

**ANALYTE:**  
**SMARCE1**

<b>NAME:</b>	SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily e, member 1
<b>SYMBOL:</b>	SMARCE1
<b>VERSION OF ORPHANET:</b>	2023-06-22 14:14:43
<b>SYNONYMS:</b>	BAF57
<b>XREF(S):</b>	<a href="#">Orphanet</a> <a href="#">Ensembl</a> <a href="#">Genatlas</a> <a href="#">HGNC</a> <a href="#">OMIM</a> <a href="#">Reactome</a> <a href="#">SwissProt</a>
<b>CREATED:</b>	13 May 2019 - 01:01
<b>CHANGED:</b>	22 Jun 2023 - 16:14

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## RELATED CONTENT

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### Related Genetic Tests

- [Congenital malformation \(gene panel - 1721 genes\)](#)
- [Congenital malformation gene panel](#)
- [Congenital structural heart defects \(gene panel\)](#)
- [Early onset epileptic encephalopathy \(gene panel - 845 genes\)](#)
- [Epilepsy 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 \(gene panel\)](#)
- [Skeletal dysplasia \(gene panel\)](#)

### Related Diseases

- [Coffin-Siris syndrome](#)
- [Familial multiple meningioma](#)
- [Meningioma](#)

### Related Gene Panels

- [Congenital malformation \(1721 genes\) - ULB](#)
- [Congenital malformation gene panel - VUB](#)
- [Congenital structural heart defects - UGent](#)

- Early onset epileptic encephalopathy (845 genes) - ULB
- Epilepsy gene panel - VUB
- Intellectual disability & Epilepsy - UGent
- Intellectual disability (gene panel)
- Intellectual disability/Epilepsy (1091 genes) - ULG
- Neurodevelopmental disorders (1300 genes) - ULB
- Neurodevelopmental disorders: developmental delay, intellectual disability, autistic disorders (1162 genes) - VUB
- Pediatric oncopredisposition - UGent
- Skeletal dysplasia (genepanel) - UZA

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