HFE-Related Hereditary Hemochromatosis

THE MOST COMMON FORM of hereditary hemochromatosis (HHC) is caused by alterations in the HFE gene. This is an autosomal recessive condition with reduced penetrance causing iron overload. HHC is clinically suspected when an individual has BOTH persistently elevated fasting ferritin AND elevated transferrin saturation (TS) levels. If untreated, HHC can cause accumulation of iron in various organ systems causing fatigue, arthralgia in the hands and fingers, bronzing of the skin, arrhythmias, low sex drive, male infertility, diabetes, liver cirrhosis, rarely liver cancer and early death. However, compliance to treatment via routine phlebotomies can arrest the disease progression and allow individuals to live relatively symptom free and have a normal lifespan. Molecular testing for the HFE gene is available to individuals showing biochemical evidence of HHC.

MOST INDIVIDUALS with HHC have two mutations in the HFE gene. The chance to develop iron overload depends on the genotype. Most affected individuals have two copies of the C282Y mutation, which is associated with a 50-90% chance of developing biochemical evidence of the disease. Individuals with one C282Y and one H63D mutations have about a 5% chance of developing the disease. Individuals with two H63D mutations have a very low chance of developing the disease (<1%). Heterozygotes (also called carriers) are not likely to have iron overload as a direct result of their HFE genotype.

ELEVATED FERRITIN in the absence of elevated TS is NOT clinically suggestive of HHC. There are many other causes of elevated ferritin, including fatty liver disease, infection, malignancy, or rheumatologic disease.

If Your Patient Has C282Y/C282Y OR C282Y/H63D Genotype:
Your patient and his/her family members may benefit from genetic counselling. Please refer your patient to genetics. We will discuss familial risks and inheritance of HHC. In the meantime, please refer to Flow Chart #2 for Surveillance and Management recommendations.

If Your Patient Has H63D/H63D, C282Y/Normal, or H63D/Normal:
Because these genotypes are associated with very low risk for HFE-related HHC, we cannot confirm this diagnosis for your patient. A genetics referral is NOT indicated.

IF this patient has elevated ferritin, consider investigating other causes as listed above. You may consider consulting the following services:
- Hematology, if ferritin is elevated but < 1000 ug/L
- Hepatology, if ferritin is > 1000 ug/L OR if there are signs of abnormal liver function (eg. elevated liver enzymes)

General Recommendations for Individuals with Iron Overload
Individuals who are at-risk for iron overload should avoid ingesting excess iron or vitamin C (ie., avoid multivitamins containing these ingredients). Individuals may also opt to donate blood with Canadian Blood Services. Diet modifications include reducing intake of red meat.

For individuals who are at-risk of abnormal liver function, alcohol consumption should be limited. These individuals should also avoid raw shellfish. Consuming raw shellfish increases the risk of exposure to Vibrio vulnificans, a bacterium that can cause life threatening complications in individuals who have liver damage.
Flow Chart #1

ADULT with family history of hereditary hemochromatosis (HHC)

IS the affected individual’s genotype and relation to the current patient known?

YES

What is the Genotype?

C282Y/C282Y

OUR PATIENT is the _____ of the affected individual

Child or Sibling

Parent

All Others

TAKE baseline ferritin & TS if not yet done

OFFER GENETIC TESTING

Refer to Flow Chart #2

C282Y/H63D

OUR PATIENT is the _____ of the affected

Sibling

All Others

TAKE baseline ferritin & TS if not yet done

OFFER GENETIC TESTING

IF Ferritin > 200 & TS >60 or >45 twice

Ferritin > 200, TS > 60 or > 45 twice in 1 mo?

No

Yes

OFFER GENETIC TESTING

Refer to Flow Chart #2

NO

ASK PATIENT to obtain further information. If affected is a sibling, may consider baseline ferritin & TS. Reassess as needed if new information arises.

The patient is NOT likely to be at increased risk for HFE-related HHC. No further work-up indicated. Reassess if symptoms arise.

IF Ferritin < 200 & TS < 60

Monitor ferritin & TS...

• Every 2 years if sibling is C282Y/C282Y
• Every 5 years if sibling is C282Y/H63D

If Ferritin > 200 & TS >60 or 45 twice

• You may consider treating your patient with phlebotomy

IF Ferritin > 200 But TS NOT > 45 twice

Patient’s high ferritin cannot necessarily be explained by HFE-related HHC

• Refer to hematology if ferritin < 1000 or hepatology if ferritin > 1000
Flow Chart #2
ADULT with DNA test results suggestive of hereditary hemochromatosis (HHC)

Genotype?

C282Y/C282Y

- IF both ferritin & TS are elevated, the patient has evidence of HFE-related HHC. Please arrange for your patient to have phlebotomy treatment with ferritin monitoring.
- IF only ferritin is elevated, consider other causes of elevated ferritin.
- IF iron indices are normal, the patient has an increased risk to develop HHC. Monitor ferritin & TS every 2 years
- REFER to hematology if ferritin < 1000 and hepatology if ferritin > 1000 for management.
- REFER PATIENT TO GENETICS
- FULL SIBLINGS: Offer genetic testing, ferritin and TS
- PARENTS: Offer baseline ferritin
- ADULT CHILDREN: Offer genetic testing, ferritin and TS. Note: If there are > 2 children and the partner is available, offer DNA testing to the partner

C282Y/H63D

- IF both ferritin & TS are elevated, the patient has evidence of HFE-related HHC. Please arrange for your patient to have phlebotomy treatment with ferritin monitoring
- IF only ferritin is elevated, consider other causes of elevated ferritin.
- IF iron indices are normal, the patient has a < 5% risk to develop HHC. Monitor ferritin & TS every 5 years
- REFER to hematology if ferritin < 1000 and hepatology if ferritin > 1000 for management
- REFER PATIENT TO GENETICS
- FULL SIBLINGS: Offer genetic testing, ferritin and TS
- PARENTS & ADULT CHILDREN: We do NOT recommend ferritin / TS screening or genetic testing for HHC

All Others: H63D/H63D, H63D/Normal & C282Y/Normal

- This genotype is NOT sufficient to diagnose HHC.
- SEE attached information sheet

This Information is Provided By:
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