

## Opis choroby \*

### Definicja

A rare non-Langerhans cell histiocytosis characterized by infiltration of lymph nodes or extranodal tissues by non-malignant histiocytes displaying emperipolesis, a non-destructive phagocytosis of lymphocytes or erythrocytes. Most typical presentation is as a massive cervical lymphadenopathy in adolescents and young adults. Most frequent sites of extranodal disease are skin, soft tissue, bones, paranasal sinuses, orbit, salivary glands, and central nervous system. Symptoms are related to mass effect in the affected organs.

### Dane

#### Klasyfikacja

Choroba

#### Synonimy

Destombes-Rosai-Dorfman disease  
Choroba Destombesa, Rosai i Dorfmana  
choroba Rosai, Dorfmana i Destombesa  
Histiocytoza zatokowa z masywną  
limfadenopatią  
SHML  
Rosai-Dorfman-Destombes disease  
SHML  
Sinus histiocytosis with massive  
lymphadenopathy

#### Kod ORPHA

158014

#### Kod OMIM

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#### Kod ICD10

D76.3

#### Kod ICD11

EK92

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#### \*Źródło

orphanet