Opis choroby *

Definicja

Werner syndrome (WS) is a rare inherited syndrome characterized by premature aging with onset in the third decade of life and with cardinal clinical features including bilateral cataracts, short stature, graying and thinning of scalp hair, characteristic skin disorders and premature onset of additional age-related disorders.

Dane

Klasyfikacja

Choroba

Synonimy Adult progeria

Progeria dorosłych

WS WS

Kod ORPHA

902

Kod OMIM

Kod ICD10

277700 E34.8

Kod ICD11 LD2B

*Źródło

orphanet