

## Opis choroby \*

### Definicja

A rare, genetic metabolic disorder characterized by reduced activity of unfractionated serum alkaline phosphatase (ALP) and various symptoms from life-threatening, severely impaired mineralization at birth to musculo-skeletal pain in adulthood.

### Dane

#### Klasyfikacja

Choroba

#### Synonimy

HPP

Choroba Rathburna

Fosfoetanolaminuria

HPP

Phosphoethanolaminuria

Rathbun disease

#### Kod ORPHA

436

#### Kod OMIM

241510

#### Kod ICD10

E83.3

#### Kod ICD11

5C64.3

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#### \*Źródło

orphanet