

DISEASE:

Familial isolated arrhythmogenic ventricular dysplasia, right dominant form

NAME:	Familial isolated arrhythmogenic ventricular dysplasia, right dominant form
ORPHACODE:	293910
SYNOMYS:	Familial isolated arrhythmogenic ventricular cardiomyopathy, classic form Familial isolated arrhythmogenic ventricular cardiomyopathy, right dominant form Familial isolated arrhythmogenic ventricular dysplasia, classic form
XREF(S):	Orphanet OMIM OMIM OMIM ICD-10 OMIM OMIM

ANALYTE(S):	<u>TTN</u> <u>PKP2</u> <u>RYR2</u> <u>TGFB3</u> <u>DSC2</u> <u>DSG2</u> <u>DSP</u> <u>JUP</u> <u>LDB3</u> <u>LMNA</u> <u>TMEM43</u> <u>CTNNA3</u> <u>CDH2</u>
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Source URL: <http://gentest.healthdata.be/disease/1935>

RELATED CONTENT

Related Genetic Tests

- [Cardiomyopathy, hereditary \(gene panel\)](#)
- [Cardiomyopathy: hypertrophic cardiomyopathy, dilated cardiomyopathy, restrictive cardiomyopathy, left ventricular non-compaction cardiomyopathy, arrhythmogenic right ventricular cardiomyopathy \(gene panel\)](#)
- [Cardiopathies, hereditary \(gene panel\)](#)
- [Primary Electrical disorders / Brugada syndrome / Long QT syndrome \(LQT\) / Short QT syndrome \(SQT\) / Arrhythmogenic right ventricular cardiomyopathy \(ARVC\) / Catecholaminergic polymorphic ventricular tachycardia \(CPVT\) \(gene panel\)](#)

Related Laboratories

- [Centrum Medische Genetica - UZ Antwerpen](#)
- [Centrum Medische Genetica - UZ Brussel VUB](#)
- [Centrum Menselijke Erfelijkhed - KUL](#)

Related Analytes

- [cadherin 2](#)
- [catenin alpha 3](#)
- [desmocollin 2](#)
- [desmoglein 2](#)
- [desmoplakin](#)
- [junction plakoglobin](#)
- [LIM domain binding 3](#)
- [lamin A/C](#)

- plakophilin 2
- ryanodine receptor 2
- transforming growth factor beta 3
- transmembrane protein 43
- titin

Related Gene Panels

- Cardiomyopathy (genepanel) - UZA
- Cardiomyopathy, hereditary (208 genes) - VUB
- Cardiopathies, hereditary (102 genes) - KUL
- Primary Electrical disorders/Brugada syndrome (genepanel) - UZA

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