

## Opis choroby \*

### Definicja

A form of acromelic dysplasia characterized by the distinctive radiological sign of angel-shaped middle phalanges, a typical metacarpophalangeal pattern profile (mainly affecting first metacarpals and middle phalanges of second, third and fifth digits, which all appear short), epiphyseal changes in the hips and, in some, abnormal dentition and delayed bone age.

### Dane

<b>Klasyfikacja</b>	<b>Synonimy</b>
Zespół wad wrodzonych ASPED	ASPED

<b>Kod ORPHA</b>	<b>Kod OMIM</b>	<b>Kod ICD10</b>
63442	105835	Q78.8

**Kod ICD11**  
LD24.8Y

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### \*Źródło

orphanet