



Patient ID SA00057552	Patient Name SAMPLEREP, VWD2N	Birth Date 1966-06-10	Gender F	Age 46
Order Number SA00057552	Client Order Number SA00057552	Ordering Physician Client, Client	Report Notes	
Account Information C7028846 DLMP Rochester		Collected 16 May 2013 00:00		

von Willebrand Disease 2N (Normandy)

VWD2N Interpretation

MCR

This individual DOES NOT have the Thr791Met, Arg816Trp or Arg854Gln mutations. The Thr791Met, Arg816Trp and Arg854Gln mutations cause >90% of cases of von Willebrand disease (vWD) Type 2N(Normandy phenotype). The incidence of the individual known mutations causing (vWD) Type 2N is as follows: Thr791Met at 15%; Arg816Trp at 10%; and the Arg854Gln at >70%. This assay will not detect other mutations in the known FVIII binding domain of the vWF gene nor will it detect other mutations in the vWF gene. Consider genetic counseling and counseling of potentially affected family members regarding laboratory testing.

ADDITIONAL INFORMATION

This assay is performed by polymerase chain reaction (PCR) followed by restriction enzyme digestion to detect the three common mutations, Thr791Met (Tai1), Arg816Trp (Aval1) and Arg854Gln (Msp1). Laboratory developed test.

Thr791Met

MCR

Negative

Reference Value
Negative

Arg816Trp

MCR

Negative

Reference Value
Negative

Arg854Gln

MCR

Negative

Reference Value
Negative

VWD2N Reviewed By

MCR

Tammy Bernatz

Received: 17 May 2013 15:50

Reported: 17 May 2013 16:05

Performing Site Legend

Code	Laboratory	Address
MCR	Mayo Clinic Dept. of Lab Med and Pathology	200 First Street SW, Rochester, MN 55905