

Opis choroby *

Definicja

Hereditary thrombophilia due to congenital histidine-rich (poly-L) glycoprotein deficiency is a rare, genetic, coagulation disorder characterized by a tendency to develop thrombosis, resulting from decreased histidine-rich glycoprotein (HRG) plasma levels. Manifestations are variable depending on location of thrombosis, but may include headaches, diplopia, progressive pain, limb swelling, itching or ulceration, and brownish skin discoloration, among others.

Dane

Klasyfikacja

Choroba

Synonimy

Hereditary thrombophilia due to congenital HRG deficiency
Dziedziczna trombofilia z powodu wrodzonego niedoboru HRG

Kod ORPHA

217467

Kod OMIM

613116

Kod ICD10

D68.5

Kod ICD11

3B61.0Y

*Źródło

orphanet