

ANALYTE:
NRAS

NAME:	NRAS proto-oncogene, GTPase
SYMBOL:	NRAS
VERSION OF ORPHANET:	2023-06-22 14:14:43
SYNONYMS:	N-ras
XREF(S):	Orphanet HGNC OMIM Reactome SwissProt Ensembl Genatlas
CREATED:	13 May 2019 - 01:01
CHANGED:	22 Jun 2023 - 16:14

Source URL: <http://gentest.healthdata.be/analyte/386>

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- [Hereditary Spastic Paraplegia \(94 genes\) - KUL](#)

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- Intellectual disability (gene panel)
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- Lymphedema / fetal hydrops (27 genes) - UCL
- Maffucci syndrome (65 genes) - KUL
- Neurodevelopmental disorders (1300 genes) - ULB
- Neurodevelopmental disorders: developmental delay, intellectual disability, autistic disorders (1162 genes) - VUB
- Overgrowth & vascular anomalies (65 genes) - KUL
- Pediatric oncopredisposition - UGent
- Primary immune deficiencies (444 genes) - KUL
- Primary immune deficiencies - UGent
- RASopathy - KUL
- Short Stature (46 genes) - IPG
- Skeletal dysplasia (genepanel) - UZA
- Skin disorders - UGent
- Sturge-Weber syndrome (65 genes) - KUL
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- Vascular malformations (somatic) (19 genes) - UCL
- cardiopathy panel - UGent
- epidermal nevus syndrome (65 genes) - KUL
- test-test

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