome position | chr19:11211025 (GRCh37) |
| dbSNP ID | rs769446356 |
| Type of mutation | Intronic (non-coding area) |
| Pathogenicity of variants based on the ACMG Guidelines [8] | Likely pathogenic Global MAF: 0.00001773 (gnomAD v2.1.1) East Asia MAF: 0.0002005 (gnomAD v2.1.1) |

1 **Abbreviations:** PCAD, premature coronary artery disease; PCI, percutaneous coronary intervention; TC, total
2 cholesterol; LDL-c, low-density lipoprotein cholesterol; TG, triglyceride; HDL-c, high-density lipoprotein
3 cholesterol; SB, Simon Broome; DLCN, Dutch Lipid Clinic Network criteria; FAMCAT, familial
4 hypercholesterolaemia case ascertainment tool; ACMG, American College of Medical Genetics and Genomics.

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1 **Table 2.** Clinical summary for the three offspring of the indexed case.

| Details | Offspring 1 | Offspring 2 | Offspring 3 |
|---|--|-------------------------|-----------------------|
| Age (Year) | 26 (born in March 1996) | 20 (born in April 2002) | 16 (born in Dec 2006) |
| Gender | Male | Male | Female |
| Personal History | | | |
| Premature coronary artery disease | No | No | No |
| ROSE Angina Questionnaire | Negative | Negative | Negative |
| Premature cerebrovascular disease | No | No | No |
| Edinburgh Claudication Questionnaire | Negative | Negative | Negative |
| Chronic kidney disease | No | No | No |
| Diabetes | No | No | No |
| Hypothyroidism | No | No | No |
| Family History | | | |
| Premature coronary artery disease (Male <55y; Female <60y) | Mother had PCAD PCI at the age of 29 and 49 years old | | |
| Premature cerebrovascular or peripheral vascular disease | No | | |

| | | | | |
|--|---------------------|---|------------------|-----------------|
| (Male <55y; Female <60y) | | | | |
| Hypercholesterolemia | | Yes Mother is genetically confirmed to have HeFH – <i>LDLR</i> gene mutation | | |
| 1 st -degree relatives with corneal arcus | | Yes Mother noticed at the age of 52 years old | | |
| 1 st -degree relatives with tendon xanthoma | | No | | |
| Physical Examinations | | | | |
| Blood pressure (mmHg) | | 138 / 74 | 109 / 70 | 112 / 58 |
| Body mass index (kg/m ²) | | 31.4 | 20.5 | 21.6 |
| Waist circumference (cm) | | 89 | 77 | 67 |
| Xanthomas | | No | No | No |
| Premature corneal arcus (<45 years old) | | No | No | No |
| Fasting Serum Lipid | Normal Range | 7.4.2023 | 17.4.2023 | 7.4.2023 |
| TC (mmol/L) | < 5.2 | 10.1 | 8.5 | 8.5 |
| LDL-c (mmol/L) | < 1.8 | 7.8 | 6.4 | 6.2 |
| HDL-c (mmol/L) | > 1.0 | 1.6 | 1.6 | 1.7 |
| TG (mmol/L) | < 1.7 | 1.5 | 1.1 | 0.9 |

| | | | |
|-------------------------------------|------------------------------------|------------------------------------|------------------------------------|
| Lipid Lowering Medications | Atorvastatin 20 mg <i>nocte</i> | Atorvastatin 20 mg <i>nocte</i> | Atorvastatin 20 mg <i>nocte</i> |
| Other Medications | Nil | Nil | Nil |
| Clinical Diagnostic Criteria | | | |
| SB Criteria | Possible FH | Possible FH | Possible FH |
| DLCN Score | 8 (Probable FH) | 6 (Probable FH) | 6 (Probable FH) |

1 **Abbreviations:** PCAD, premature coronary artery disease; PCI, percutaneous coronary intervention; TC, total
2 cholesterol; LDL-c, low-density lipoprotein cholesterol; TG, triglyceride; HDL-c, high-density lipoprotein
3 cholesterol; SB, Simon Broome; DLCN, Dutch Lipid Clinic Network criteria; LLM, Lipid-Lowering Medication

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ACCEPTED MANUSCRIPT

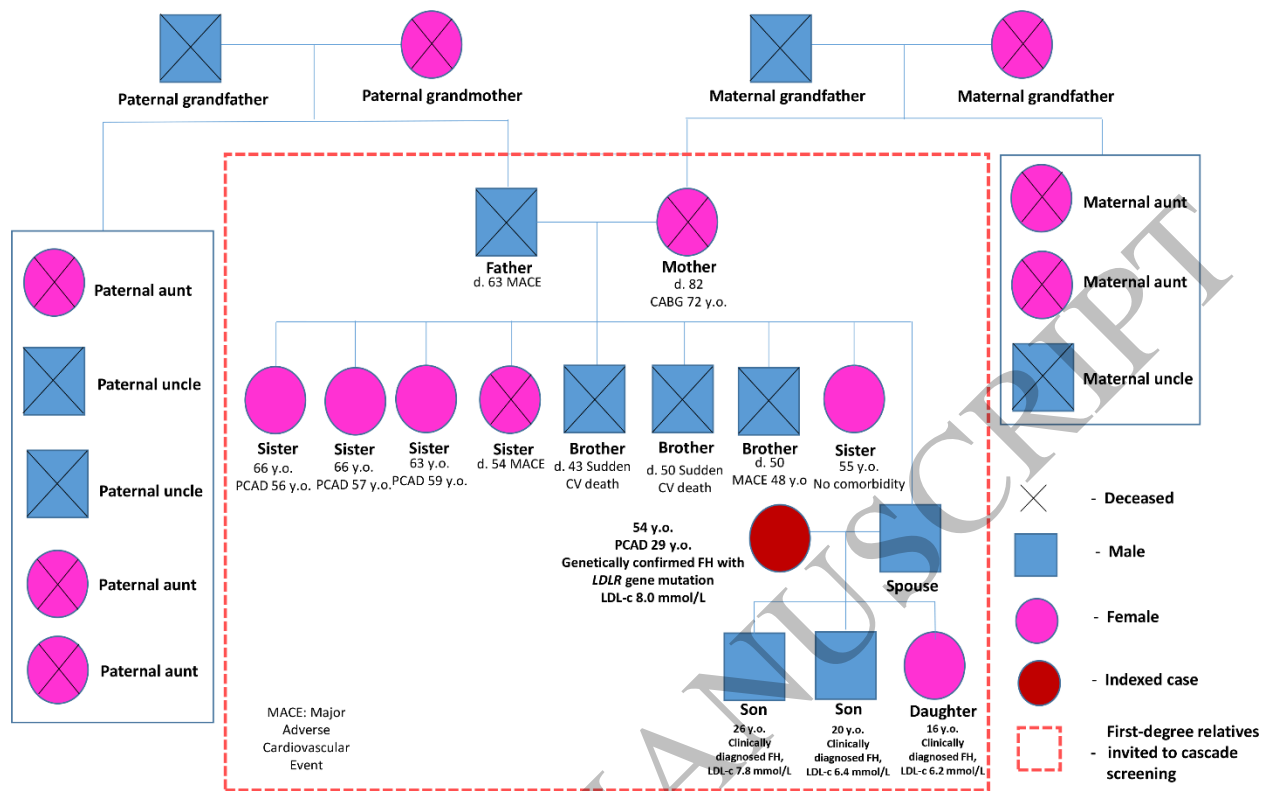


Figure 1
302x189 mm (DPI)



Figure 2
297x111 mm (x DPI)

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