Lehigh Valley Health Network LVHN Scholarly Works

Department of Surgery

# The Clue to his Pancreatitis!

Malia Eischen MD Lehigh Valley Health Network, Malia.Eischen@lvhn.org

Dale A. Dangleben MD, FACS Lehigh Valley Health Network, Dale\_A.Dangleben@lvhn.org

Follow this and additional works at: https://scholarlyworks.lvhn.org/surgery

Part of the Other Medical Specialties Commons, and the Surgery Commons Let us know how access to this document benefits you

#### Published In/Presented At

Eischen, M., & Dangleben, D. A. (2011). The Clue to his Pancreatitis!. *LVHN Scholarly Works*. Retrieved from https://scholarlyworks.lvhn.org/surgery/26

This Poster is brought to you for free and open access by LVHN Scholarly Works. It has been accepted for inclusion in LVHN Scholarly Works by an authorized administrator. For more information, please contact LibraryServices@lvhn.org.

# The Clue to his Pancreatitis!

Malia Eischen, MD, Dale Dangleben, MD, FACS, Lehigh Valley Health Network, Allentown, Pennsylvania

## Introduction

Xanthomas are lesions characterized by accumulations of lipid-laden macrophages. They are a common manifestation of lipid metabolism disorders, including hyperliproteinemia (HLP). HLPs have been classified according to 5 major phenotypes described in table 1. These phenotypes are based on specific electrophoretic patterns seen in patients with elevated lipoprotein levels. We present a case of a patient presenting with gallstones and pancreatitis, initially misdiagnosed as gallstone pancreatitis. His skin manifestations in combination with an abnormal lab specimen clued us in to the correct diagnosis.

#### Case

A 28-year-old male presented to the Emergency department with abdominal pain. He had a CAT scan which showed extensive peripancreatic edema and cholelithiasis. His amylase and lipase levels were markedly elevated, but his LFTs were normal. A diagnosis of gallstone pancreatitis was made and the surgical service was consulted. Prior to entering the room, we were cautioned that the patient had a, "nasty" rash. On examination, he was found to have the lesions shown on images 1 and 2. Subsequently, we were called by the lab stating that the laboratory specimen was lipemic (Image 3), and his triglyceride level was >5000 mg/ dL. This sent us to the textbooks, and we were quick to realize that he had tuberoeruptive xanthomas, most likely related to a form of HLP. He was later diagnosed with type III HLP, which was considered the cause of his pancreatitis. Both medicine and endocrine were consulted and the patient was transferred to the ICU.

### Discussion

The patient was managed in the ICU with IV fluids, pain control, gemfibrozil/statin, and tight glycemic control for approximately 3 weeks. He recovered from his pancreatitis and was discharged home.

The plan is for an eventual cholecystectomy, but for now, he requires tight control of his HLP.

He was diagnosed with Type III HLP, a serious disorder, with lifelong consequences of premature vascular disease and recurrent pancreatitis. Drug treatment (such as HMG-CoA reductase inhibitors/other lipid-lowering drugs) is indicated since dietary modifications alone are insufficient in managing these patients.





## Types of Hyperlipoproteinemia

Types	Causes and incidence	Diag
I		
(Frederickson's hyperlipoproteinemia, fat-induced hyperlipemia, idiopathic familial)	<ul> <li>Deficient or abnormal lipoprotein lipase, resulting in decreased or absent post heparin lipolytic activity</li> <li>Relatively rare</li> <li>Present at birth</li> </ul>	<ul> <li>Ch de ho</li> <li>Hig lev</li> <li>Lo</li> <li>Le</li> </ul>
II		
(Familial hyperbetalipoproteinemia, essential familial hypercholesterolemia)	<ul> <li>Deficient cell surface receptors that regulate LDL, degradation and cholesterol synthesis, resulting in increased levels of plasma LDL over joints and pressure points</li> <li>Onset between ages 10 and 30</li> </ul>	<ul> <li>Inc</li> <li>Inc</li> <li>Arr</li> </ul>
ш		
(Familial broad beta disease, dysbetalipoproteinemia, remnant removal disease, xanthomia tuberosum)	<ul> <li>Unknown underlying defect results in deficient conversion of triglyceride rich VLDL to LDL</li> <li>Uncommon: usually occurs after age 20 but can occur earlier in men</li> </ul>	• Ab • Ele • Sli • Hy
IV		
(Endogenous hypertriglyceridemia, hyperbetalipoproteinemia)	<ul> <li>Usually occurs secondary to obesity, alcoholism, diabetes, or emotional disorders</li> <li>Relatively common, especially in middle-aged men</li> </ul>	<ul> <li>Elé</li> <li>Ab inc</li> <li>No</li> <li>Mi</li> <li>Fa</li> <li>Ea</li> </ul>
V		
(Mixed hypertriglyceridemia mixed hyperlipidemia)	<ul> <li>Defective triglyceride clearance causes pancreatitis usually secondary to another disorder, such as obesity or nephrosis</li> <li>Uncommon: onset usually occurs in late adolescence or early adulthood</li> </ul>	• Ch • Ele • Ele

#### References

 Morganroth J, Levy R, Fredrickson D. The biochemical, clinical, and genetic features of type III hyperlipoproteinemia. A nn Intern Med February 1975;82(2):158-174.

 Cameron JL, Capuzzi DM, Zuidema GD, Margolis S. Acute pancreatitis with hyperlipemia: the incidence of lipid abnormalities in acute pancreatitis. Ann Surg 1973 Apr;177(4):483-9. 3. Cameron JL, Crisler C, Margolis S, DeMeester TR, Zuidema GD. Acute pancreatitis with hyperlipemia. Surgery 1971 Jul;70(1):53-61.

4. Gotto, A M Jr. Clinical diagnosis of hyperlipoproteinemia. Am J Med 1983 74(5A):5-9 5. Yadav, D., & Pitchumoni, C. (2003). Issues in hyperlipidemic pancreatitis. J Clin Gastroenterol 36(1), 54-62.

#### gnostic findings

- nylomicrons (very low density lipoprotein (VLDL), low ensity lipotein (LDL), high density lipopro plasma 14 burs or more after last meal
- ghly elevated serum chylomicrons and triglyceride vels, slightly elevated serum cholesterol levels
- ower serum lipoprotein lipase levels
- eukocytosis

creased plasma concentrations of LDL creased serum LDL and cholesterol levels mniocentesis shows increased LDL levels

- onormal serum beta lipoprotein
- evated cholesterol and tryglyeride
- ightly elevated glucose tolerance
- peruricemia
- evated VLDL levels
- phormal levels of triglycerides in plasma: variable crease in serum
- ormal or slightly elevated serum cholesterol levels
- ildly abnormal glucose tolerance
- mily history
- arly coronary artery disease

hylomicrons in plasma

- evated plasma VLDV levels
- evated serum cholesterol and triglyceride levels



A PASSION FOR BETTER MEDICINE.