

ANALYTE:
PDSS2

NAME:	decaprenyl diphosphate synthase subunit 2
SYMBOL:	PDSS2
VERSION OF ORPHANET:	2022-05-01 04:55:23
SYNONYMS:	COQ1B bA59I9.3
XREF(S):	<u>Orphanet</u> <u>Ensembl</u> <u>Genatlas</u> <u>HGNC</u> <u>OMIM</u> <u>Reactome</u> <u>SwissProt</u>
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- [Leigh syndrome with nephrotic syndrome](#)

Related Gene Panels

- [Ataxia \(348 genes\) - ULB](#)
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- [Epilepsy gene panel - VUB](#)
- [Intellectual disability & Epilepsy - UGent](#)

- Intellectual disability (gene panel)
- Intellectual disability/Epilepsy (1091 genes) - ULG
- Nephropathies, hereditary (219 genes) - KUL
- Nephropathy panel - UGent
- Nephrotic syndrome, FSGS, Alport syndrome (76 genes) - IPG
- Neurodevelopmental disorders (1300 genes) - ULB
- Neurodevelopmental disorders: developmental delay, intellectual disability, autistic disorders (1162 genes) - VUB
- Panel Nephro-ULG-V1
- mitochondrial disease, nuclear based (343 genes) - VUB

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