## **Opis choroby \***

## Definicja

A rare autosomal recessive primary immunodeficiency characterized by susceptibility to Epstein-Barr virus (EBV)-associated lymphoproliferative disorders such as malignant B-cell proliferation, Hodgkin lymphoma, B-cell lymphoma, lymphoid granulomatosis, hemophagocytic lymphohistiocytosis, and smooth muscle tumor. Patients present persistent symptoms of infectious mononucleosis including recurrent febrile episodes, lymphadenopathies, and hepatosplenomegaly, accompanied by high EBV viral load in the blood. Additional manifestations are autoimmune diseases like hemolytic anemia or renal disease.

Dane

<b>Klasyfikacja</b> Choroba	due to ITK deficier Autosomal recess	Autosomal recessive lymphoproliferative disease due to ITK deficiency Autosomal recessive lymphoproliferative disease due to ITK deficiency	
<b>Kod ORPHA</b>	Kod OMIM	<b>Kod ICD10</b>	
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Kod ICD11

\*Źródło

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orphanet