

ANALYTE:
RAF1

NAME:	Raf-1 proto-oncogene, serine/threonine kinase
SYMBOL:	RAF1
VERSION OF ORPHANET:	2023-06-22 14:14:43
SYNONYMS:	C-Raf proto-oncogene, serine/threonine kinase CRAF Raf-1 c-Raf
XREF(S):	Orphanet Ensembl Genatlas HGNC OMIM Reactome SwissProt
CREATED:	13 May 2019 - 01:01
CHANGED:	22 Jun 2023 - 16:14

RELATED CONTENT

Related Genetic Tests

- [Arteriovenous malformation \(gene panel\)](#)
- [Capillary malformation - arteriovenous malformation \(2 genes\)](#)
- [Cardiomyopathy, hereditary \(gene panel\)](#)
- [Cardiomyopathy: hypertrophic cardiomyopathy, dilated cardiomyopathy, restrictive cardiomyopathy, left ventricular non-compaction cardiomyopathy, arrhythmogenic right ventricular cardiomyopathy \(gene panel\)](#)
- [Cerebral cavernous malformation \(gene panel\)](#)
- [Congenital malformation \(gene panel - 1721 genes\)](#)
- [Congenital malformation gene panel](#)
- [Congenital structural heart defects \(gene panel\)](#)
- [Dermatogenetic panel, severe, rare and hereditary genodermatoses \(gene panel - 394 genes\)](#)
- [Dilated Cardiomyopathy \(Gene panel\)](#)
- [Early onset epileptic encephalopathy \(gene panel - 845 genes\)](#)
- [Epilepsy gene panel](#)
- [Heart / Cardio disorders / Cardiopathy \(gene panel\)](#)
- [Hereditary Spastic Paraparesis \(94 genes\)](#)
- [Hypertrophic cardiomyopathy \(gene panel\)](#)
- [Intellectual disability & Epilepsy \(gene panel\)](#)
- [Intellectual disability \(gene panel\)](#)
- [Intellectual disability \(virtual gene panel\)](#)
- [Neurodevelopmental disorders \(1300 genes\)](#)
- [Neurodevelopmental disorders gene panel](#)
- [Pediatric oncopredisposition \(gene panel\)](#)
- [Primary ciliary dyskinesia \(PCD\) Heterotaxyies \(gene panel\)](#)
- [Primary lymphedema / fetal hydrops \(gene panel\)](#)
- [RASopathy \(gene panel\)](#)
- [Rendu-Osler-Weber disease \(4 genes\)](#)

- [Short Stature \(gene panel\)](#)
- [Short stature/ Growth retardation/ \(gene panel\)](#)
- [Skin disorders \(gene panel\)](#)
- [Venous malformation \(3 genes\)](#)

Related Diseases

- [Familial isolated dilated cardiomyopathy](#)
- [Noonan syndrome](#)
- [Noonan syndrome with multiple lentigines](#)
- [Pilomyxoid astrocytoma](#)

Related Gene Panels

- [Cardiomyopathy \(genepanel\) - UZA](#)
- [Cardiomyopathy, hereditary \(208 genes\) - VUB](#)
- [Congenital malformation \(1721 genes\) - ULB](#)
- [Congenital malformation gene panel - VUB](#)
- [Congenital structural heart defects - UGent](#)
- [Dermatogenetic / severe, rare and hereditary genodermatoses \(394 genes\) - ULB](#)
- [Dilated Cardiomyopathy \(79 genes\) - IPG](#)
- [Early onset epileptic encephalopathy \(845 genes\) - ULB](#)
- [Epilepsy gene panel - VUB](#)
- [Growth retardation/short stature \(genepanel\) - UZA](#)
- [Hereditary Spastic Paraparesis \(94 genes\) - KUL](#)
- [Heterotaxia PCD - UGent](#)
- [Hypertrophic cardiomyopathy \(75 genes\) - IPG](#)
- [Intellectual disability & Epilepsy - UGent](#)
- [Intellectual disability \(gene panel\)](#)
- [Intellectual disability/Epilepsy \(1091 genes\) - ULG](#)
- [Lymphedema / fetal hydrops \(27 genes\) - UCL](#)

- Neurodevelopmental disorders (1300 genes) - ULB
- Neurodevelopmental disorders: developmental delay, intellectual disability, autistic disorders (1162 genes) - VUB
- Pediatric oncopredisposition - UGent
- RASopathy - KUL
- Short Stature (46 genes) - IPG
- Skin disorders - UGent
- Vascular malformations (germline) (38 genes) - UCL
- cardiopathy panel - UGent
- test-test

Source URL: <http://gentest.healthdata.be/analyte/385>