Opis choroby *

Definicja

A rare hereditary angioedema characterized by potentially life-threatening episodes of subcutaneous and/or submucosal edema without urticaria, associated with C1 esterase inhibitor (C1-INH) deficiency. Hereditary angioedema (HAE) type 1 is caused by quantitative, HAE type 2 by qualitative defects of C1-INH. The two subtypes are clinically indistinguishable. Patients may present at any age (but most commonly in childhood) with recurrent attacks of nonpitting edema of the skin, severe abdominal symptoms such as pain and swelling, and/or respiratory distress due to upper respiratory airways involvement. Genital, bladder, muscle, or joint swelling may occur in some cases.

Dane

Klasyfikacja

Synonimy

Choroba

HAE with C1 inhibitor deficiency

HAE z niedoborem C1Inh

HAE z niedoborem inhibitora C1

Dziedziczny obrzęk angioneurotyczny z

niedoborem C1Inh

Dziedziczny obrzęk angioneurotyczny z

niedoborem inhibitora C1 HAE with C1Inh deficiency

Hereditary angioneurotic edema with C1

inhibitor deficiency

Hereditary angioneurotic edema with C1Inh

deficiency

Kod ORPHA

Kod OMIM

Kod ICD10

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D84.1

Kod ICD11

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*Źródło

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