

Opis choroby *

Definicja

A rare teratogenic disorder due to acitretin or etretinate exposure during the first trimester of pregnancy, carrying a risk of fetal malformations of approximately 20%, including central nervous system, craniofacial, ear, thymic, cardiac and limb anomalies.

Dane

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| Klasyfikacja | Synonimy |
| Zespół wad wrodzonych | Fetal acitretin/etretinate syndrome |
| | Embriopatia retinoidowa |
| | Płodowy zespół acytretynowy/etretynatowy |
| | Retinoid embryopathy |

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|------------------|-----------------|------------------|
| Kod ORPHA | Kod OMIM | Kod ICD10 |
| 40366 | - | Q86.8 |

Kod ICD11
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*Źródło

orphanet