

Resource Summary Report

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RISCALW

RRID:SCR_009352

Type: Tool

Proper Citation

RISCALW (RRID:SCR_009352)

Resource Information

URL: <http://www.riscalw.uni-hd.de>

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Description: Windows program for risk calculation in families with Duchenne muscular dystrophy. It is based on an extended genetic model which includes germline mosaicism and different new mutation rates depending on sex and mutation type. Arbitrary family structures and additional diagnostic information like genotypes from intragenetic and flanking genetic markers of the dystrophin gene, creatin kinase values and female deletion test results can be taken into account. (entry from Genetic Analysis Software)

Abbreviations: RISCALW

Synonyms: RISk CALculation in Windows

Resource Type: software application, software resource

Keywords: gene, genetic, genomic, delphi 4, ms-windows

Related Condition: Duchenne muscular dystrophy

Resource Name: RISCALW

Resource ID: SCR_009352

Alternate IDs: nlx_154567

Record Creation Time: 20220129T080252+0000

Record Last Update: 20260905T133250+0000

Ratings and Alerts

No rating or validation information has been found for RISCALW.

No alerts have been found for RISCALW.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We have not found any literature mentions for this resource.