

# Dystrofia czopków i pręcików

## Kod Orpha: 1872 Kod OMIM: 616502

### Opis choroby \*

#### Definicja

A rare genetic isolated inherited retinal disorder characterized by primary cone degeneration with significant secondary rod involvement, with a variable fundus appearance. Typical presentation includes decreased visual acuity, central scotoma, photophobia, color vision alteration, followed by night blindness and loss of peripheral visual field.

#### Dane

##### Klasyfikacja

Choroba

**Kod ORPHA**  
1872

**Kod OMIM**  
616502

**Kod ICD10**  
H35.5

**Kod ICD11**  
9B70

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\*Źródło

orphanet

### Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.