04 SERVICE FLOW

STEP 1

A specimen of 3ml EDTA anticoagulant blood or saliva is collected*



STEP 2

The specimen will be delivered directly to our testing centre for genetic testing



STEP 3

Your health report will be completed in 22 working days



STEP 4

If a disease-causing mutation is found, your family members can undergo genetic testing at a special rate to confirm whether they carry the same mutation

05 ABOUT US

GemVCare - committed to helping the public prevent chronic diseases and improving the health of patients.

With professional genetic testing technology for chronic diseases, medical professionals can customize precise treatments and management models according to your personal genetic profile and health conditions so that you can live your best life

OUR PARTNERS

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General Disclaime

Gemomic® Technology is a patented technology adopted by GemVCare® Ltd. ('GemVCare') to assess the risks of diabeter & chronic diseases and its complications.

As the causes of metabolic diseases and its complications are complex, all related tests require professional interpretation. The Genomics' Technology and any other Cent/Care services mentioned in this document (collectively interpretation. The Genomics' Technology and any other Cent/Care services' shall be used for reference purpose only. The service is not medical advice, diagnosis, therapeutic or prophylactic in general or for any particular individual case or patient and should not be treated as a substitute for professional medical diagnosis, advice, therapeutic or prophylactic. Users are reminded to seek professional advice and shall rely or make decisions based on the Service at their own time.

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FAMILIAL HYPERCHOLESTEROLEMIA TEST



^{*}Please check with your doctor or our service provider for sampling arrangements

[^] Please refer to the terms and conditions on page 3

■ EFFECTS OF FAMILIAL HYPERCHOLESTEROLEMIA (FH)

- **20 times** more likely to have sudden heart attacks and develop coronary arterial disease 20 years earlier
- X Your children have a 50% chance of inheriting the same condition
- **▼ 1** in **200** people has FH ²

General **High Cholesterol**

Familial Hypercholesterolaemia

Cause of disease...

Acquired Factors (E.g. Unhealthy lifestyle)



Congenital gene mutation

(Born with high cholesterol levels)

Life expectancy...

Reduced by 10-15 years ³ (If there are additional risks such as smoking and high blood pressure)



Reduced by 20-30 vears 4

(Even without additional risks)

Risk of heart disease...

Between ages 35-55, risk increases by 40% every 10 years 5



20 times higher than a healthy person 6

How is the disease controlled...

For some patients, it can be controlled by diet and exercise

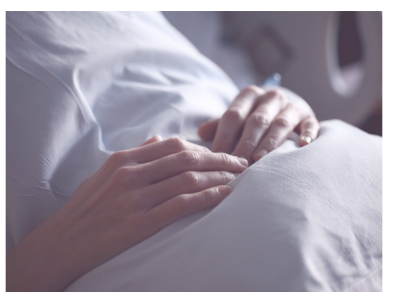


Most patients need medication

Sources:

- 1.What Are the Risks with FH?" March 27, 2020. https://thefhfoundation.org/familialhypercholesterolemia/what-are-the-risks-with-fh.
- Familial Hypercholesterolemia (FH). (n.d.). Retrieved from https://www.heart.org/en/health topics/cholesterol/causes-of-high-cholesterol/familial-hypercholesterolemia-fh
- 3.Clarke, R., Emberson, J., Fletcher, A., Breeze, E., Marmot, M., & Shipley, M. J. (2009). Life expectancy in relation to cardiovascular risk factors: 38 year follow-up of 19 000 men in the Whitehall study. Bmj. 339(Sep16 3)
- 4.Mortensen, G. L., Madsen, I. B., Kruse, C., & Bundgaard, H. (2016). Familial hypercholesterolaemia reduces the quality of life of patients not reaching treatment targets. Danish medical journal, 63(5), A5224, 5.Ann Marie Navar-Boggan, Eric D. Peterson, Ralph B. D'agostino, Benjamin Neely, Allan D. Sniderman, and
- Michael J. Pencina. Hyperlipidemia in Early Adulthood Increases Long-Term Risk of Coronary Heart Disease. Circulation, January 2015 DOI: 10.1161/CIRCULATIONAHA.114.012477

6 Kim, Y. R. & Han, K. H. (2013). Familial hypercholesterolemia and the atherosclerotic disease. Korea circulation journal, 43(6), 363-367. https://doi.org/10.4070/kcj.2013.43.6.36



At present, the traditional diagnosis of FH is based on testing blood lipid levels, observing physical symptoms and checking the family history. However, this method may not be accurate, and can lead to misdiagnosis. 80% of affected individuals only learn that they have FH after experiencing a serious cardiac event. 1

GENETIC TESTING CAN DETERMINE WHETHER YOU HAVE FH

Please pay attention if you meet the following criteria:





Early detection and treatment can reduce cholesterol and cardiovascular disease. The new generation of PCSK9 (Proprotein convertase subtilisin/kexin type 9) inhibitor reduces cardiovascular disease by up to 60% compared with traditional drugs.

Sources:

- 1.Brumit, M. L. (2013, January 21). Familial Hypercholesterolemia Often Misdiagnosed. Retrieved from https://thefh foundation.org/familial-hypercholesterolemia-is-often-misdiagnosed-by-physicians-unaware-of-by-physicians
- 2.PCSK9 Inhibitors. (2020, June 02). Retrieved from https://thefhfoundation.org/fh-treatments/pcsk9-inhibito

GEMVCAPE GENETIC TESTING



Test 9 up-to-date genes related to FH

(APOR APOF IDIR IDIRAPI PCSK9 ABCG5 ABCG8 STAP1 LIPA)



Comprehensive detection of disease-causing genes: Mutations in genetic sequences and gene copy-number variations

(Includes major gene LDLR)



Double verification of results



Technical Accuracy - confirmed by a local university



Ouality Assurance-GemVCare is a participant of the EMQN* programme



If a disease-causing mutation is found, your family members can undergo genetic testing at a special rate to confirm whether they carry the same mutation[^]

- * EMON: The European Molecular Genetics Quality Network
- ^ Terms and conditions:
- 1. This test only includes the specific likely pathogenic genetic mutation of the first patient, must be confirmed
- and referred by a doctor, and does not include other fees (e.g. doctor consultation fee).

 2. The patient must provide a copy of the first patient's report and a doctor's referral letter to verify the need for testing.

 3. All family members receiving the test must be referred by the same clinic or hospital as the first patient.
- 4.The company reserves the right to make the final decision on the offer and any disputes and can change the terms and conditions of this offer without prior notice.