

Chłoniak angioimmunoblastyczny z komórek T

Kod Orpha: 86886 Kod OMIM:

Opis choroby *

Definicja

A rare T-cell non-Hodgkin lymphoma characterized by infiltration of lymph nodes by neoplastic cells of T follicular helper cell origin with a polymorphous inflammatory background including markedly increased follicular dendritic cells and EBV-positive B-cells, as well as prominent proliferation of high endothelial venules. The spleen, liver, skin, and bone marrow are also frequently involved. Patients typically present with generalized lymphadenopathy, hepatosplenomegaly, systemic symptoms, and polyclonal hypergammaglobulinemia. Pruritic skin rash, arthritis, pleural effusion, and ascites may also be observed. The condition is aggressive with generally poor prognosis.

Dane

Klasyfikacja	Synonimy
Choroba	AILT
	AILT
	Chłoniak z komórek T, typ AILD
	Limfadenopatia limfoblastyczna
	Limfogranulomatoza X
	Immunoblastic lymphadenopathy
	Lymphogranulomatosis X
	T-cell lymphoma, AILD type

Kod ORPHA	Kod OMIM	Kod ICD10
86886	-	C86.5

Kod ICD11
2A90.9

*[Źródło](#)

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

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