

Opis choroby *

Definicja

A rare autosomal dominant autoinflammatory syndrome characterized by early onset systemic inflammation with autoimmune manifestations and more rarely, humoral immune deficiency and increased production of circulating proinflammatory cytokines, variably manifesting with recurrent oral aphthous ulcers, genital ulcers, arthralgia or arthritis, periodic fever, uveitis, and severe gastrointestinal involvement (pain, diarrhea, vomiting, rectal bleeding).

Dane

Klasifikacja

Choroba

Synonimy

Behçet-like disease due to HA20

Choroba podobna do choroby Behçeta

spowodowana HA20

Choroba podobna do choroby Behçeta

spowodowana haploinsuficjencją A20

Behçet-like disease due to haploinsufficiency of
A20

Kod ORPHA

476102

Kod OMIM

301074

Kod ICD10

D89.8

Kod ICD11

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*Źródło

orphanet